DATA NOTE

Pan-cancer repository of validated natural and cryptic mRNA splicing mutations [version 3; peer review: 2 approved, 1 approved with reservations]

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Abstract
We present a major public resource of mRNA splicing mutations validated according to multiple lines of evidence of abnormal gene expression. Likely mutations present in all tumor types reported in the Cancer Genome Atlas (TCGA) and the International Cancer Genome Consortium (ICGC) were identified based on the comparative strengths of splice sites in tumor versus normal genomes, and then validated by respectively comparing counts of splice junction spanning and abundance of transcript reads in RNA-Seq data from matched tissues and tumors lacking these mutations. The comprehensive resource features 341,486 of these validated mutations, the majority of which (69.9%) are not present in the Single Nucleotide Polymorphism Database (dbSNP 150). There are 131,347 unique mutations which weaken or abolish natural splice sites, and 222,071 mutations which strengthen cryptic splice sites (11,932 affect both simultaneously). 28,812 novel or rare flagged variants (with <1% population frequency in dbSNP) were observed in multiple tumor tissue types. An algorithm was developed to classify variants into splicing molecular phenotypes that integrates germline heterozygosity, degree of information change and impact on expression. The classification thresholds were calibrated against the ClinVar clinical database phenotypic assignments. Variants are partitioned into allele-specific alternative splicing, likely aberrant and aberrant splicing phenotypes. Single variants or chromosome ranges can be queried using a Global Alliance for Genomics and Health (GA4GH)-compliant, web-based Beacon "Validated Splicing Mutations" either separately or in aggregate alongside other Beacons through...

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Competing interests: PKR cofounded and BCS is an employee of CytoGnomix Inc., which hosts the interactive webpage described in this study. CytoGnomix markets subscriptions to and services based on the software that generated the ValidSpliceMut database. EJM has no conflict of interest.

Grant information: PKR is supported by The Natural Sciences and Engineering Research Council of Canada (NSERC) [RGPIN-2015-06290], Canadian Foundation for Innovation, Canada Research Chairs, and CytoGnomix. Compute Canada and Shared Hierarchical Academic Research Computing Network (SHARCNET) provided high performance computing and storage facilities.

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How to cite this article: Shirley BC, Mucaki EJ and Rogan PK. Pan-cancer repository of validated natural and cryptic mRNA splicing mutations [version 3; peer review: 2 approved, 1 approved with reservations] F1000Research 2019, 7:1908 https://doi.org/10.12688/f1000research.17204.3

Amendments from Version 2

1. ValidSpliceMut now quantifies information analyses, expression pattern evidence and allele frequencies for each variant, and uses this information with an algorithm to classify the splicing molecular phenotype of each tumour. This was done to make it easier for the user to understand the information provided by ValidSpliceMut. The following changes have been made:
   - Methods have been rewritten to describe the logic and justification for the classification algorithm.
   - A categorical analysis of the splicing molecular phenotype classifications is reported in ‘Dataset validation and discussion’.
   - Added a new figure (Figure 2), which consists of the workflow diagram of the aforementioned classification algorithm.
   - A new dataset has been added (Dataset 2) which includes 6 histograms illustrating the consistency of the splicing molecular phenotype classifier for the same mutation across increasing numbers of tissues.
   - Figure 1 has been updated as there is now new information provided by ValidSpliceMut to the user: a) Consensus splicing molecular phenotypes of all patients with the mutation; b) A classification evidence bar for each case studied, constructed using a point-based system which considers all evidence types relevant to the given tumor type.

2. A new analysis evaluates expression in the normal control population in genomic regions carrying variants. This involved the analysis of variants that were not statistically significant with Veridical. This was performed completely independently from the previously-described classification. A new table (Table 3) was added indicating the statistics of alternative splicing detected in controls.

3. Descriptions of information theory-based terms were expanded. The website now includes a link to a glossary of terms consisting of all information theory-based terms and terms used to describe evidence of aberrant splicing based on the output of the Veridical program. ValidSpliceMut now also displays a “Fold Change” field, to relate R changes to splicing binding site strength.

Any further responses from the reviewers can be found at the end of the article.

Introduction

Next generation sequencing continues to reveal large numbers of novel variants whose impact cannot be interpreted from curated variant databases, or through reviews of peer-reviewed biomedical literature. This has created a largely unmet need for unequivocal sources of information regarding the molecular phenotypes and potential pathology of variants of unknown significance (VUS); in cancer genomes, such sources are critically needed to assist in distinguishing driver mutations from overwhelming numbers of bystander mutations. VUS classification criteria highlight the limitations in genome interpretation due to ambiguous variant interpretation. Of the 458,899 variant submissions in NCBI’s ClinVar database with clinical interpretations, nearly half (n=221,271) are VUS (as of November 5th 2018). Only 10,784 variants in ClinVar have been documented to affect mRNA splicing at splice donor or acceptor sites, with 1,063 of these being classified as VUS, and cryptic mRNA splicing mutations are not explicitly described. The current ACMG criteria2 for variant pathogenicity prevent clinical classification of most VUS. Functional evidence that VUS either disrupt or abolish expression of genes has been sought to improve classification and provide insight into the roles, if any, of individual VUS in predisposing or causing disease. We present a comprehensive data repository for a relatively common mutation type (cis-acting variants that alter mRNA splicing). Mutations are predicted with information theory-based analyses3, and supported with functional evidence that variants in tumor genomes are specifically associated with abnormally spliced mRNAs that are infrequent or absent in transcriptomes lacking these variants5.

Information theory (IT) has been proven to accurately predict impact of mutations on mRNA splicing, and has been used to interpret coding and non-coding mutations that alter mRNA splicing in both common and rare diseases1,2,4–5. We have described an IT-based framework for the interpretation and prioritization of non-coding variants of uncertain significance, which has been validated in multiple studies involving novel variants in patients with history or predisposition to heritable breast and/or ovarian cancer1,4–5.

The Cancer Genome Atlas (TCGA) Pan-Cancer Atlas (PCA) is a comprehensive integrated genomic and transcriptomic resource containing data from >10,000 tumors across 33 different tumor types10. Here, we utilized IT-based tools for assessment of high quality sequenced variants in TCGA patients, as well as patients from tumor datasets provided by the International Cancer Genome Consortium (chronic lymphocytic leukemia, esophageal adenocarcinoma, malignant lymphoma, pancreatic cancer endocrine neoplasms, as well as liver, ovarian, and renal cell cancers), for their potential impact on mRNA splicing. The accuracy of predicted mutations was evaluated with the algorithm we previously developed4 that compares transcripts from cases carrying these variants with others lacking them. The results of these genome-wide analyses are presented using an online internet resource, ValidSpliceMut, which can also be queried through the Beacon Network12,14.

Methods

TCGA and ICGC data acquisition and processing

Controlled-access data was obtained with permission from the Data Access Committee at NIH for TCGA and from the International Cancer Genomics Consortium. Patient RNA sequencing BAM files (tumor and normal, when available) and their associated VCF files (GRCh37) were initially obtained from the CancerGenomeHub (Cghub). Files were later downloaded through Genomic Data Commons using the GDC Data Transfer Tool (version 1.3.0), as Cghub was decommissioned mid-project. Genomic data from ICGC was downloaded through the Score client (version 1.5.0). Variants in VCF files which did not pass quality control (QC) were not analyzed.
Information analysis and RNA-Seq validation of splicing variants

We used the **Shannon Pipeline software** (SP; which applies IT to rapidly perform high-throughput, in silico prediction of the impacts of variants on mRNA splicing)\(^1\) to analyze all QC-passing variants in VCFs from TCGA and ICGC (>168 million TCGA and >41 million ICGC variants) to evaluate their potential impact on splice site binding strength (changes in information content, \(R\)). \(R\) is a measure of binding site strength; it is related to affinity through the second law of thermodynamics and is measured in bits (a glossary of IT and Veridical terms can be found here). According to Shannon information theory, only sites with \(R\) values exceeding zero bits can be bound. The minimum fold change in affinity is exponentially related to the difference in \(R\) values of wild type and mutant binding sites (\(\geq 2^\Delta R\)). For example, a 3 bit change would result in at least an 8-fold change in binding affinity. Variants which were predicted to strengthen known natural sites or weaken cryptic splice sites were excluded from all subsequent analyses. We also required novel cryptic splice sites to be strengthened by \(\geq 2\) bits (at least 4 fold), and with final strengths \((R_{\text{final}})\) exceeding that of the nearest natural site of the same polarity.

To validate the potential impact of SP-flagged mutations, **Veridical software** analyzed genomic variants (including insertions and deletions) by comparing the RNA-Seq alignment in the region surrounding the variant in the index case with the corresponding interval in control transcriptomes (normal and tumor tissue of the same type) lacking the same variant\(^2\). The Veridical algorithm: a) counts abnormally spliced reads in RNA-Seq data (categorized as either cryptic site use, exon skipping, or intron inclusion [containing or adjacent to the flagged mutation]), b) applies the Yeo-Johnson transformation to these results, and c) determines the null hypothesis probability (p-value) that the transformed read count corresponds to normal splicing. In tumor types where normal controls were not available, a set of RNA-Seq datasets from 100 different normal tissues from TCGA were used (e.g. a combination of 5 tissue types: BRCA, BLCA, LUAD, KIRC, PRAD). Variants not flagged for any particular evidence type (p-value > 0.05) were inconsistent with being splicing mutations, and were considered alternatively spliced and most likely, benign. After this analysis, Veridical validated 341,486 unique mutations for their direct impact on mRNA splicing (Table 1).

The Shannon pipeline-flagged and Veridical-filtered results were combined into a single large table (Dataset 1)\(^3\) that contains the source data for the ValidSpliceMut SQL database and the associated Beacon application.

### Development of the ValidSpliceMut database and Beacon

We created a publicly accessible Application Programming Interface (API) (https://beacon.cytognomix.com) that can be utilized to programmatically query variants passing filter thresholds

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#### Table 1. Unique Flagged Variants by TCGA and ICGC Tumor Tissue Type*

<table>
<thead>
<tr>
<th>TCGA-ACC</th>
<th>TCGA-BLCA</th>
<th>TCGA-BRCA</th>
<th>TCGA-CESC</th>
<th>TCGA-CHOL</th>
<th>TCGA-COAD</th>
<th>TCGA-DLBC</th>
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<tr>
<td>1,717</td>
<td>9,865</td>
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<td>25,822</td>
<td>9,817</td>
<td>7,512</td>
<td>6,036</td>
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<tr>
<td>TCGA-ESCA</td>
<td>TCGA-GBM</td>
<td>TCGA-HNSC</td>
<td>TCGA-KICH</td>
<td>TCGA-KIRC</td>
<td>TCGA-KIRP</td>
<td>TCGA-LAML</td>
</tr>
<tr>
<td>19,361</td>
<td>935</td>
<td>2,840</td>
<td>26,519</td>
<td>6,711</td>
<td>4,892</td>
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<tr>
<td>TCGA-LGG</td>
<td>TCGA-LHIC</td>
<td>TCGA-LUAD</td>
<td>TCGA-LUSC</td>
<td>TCGA-MESO</td>
<td>TCGA-OV</td>
<td>TCGA-PAAD</td>
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<td>1,346</td>
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<td>2,628</td>
<td>303</td>
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<tr>
<td>TCGA-PCPG</td>
<td>TCGA-PRAD</td>
<td>TCGA-READ</td>
<td>TCGA-SARC</td>
<td>TCGA-SKCM</td>
<td>TCGA-STAD</td>
<td>TCGA-TGCT</td>
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<tr>
<td>90</td>
<td>944</td>
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<td>20,024</td>
<td>12,515</td>
<td>20,245</td>
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<tr>
<td>TCGA-THCA</td>
<td>TCGA-THYM</td>
<td>TCGA-UCEC</td>
<td>TCGA-UCS</td>
<td>TCGA-UVM</td>
<td>ICGC-CLLE</td>
<td>ICGC-ESAD</td>
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<tr>
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<td>16,599</td>
<td>28,524</td>
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<td>2,498</td>
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<tr>
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<td>ICGC-MALY</td>
<td>ICGC-OV</td>
<td>ICGC-PACA</td>
<td>ICGC-RECA</td>
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<td>2,818</td>
<td>3,182</td>
<td>4,255</td>
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*The number of Veridical-flagged mutations in each TCGA and ICGC cancer data sets. Variants between multiple tissue types were counted for each category. Variant and RNA-Seq data were provided by The Cancer Genome Atlas Pan-Cancer Analysis Project.\(^1\) TCGA: ACC [Adrenocortical carcinoma], BLCA [Bladder Urothelial], BRCA [Breast Cancer], CESC [Cervical Squamous Cell Carcinoma], CHOL [Cholangiocarcinoma], COAD [Colon Adenocarcinoma], DLBC [Lymphoid Neoplasm Diffuse Large B-cell Lymphoma], ESCA [Esophageal Cancer], GBM [Brain Glioblastoma Multiforme], HNSC [Head and Neck Squamous Cell Carcinoma], KICH [Kidney Chromophobe], KIRC [Kidney Renal Clear Cell Carcinoma], KIRP [Kidney Renal Papillary Cell Carcinoma], LAML [Acute Myeloid Leukemia], LGG [Brain Lower Grade Glioma], LiHC [Liver Hepatocellular carcinoma], LUAD [Lung Adenocarcinoma], LUSC [Lung Squamous Cell Carcinoma], MESO [Mesothelioma], OV [Ovarian Serous Cystadenocarcinoma], PAAD [Pancreatic Cancer], PCPG [Pheochromocytoma and Paraganglioma], PRAD [Prostate Adenocarcinoma], READ [Rectum Adenocarcinoma], SARC [Sarcoma], SKCM [Skin Cutaneous melanoma], STAD [Gastric Adenocarcinoma], TGCT [Testicular Germ Cell Tumors], THCA [Head and Neck Thyroid Carcinoma], THYM [Thymoma], UCEC [Uterine Corpus Endometrial Carcinoma], UCS [Uterine Carcinosarcoma], UVM [Uveal Melanoma]. ICGC: CLLE [Chronic Lymphocytic Leukemia], ESAD [Esophageal Adenocarcinoma], LIRI [Liver Cancer], MALY [Malignant Lymphoma], OV [Ovarian Cancer], PACA [Pancreatic Cancer Endocrine Neoplasms], RECA [Renal Cell Cancer].
described above (Dataset 1)). It was built in accordance with the GA4GH Beacon v1.0.0 specification, which describes a Representative State Transfer (REST) API for genetic data sharing. A Beacon accepts queries using an HTTP request and returns JavaScript Object Notation (JSON). Our Beacon implementation is coded in PHP 7.0 and utilizes a MySQL database (version: 5.7.24) with indexes applied to variant ID, chromosome, and coordinate fields (GRCh37). The returned JSON object reports whether the variant was found within our Beacon dataset as well as metadata including splice site coordinate, splice type, site type, the IT-based measures $R_{\text{natural}}$ and $R_{\text{final}}$, affected individual IDs, tumor type, Veridical evidence by type annotated with significance level, and, if known, the corresponding rsID with its average heterozygosity (dbSNP 150). The metadata for each variant sent to the Beacon Network is a concise subset of available results in our database. It includes the first relevant database entry, meaning that if the variant exists within multiple cases only the first will contribute fields to the metadata. However, among this metadata is a hyperlink to our local website containing results for any remaining tumors.

We developed the website, ValidSpliceMut (example output indicated in Figure 1) to serve as a local interface to our Beacon, allowing users to manually search for a variant, by gene name or genome coordinate range. ValidSpliceMut automatically queries our Beacon, and formats the results of the search, if any. This website provides a complete view of variants, including Veridical-based evidence on all data related to every affected individual. If a variant is associated with multiple splice sites, the user is presented with a brief overview of all affected sites and must select a desired site to continue. To obtain the coordinate of the queried variant in gene-centric notation, a link is provided which queries the Mutalyzer API and generates coordinates for all available transcripts. ValidSpliceMut only reports transcripts for the gene affected by the variant.

A results page presents variant-specific data in tabular format and an expandable list of panels describing the affected cases. Each of these panels contains Veridical output in tabular format for the selected tumor, a link to the tumor metadata at US National Cancer Institute (by querying the GDC API to obtain a UUID which is used to construct a link to the GDC data portal), the predicted molecular phenotype for that case, an Integrative Genome Viewer (IGV) screenshot containing the variant (IGV screenshots are available for selected variants, see below), and a histogram which presents the expression levels of the variant-containing gene compared to all other gene expression levels across a selected normal tissue type (created dynamically using gnuplot 5.0). The tissue expression data is provided by GTEx (downloaded on 10/22/18). However, several tumor types did not have a GTEx equivalent (TCGA: CHOL, DLBC, MESO, READ, SARC, THYM and UVM; ICQC: MALY). The GNF Expression Atlas 2 was downloaded from the UCSC Genome Browser and was used for expression data for both lymph nodes (DLBC; MALY) and the thymus (THYM). For the remaining tissues, expression data from the following studies were obtained from the Genome Expression Omnibus (GEO): GSE76297 (CHOL), GSE2549 (MESO), GSE15781 (READ) GSE44426 (SARC), and GSE44295 (UVM). ValidSpliceMut quantifies information analyses, expression pattern evidence and allele frequencies for each variant. We derived an algorithm to classify the consequences of each case, i.e. a tumour with a particular mutation, based on these properties. The molecular phenotypes assigned can be either (1) aberrantly spliced, (2) likely aberrantly spliced, or (3) result in allele-specific alternative splicing. The thresholds and parameters of this classification system (Figure 2) were first calibrated against the clinically-validated phenotypes of variants in the ClinVar database. ClinVar contains 34,671 variants that are also present in ValidSpliceMut. In ClinVar, 1,948 of these variants are designated as pathogenic (1,180 natural; 768 cryptic) and 26,056 variants as benign (afflicting 9,601 natural and 16,455 cryptic splice sites). We determined if discernible characteristics of these splicing mutations in ValidSpliceMut corresponded to either pathogenic or benign designations in ClinVar. Pathogenic variants in ClinVar were considered to be equivalent to aberrant splice molecular phenotypes in ValidSpliceMut. Benign variants in ClinVar were considered to be equivalent to allele-specific alternative splicing in ValidSpliceMut. From our previous work on information-theory based analysis of splicing mutations, the characteristics evaluated for splice sites associated or altered by each variant included: 1) natural site $\Delta R$ value, 2) cryptic site $\Delta R$, 3) cryptic site $R$ strength relative to the $R$ value of the nearest cognate natural site, 4) the number of supporting evidence types detected by Veridical, as well as their respective p-values, 5) the heterozygosity of variant in dbSNP150, if present in this database, 6) the overall level of expression in the tissue type from which the tumor was derived, and 7) the distance of the cryptic site, if present, from the closest cognate natural splice site. All of the characteristics were assigned a weight between 0 and 1 for each variant; natural and cryptic splicing mutations were examined separately. For characteristics relevant to a particular site type (4 of these characteristics were relevant for natural splice sites, 6 were for cryptic sites), all potential combinations of weights divisible by 0.2 were examined. All characteristics were initially classified as either aberrant or allele-specific alternatively spliced; subsequently, the weighted majority of classifications were computed for each variant in each tumour. The Matthews correlation coefficient (MCC) was chosen to measure adherence to ClinVar classifications for each weighting scheme as it can account for the imbalance in ClinVar between number of variants classified as benign versus those indicated as pathogenic in assessing the performance of each characteristic. Examination of natural splice site variants alone demonstrated that characteristics 1 and 5 together were consistently the best predictors of ClinVar phenotypes (MCC: 0.777) with $\Delta R < -2.7$ bits [aberrant characteristic] and average heterozygosity $\leq 0.002$ [aberrant characteristic] as the optimal thresholds. Other variant characteristics, either individually or in combination with characteristics 1 and 5, had significantly lower concordance with ClinVar designations (for example, inclusion of tissue specific expression levels reduced MCC to 0.424, and inclusion of aberrant expression patterns based on Veridical results decreased this to 0.446).

Variants known to be frequent in dbSNP150 ($>0.01$ average heterozygosity) are initially partitioned by the algorithm into allele-specific alternative splicing phenotype, since they are
Figure 1. Screenshot of ATM:g.108214098G>T Results Provided By ValidSpliceMut Website. (A) At the top of the page, the predicted molecular phenotype of the mutation for all cases is presented (prediction algorithm is shown in Figure 2). Then, the ‘Variant Position’ heading displays the variant of interest in g. notation, and provides a link which queries the Mutalyzer API to obtain the variant coordinate in a gene-centric c. mutation format. Variant-specific and splice-site-specific tabular results are presented under the headings “Splice Site Information” and “Variant Data”. Results are then organized by TCGA and ICGC sample IDs (‘cases’) harboring the mutation within a series of expandable panels. A link is provided to patient tumor metadata on the GDC data portal. Each panel consists of the molecular phenotype classification for that particular sample, and the read counts and p-values for each Veridical evidence type. Significant p-values (≤0.05) are highlighted in bold. Evidence types deemed “strongly corroborating” (Viner et al. 2014) are color coded and correspond to the dynamically generated text appearing above the table. (B) An integrative genome viewer (IGV) image showing alignment of expressed sequence reads. IGV screenshots are provided only for mutations present <1% of population (in dbSNP 150), with ≥5 junction-spanning reads, and are highly significant (p < 0.01) for cryptic splicing, exon skipping, and/or intron inclusion with mutation. A specific IGV screenshot for this sample captures the region surrounding the mutation. Here, several RNA-Seq reads show skipping of the affected exon. (C) A dynamically generated histogram presents expression levels of all genes for a selected normal tissue type. Genes are grouped into bins based on expression level, denoted on the x-axis. The number of genes present in each bin is shown on the y-axis (log10 scale). The histogram key indicates the expression range which contains the variant-containing gene (purple). Tissue type can be changed via a drop-down list.
indistinguishable from germline polymorphisms. Indeed, the ClinVar database indicates that such variants also present in ValidSpliceMut were 182-fold more likely to be benign than pathogenic, in contrast with 3-fold for mutations with <0.01 average heterozygosity. Since variants with benign phenotypes may also be infrequent (heterozygosity ≤ 0.01), a threshold $\Delta R_i$ value that optimally distinguished benign from pathogenic natural splice site variants was determined among those also present in ValidSpliceMut. Variants that decrease natural splice site strength ($\Delta R_i$) by ≥4 bits (or ≥16 fold) nearly completely exclude all benign variants in ClinVar (< 4.7%). The threshold is robust, as the percentage of benign variants steeply decreases from $\Delta R_i < -1$ (44.9%) to $\Delta R_i < -4$ bits (4.7%), then level out with larger reductions in $\Delta R_i$. This criterion prevented many benign variants from being classified as causing either aberrant or likely aberrant molecular phenotypes (Figure 2).

Inferred molecular phenotypes in ValidSpliceMut are corroborated by previously published information theory-based analyses. Natural splice sites containing variants with modestly decreased $R_i$ values (of ≤ 2 bits) do not detectably alter mRNA splicing. Such small changes in $R_i$ values have been consistently associated with benign genetic polymorphisms. Conversely, splicing mutations with $\Delta R_i < -7$ bits result in severe phenotypes in inherited disorders. Activated cryptic splice sites that result in novel mRNA isoforms exhibit $R_i$ values competitive with adjacent natural splice sites of the same polarity. Exonic cryptic splice sites are more likely to be classified as either aberrant or

Figure 2. Evidence-Based Case Classification Flowchart. Flowchart depicting steps taken to classify the molecular phenotype of each case. Cryptic site (left) and natural site changes (right) have differing categorization criteria, which involve the combination of information theory-based predictions and Veridical evidence. SNPs common in dbSNP150 (>0.01 average heterozygosity) are immediately considered allele-specific alternative splicing.
After partitioning variants based on heterozygosity and $\Delta R$ values, the algorithm applies supporting Veridical evidence types to molecular phenotype assignments. Evidence for aberrant splicing deemed “strongly corroborating” (for example, junction read types) is weighted more heavily than other evidence types, thus, an “aberrant” or “likely aberrant” classification is more likely in the presence of a strongly corroborating evidence type with $p$-value $\leq 0.05$. Multiple Veridical evidence types (at least one of which is required to be strongly corroborating) with $p$-values $\leq 0.05$, or the presence of a strongly corroborating evidence type with $\leq 0.005$, reinforces a Veridical prediction of aberrant splicing.

When displaying the predicted molecular phenotype classification for each case, the classifications of other sites affected by the same variant in the same tumour type are also examined. If any of these other sites cause a case to be classified as aberrant, all sites affected by this variant are uniformly classified as aberrant. Reclassification affected a minor fraction of variants in ValidSpliceMut. Slightly over 10-fold more of the reclassified variants were upgraded from likely aberrant to aberrant compared to variants reclassified allele-specific to aberrant, where $n=30,733$ [11% of all aberrant cases] were reclassified as aberrant from likely aberrant, while $n=3,026$ [1% of aberrant cases] were reclassified from allele-specific alternative splicing. The majority of reclassified mutations affect cryptic sites, relative to natural site-altering mutations (11.3% and 0.7% of all aberrant cases, respectively). In the event reclassification has occurred, this is denoted below the classification evidence bar associated with the case.

Although individual cases are assigned according to their distinct molecular phenotypes, there are many combinations of evidence types which can lead to the same phenotypic assignment. During classification, all evidence types are calculated first, then the algorithm applies a series of rules are applied to arrive at an unambiguous classification of each case. This is depicted with a contiguous evidence bar (shown in Figure 1), which linearly arranges the classifications of each variant according to phenotypic severity of the effect on mRNA splicing. The evidence bar is constructed using a point-based system which considers all sources of evidence pertaining to the type of splice site affected by the variant. The position of the bar within a phenotypic classification reflects the balance between evidence supporting and contrasting the assigned classification. Supporting evidence moves the bar to the right, while contrasting evidence moves the bar to the left within a classification category. Equal levels of supporting and contrasting evidence types will cause the evidence bar to appear in the middle of the classification category.

After classifying all cases containing the specified variant (regardless of tumour type), the most frequent classification among all cases, termed the consensus molecular phenotype, is summarized and displayed at the top of the Results page (example in Figure 1). Adjacent to the consensus molecular phenotype, the number of cases assigned the most common classification is indicated alongside the total number of cases. Multiple consensus phenotypes are presented if there is an identical number of cases in each of the classification categories.

To generate IGV images presented on the webpage, a bash script was written to automatically load the RNA-Seq BAM file of a patient with a mutation of interest into IGV, set the viewing window within the region of interest (300nt window, centered on the variant), sorted to bring reads containing the variant of interest to the top of the screen (to increase chance of visualizing mutant splice form), followed by a screen capture. The generation and storage of IGV images for all patient-mutation pairs would be prohibitive due to limitations in time and server space requirements. Therefore IGV images showing evidence of splicing abnormalities were generated only for patient-mutation pairs which met the most stringent criteria: the mutation was required to be flagged for junction-spanning cryptic site use, exon skipping, or intron inclusion (with mutation); the flagged category must include 5 or more reads in this category; if the variant is present in the dbSNP database (release 150), the frequency was required to be $<1\%$ of the population; and the Veridical results, in which the mutations flagged were required to exhibit $p \leq 0.01$ for at least one form of evidence of a splicing abnormality. In some cases, the splicing event observed by Veridical may not be present (or displayed in its entirety) within the image window as the automated procedure used to create these images does not present all evidential sequence reads due to limitations on the number of reads that can be shown. Additionally, reads appearing as exon skipping may instead indicate a pre-existing cryptic site outside of the viewing window (for examples, see Table 2; $FAT1$: g.187521515C>A [c.11641-1G>T] and $SMAD3$: g.67482748C>G [c.1155-3C>G]).

**Dataset validation and discussion**

We have derived a GA4GH-standardized searchable resource for a large set of validated mRNA splicing mutations present in diverse tumor types. All variants passing QC in TCGA and ICGC cancer patients were analyzed with the Shannon pipeline. This revealed that 1,094,749 variants were predicted to have significant impacts on normal mRNA splicing (380,852 natural and 752,472 cryptic splice sites; 38,575 affecting both types). Subsequent RNA-Seq analysis with Veridical provided evidence of abnormal gene expression specifically associated with a subset of these variant(s), identifying 341,486 unique mutations.

The molecular phenotype of each case containing a variant is classified in the ValidSpliceMut database and these results are collated for multiple tumors as the consensus phenotype. The molecular phenotype algorithm for Veridical-flagged natural splice site mutations finds 34.7% (87,382) of cases are classified as ‘aberrant’ mRNA isoforms, while 35.3% (88,993) and 30.0%
Table 2. Representative Validated Splicing Mutations in COSMIC Cancer Gene Census genes.

<table>
<thead>
<tr>
<th>Gene</th>
<th>Splice Mutation</th>
<th>R(bits)</th>
<th>Tumor</th>
<th>Observed Splicing Event</th>
</tr>
</thead>
<tbody>
<tr>
<td>CASC5</td>
<td>15:40942786G&gt;A</td>
<td>4.8 &gt; 1.7</td>
<td>AML</td>
<td>The natural donor site of CASC5 exon 19 (NM_144508.4) is weakened, leading to a significant increase in intron inclusion.</td>
</tr>
<tr>
<td>DNMT3A</td>
<td>2:25467022A&gt;G</td>
<td>3.6 &gt; -3.5</td>
<td>AML</td>
<td>The natural donor site of DNMT3A exon 15 (NM_022552.4) is abolished, resulting in a significant increase in total exon skipping and intron inclusion.</td>
</tr>
<tr>
<td>STAG2</td>
<td>X:123176495G&gt;A</td>
<td>6.5 &gt; 3.5</td>
<td>BLCA</td>
<td>The natural donor of STAG2 exon 6 (NM_006603.4) is weakened, and a significant amount of exon 6 skipping is observed.</td>
</tr>
<tr>
<td>STAG2</td>
<td>X:123200024G&gt;A</td>
<td>19.5 &gt; 8.6</td>
<td>BLCA</td>
<td>The natural acceptor of STAG2 exon 21 (NM_006603.4) is weakened, resulting in a significant increase in exon 21 skipping.</td>
</tr>
<tr>
<td>ATM</td>
<td>11:108214098G&gt;T</td>
<td>8.7 &gt; 5.1</td>
<td>BRCA</td>
<td>A natural donor site is weakened, leading to a significant increase in ATM exon 57 (NM_000515.1) skipping events. Some reads with mutation depict wildtype, leaky splicing.</td>
</tr>
<tr>
<td>BARD1</td>
<td>2:215645882A&gt;T</td>
<td>0.9 &gt; 3.1</td>
<td>BRCA</td>
<td>The mutation strengthens a cryptic site within BARD1 exon 4 (NM_000465.2). Reads which use this activated cryptic site contain the mutation (one exception). Some reads with mutation depict wildtype, leaky splicing.</td>
</tr>
<tr>
<td>GATA3</td>
<td>10:8115701G&gt;C</td>
<td>0.9 &gt; -10.7</td>
<td>BRCA</td>
<td>The mutation abolishes the natural acceptor of GATA3 exon 6 (NM_002051.2). This both increases the use of a pre-existing exonic cryptic splice site (4.2 &gt; 5.6 bits; leads to an 8nt deletion) and significantly increases overall intron inclusion.</td>
</tr>
<tr>
<td>TP53</td>
<td>17:7577609C&gt;T</td>
<td>6.0 &gt; -4.9</td>
<td>BRCA</td>
<td>A natural acceptor site is abolished, activating a cryptic site 49nt upstream (R=5.2 bits) of TP53 exon 7 (NM_005465.5).</td>
</tr>
<tr>
<td>POLD1</td>
<td>19:50920353A&gt;G</td>
<td>8.6 &gt; 6.1</td>
<td>COAD</td>
<td>The natural donor of POLD1 exon 25 (NM_002691.3) is weakened, leading to a significant increase in overall exon skipping.</td>
</tr>
<tr>
<td>SMAD3</td>
<td>15:67482748C&gt;G</td>
<td>119 &gt; 4.0 &gt; 7.7</td>
<td>COAD</td>
<td>This mutation weakens the natural acceptor of SMAD3 exon 9 (NM_005592.3) and predicts a cryptic site that does not appear to be used. A significant number of intron inclusion reads are observed. A distant pre-existing exonic acceptor (9.6 bits; 3598nt from natural acceptor) was used in multiple reads.</td>
</tr>
<tr>
<td>PIK3R1</td>
<td>5:67591246A&gt;G</td>
<td>7.5 &gt; -7.3</td>
<td>GBM</td>
<td>The natural acceptor of PIK3R1 exon 8 (NM_181504.3) is weakened, which promotes a significant increase in exon 8 skipping.</td>
</tr>
<tr>
<td>FAT1</td>
<td>4:187521515C&gt;A</td>
<td>5.3 &gt; -2.4</td>
<td>HNSC</td>
<td>The natural acceptor of FAT1 exon 22 (NM_005245.3) is abolished, resulting in both intron inclusion (overall intron inclusion and the use of a 2.3 bit cryptic site 82nt upstream of natural acceptor) and use of two exonic cryptic sites (237nt and 234nt from the natural acceptor; R=1.0 bits and -0.2 bits, respectively).</td>
</tr>
<tr>
<td>TGFBR2</td>
<td>3:30729875G&gt;A</td>
<td>8.4 &gt; -2.5</td>
<td>HNSC</td>
<td>TGFBR2 exon 6 natural acceptor (NM_003242.5) is abolished, leading to multiple splicing events: intron inclusion, use of three cryptic sites (35nt exonic [R=3.7 bits], 30nt and 97nt intronic [R=0.4 bits and 11.2 bits, respectively]), and exon 6 and 7 skipping (activates a novel pseudo exon -55kb downstream of exon 7).</td>
</tr>
<tr>
<td>PBRM1</td>
<td>3:52682355C&gt;G</td>
<td>6.8 &gt; 2.9</td>
<td>KIRC</td>
<td>The natural donor of PBRM1 exon 8 (NM_018313.4) is weakened, which leads to a significant increase in exon 8 skipping.</td>
</tr>
<tr>
<td>PBRM1</td>
<td>3:52685756A&gt;G</td>
<td>7.7 &gt; 0.7</td>
<td>KIRC</td>
<td>The natural donor of PBRM1 exon 7 (NM_018313.4) is abolished, resulting in a significant increase in exon skipping.</td>
</tr>
<tr>
<td>SETD2</td>
<td>3:47097629T&gt;A</td>
<td>9.8 &gt; 2.1 &gt; 6.4 &gt; 9.0</td>
<td>KIRC</td>
<td>This mutation both significantly weakens the natural acceptor of SETD2 exon 18 (NM_014159.6) while strengthening a 4nt exonic cryptic site, which is used.</td>
</tr>
<tr>
<td>RB1</td>
<td>13:49027249T&gt;A</td>
<td>4.9 &gt; 13.7</td>
<td>LUAD</td>
<td>The natural donor of RB1 exon 18 (NM_003212.1) is abolished, leading to a significant increase in both exon skipping and intron inclusion. All intron inclusion reads contain the mutation of interest.</td>
</tr>
<tr>
<td>RBM10</td>
<td>X:47006900G&gt;T</td>
<td>7.8 &gt; 4.1</td>
<td>LUAD</td>
<td>The natural donor of RBM10 exon 2 (NM_005676.4) is weakened, leading to a significant increase in exon 2 skipping.</td>
</tr>
</tbody>
</table>
The natural acceptor of UVM Splice Mutation
This mutation abolishes the natural donor of PRAD (NM_004396.3), which would lead to a 4nt deletion of exon 4. Note that wildtype splicing is still the dominant isoform observed.

<table>
<thead>
<tr>
<th>Gene</th>
<th>Splice Mutation</th>
<th>$R$(bits)</th>
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</tr>
</thead>
<tbody>
<tr>
<td>RBM10</td>
<td>X:47028898G&gt;T</td>
<td>8.7 &gt; -9.9</td>
<td>LUAD</td>
</tr>
<tr>
<td></td>
<td>(c.201+1G&gt;T)</td>
<td>(Natural Site)</td>
<td></td>
</tr>
<tr>
<td>DDX5</td>
<td>17:62500098 TACAG&gt;T</td>
<td>-3.4 &gt; 5.4</td>
<td>PRAD</td>
</tr>
<tr>
<td></td>
<td>(c.441+2delACAG)</td>
<td>(Cryptic Site)</td>
<td></td>
</tr>
<tr>
<td>PTEN</td>
<td>10:89690802G&gt;A</td>
<td>8.5 &gt; -2.3</td>
<td>PRAD</td>
</tr>
<tr>
<td></td>
<td>(c.210-1G&gt;A)</td>
<td>(Natural Site)</td>
<td></td>
</tr>
<tr>
<td>NRAS</td>
<td>1:115258669A&gt;G</td>
<td>8.1 &gt; 1.1</td>
<td>SKCM</td>
</tr>
<tr>
<td></td>
<td>(c.111+2T&gt;G)</td>
<td>(Natural Site)</td>
<td></td>
</tr>
<tr>
<td>PPP6C</td>
<td>9:127933364C&gt;T</td>
<td>6.7 &gt; 3.7</td>
<td>SKCM</td>
</tr>
<tr>
<td></td>
<td>(c.171G&gt;A)</td>
<td>(Natural Site)</td>
<td></td>
</tr>
<tr>
<td>PPP6C</td>
<td>9:127923119C&gt;G</td>
<td>6.8 &gt; -11.8</td>
<td>SKCM</td>
</tr>
<tr>
<td></td>
<td>(c.237+1G&gt;C)</td>
<td>(Natural Site)</td>
<td></td>
</tr>
<tr>
<td>BAP1</td>
<td>3:52442512T&gt;C</td>
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</tr>
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</tr>
</tbody>
</table>

Example mutations which alter splicing in tumor-associated genes found in patients with these tumor types. Mutations are hyperlinked to their ValidSpliceMut Beacon page, which provides additional material such as IGV images of the RNASeq evidence for the regions of interest. GRCh37 coordinates are indicated.

The vast majority of variant classifications of cases, when present in multiple tumors, are highly consistent across the same and distinct cancer types (Dataset 2)\(^6\). Nearly 75% of all mutations found in two or more tissues had a consistent molecular phenotype between >92% of samples with said mutation. The high proportion of samples with consistent molecular phenotypes across samples remains true when considering mutations that occur more frequently in ValidSpliceMut. For example, 77% of the variants present in >10 distinct tumor types have the same molecular phenotype across 94% of patients with said mutation. The consistency of molecular phenotypes of natural and cryptic splicing mutations suggest pleiotropy for this mutation category across multiple neoplastic tissues. Thus, the effects of splicing mutations on mRNA detected in one cancer diagnosis will likely be similar to those found in other tumour types.

Our results contrast with an earlier TCGA study that investigated alternative mRNA splicing\(^7\) and demonstrated a limited set of non-constitutive exon-exon junctions attributable to cis-acting splicing mutations (n=32). The 3,311 novel or rare variants from TCGA patients that we report specifically activate abnormal cryptic splicing (significant ‘junction-spanning cryptic site use’ reads found by Veridical). This exceeds the number reported in cited study that analyzed all available TCGA tumor transcriptionomes (n=1,914)\(^8\). When ICGC datasets were included, a total of 3,650 variants were found to activate cryptic splicing. Comparing the validated cryptic splicing mutations we found with this previous report\(^9\), 1,176 variants fulfilled our IT-based filtering criteria for constitutive splicing mutations. Veridical validated mRNA splicing effects for 824 of these variants (70.1%). The remaining 738 variants were reanalyzed for changes within the binding sites of regulatory splicing factors (SRF) that might affect normal mRNA processing. Together, including the effects on constitutive splicing, IT analysis of SRFs (SRSF1, SRSF2, SRSF5, SRSF6, hnRNP A1, ELAVL1, PTB and TIA1) cumulatively identified changes in binding strength in 1746 (91.2%) sites affected by these variants.

Veridical validated splicing variants, which we define as mutations, were also tallied by tumor tissue type (Table 1). 38.5% of unique mutations in TCGA (n=131,347) significantly weaken natural splice sites, while 65.0% (n=222,071) strengthen novel or pre-existing cryptic sites. 238,570 of these mutations (69.9%) are absent from dbSNP 150. 72,615 mutations (21.3%) are rare (found in <1% of the population), of which 28,812 (and those not present in dbSNP) were detected in multiple tumor types and cases. Valid mutations lacking rsIDs, by definition, represent either novel or recently observed variants. This low level of dbSNP saturation in TCGA is consistent with the possibility that many currently unknown mRNA splicing mutations may yet be discovered through additional sequencing studies.

### Table 1

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</tr>
</tbody>
</table>
The ValidSpliceMut database consists of variants from both TCGA and ICGC, however the vast majority of variants were sourced from TCGA (329,758 of 341,597 flagged, unique variants were found only in TCGA patients; 96.5%). This had been expected, as the ICGC datasets were smaller (492 patients with available RNAseq data analyzed across 7 tumor types). There were 7,380 Veridical-flagged ICGC variants that were absent from TCGA patients; 4,459 variants were flagged in both TCGA and ICGC datasets (of which, 287 were not found in dbSNP 150). To evaluate the frequency of flagged TCGA and ICGC variants, we compared those shared between the two datasets that met SP criteria (n=9,485). We computed that for, a meaningful comparison (with 95% confidence interval [CI]), at least 9 ICGC and 24 TCGA patients should possess the shared mutation (typically, these correspond to common SNPs) and 1,379 shared variants met this criteria. We determined that on average, a higher average proportion of ICGC patients with shared mutations were flagged by Veridical compared to the TCGA cohort (Figure 3). We observed that a higher fraction of SP-flagged variants are natural splice site alterations in the ICGC dataset compared to TCGA (49.7% to 38.3% of total SP-flagged variants, respectively), with fewer affecting cryptic sites (50.7% to 65.3%, respectively). A similar fraction of these sites were predicted to abolish natural splicing (16.8% ICGC and 14.4% TCGA of total SP-flagged variants). A higher percentage of ICGC variants compared to TCGA were confirmed with Veridical (49.1% to 30.9%, respectively), which may possibly be due to higher overall coverage in these regions in the RNAseq results for ICGC relative to TCGA. Interestingly, the fractions of novel variants i.e. not recorded in dbSNP, between TCGA and ICGC are inconsistent (70.0% vs. 42.4%, respectively). We speculate that this may be related to differences in sequence coverage, since TCGA variants were identified from a mixture of whole genome (WGS) and exomes14, while ICGC variants were exclusively derived by WGS.

To assess the predictive accuracy of SP, we also analyzed an independent set of experimentally-validated splicing mutations that altered exon definition (1,050 variants), originally sourced from the Genome Aggregation Database; gnomAD11 and validated with a high throughput, multiplexed splicing minigene reporter assay12. Significant changes in constitutive splice site and/or (≥ 3 bit) SRF binding site strengths were found with SP for 1017 of these 1050 mutations (96.9%). Based on changes at constitutive splice sites alone, 447 variants were flagged (435 weaken natural sites, and 14 strengthen cryptic sites exceeding that of the most proximate natural splice site of the same polarity).

In Table 2, we highlight a representative subset of validated splicing mutations which were identified in known driver genes implicated in the COSMIC (Catalogue Of Somatic Mutations In Cancer) Cancer Gene Census catalog (CGC)16. In total, 543 ‘Tier 1’ CGC genes exhibited at least one Veridical-flagged variant present in the ValidSpliceMut database. These mutations were associated with either increased exon skipping, intron inclusion, and/or cryptic site use. Mutations in Table 2 are hyperlinked to their corresponding ValidSpliceMut webpage, which provides additional information, including specific expression evidence for each of the tumors in the database that support predictions made by SP.

Many mutations generated multiple types of abnormal read evidence present in mis-spliced transcripts. Interestingly, a subset of TCGA mutations (n=33) produced evidence for every type of abnormal splicing reported by Veridical. Dataset 313 (see Data Availability) describes 11 representative mutations that simultaneously increase exon skipping, intron inclusion, and activate (or significantly increase use of) a strengthened cryptic site. In all but one instance, the mutation weakens the natural site, while simultaneously strengthening an adjacent cryptic site. The only exception involves the gene SAP30BP, where simultaneously occurring mutations in the same read (in linkage disequilibrium; separated by 4 nucleotides) independently cause two separate splicing changes: g.73702087G>A (c.661-1G>A; abolishes the natural acceptor of exon 10) and g.73702091G>A (c.664G>A; creates a weak cryptic acceptor site). The combined splicing impact of these variants is significant exon skipping, intron inclusion, and use of the activated cryptic site.

While ValidSpliceMut was designed to find evidence of variant-directed aberrant splicing, some variants with significant IT changes in splice sites are also associated with naturally-occurring, alternative splicing in controls. These are indicated by Veridical as non-wildtype splice junction reads in the index
case that are not flagged as significantly different from controls (p-value ≥ 0.05). The relatively higher frequencies of these read in control samples is consistent with alternative mRNA splicing. The presence of the splicing mutation could theoretically alter the abundance of alternatively spliced isoforms. However, without evidence that disruption of the balance between constitutive and alternative forms would be deleterious, such variants are considered to exhibit a benign or likely benign molecular phenotype. The pre-filtered Veridical output was analyzed to identify variants that did not meet our p-value threshold to support validation of splicing mutation. We applied increasingly stringent thresholds to the average levels of alternative splicing in controls to identify alternatively spliced exons masquerading as predicted splicing mutations (Table 3). A higher average number of reads in controls indicates that the alternative splicing events are more common in the control samples. Variants within the dinucleotides of intronic sequences at exon-intron boundaries have frequently been assumed to be pathogenic, and this analysis identifies those variants where this assumption may be incorrect. As expected, the most common alternative splicing events masquerading as splicing mutations were associated with exons that are commonly skipped or have reads which cross the exon-intron boundary (Table 3). The ‘junction-spanning cryptic splice site use’ evidence type can only be computed as insignificant when the cryptic site is active in control samples. In all cases where this occurred (N=688), the cryptic site was functional in the wildtype sequence (R_{\text{signal}} > 1.6 bits [R_{\text{rand}}]). These variants represent another type of alternately spliced mRNA via junction-spanning cryptic and exon skipping reads (average of ≥1 read per control), distinct from those in ValidSpliceMut, which have a similar classification, but does not explicitly determine read counts in control samples.

This resource presents a set of splicing abnormalities in which we have the highest confidence because expression validation is required. We anticipate that some correct predictions of the Shannon pipeline may have not been validated by Veridical due to the limitations of mRNA detection; for example, either low expression of the gene harboring the mutation or nonsense-mediated decay of the corresponding transcript could be consistent with the effects of a valid splicing mutation, but in the absence of a sufficient number of abnormal reads, the mutation could not be confirmed. Indeed, expression of genes in tumours with validated splicing mutations exhibited greater overall gene expression than in other variants that were not validated in the same gene. The overall difference in gene expression between these two variant groups was statistically significant for 69.3% of genes based on the Student’s t-test (90% C.I.). Furthermore, overall gene expression in the group with Veridical-flagged variants exceeded the non-flagged group for nearly all other genes (99.7%). Differences between expression levels of Veridical-flagged splicing mutations and other predicted mutations suggests the possibility that unverified SP predictions may arise from lack of or low levels of gene expression of the genes containing this subset of variants.

The original SP version that processed most of the TCGA data did not report regulatory splicing variants which influence exon definition. The Automated Splice Site and Exon Definition Analysis (ASSEDA) server analyses individual variants for regulatory and constitutive IT changes. Due to time constraints, it was not feasible to perform a reanalysis with the upgraded SP of the entire set of ~209 million unique variants present in the TCGA and ICGC datasets. However, the SP upgrade did verify improvements in detection of both constitutive and regulatory splicing mutations using the set of validated mutations reported in a recent study. This version of SP (available through our MutationForecaster system) is capable of predicting mutations in splicing regulatory sequences at higher throughput than ASSEDA and, like ASSEDA, accounts for relative abundance of mRNA isoforms by exon definition.

The Validated Splicing Mutation resource should significantly contribute to reducing the number of outstanding VUS in tumor (and possibly some germline) genomes, and substantially increases the number of functional variants with previously unappreciated consequences to mRNA splicing, in particular, those which activate cryptic splice sites. In our earlier study, a subset of the TCGA breast cancer patient data was evaluated with IT-based tools, identifying 988 mutations as significantly altering

<table>
<thead>
<tr>
<th>Type of Junction Spanning mRNA Read Evidence</th>
<th>Average Number of Reads per Set of Control Samples</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cryptic Splice Site Use</td>
<td>≥5</td>
</tr>
<tr>
<td>Exon Skipping Reads</td>
<td>313</td>
</tr>
<tr>
<td>Intron Inclusion</td>
<td>14,937</td>
</tr>
<tr>
<td>Intron Inclusion with Variant Present</td>
<td>232,276</td>
</tr>
</tbody>
</table>

1 Standard deviation of control sample read average vary significantly between each variant. In some cases, a significant proportion of control samples will not have the read-type of interest despite a high average.

2 Includes variants that were not flagged by Veridical
normal splicing by Veridical (19% of total mutations flagged by IT). This database greatly expands the size of the repository. Here, a higher ratio of rare or novel mutations have been validated by Veridical (31% of total mutations were flagged by IT). The higher yields seen here could be related to mutations present in multiple samples from the same tumor type and other tumor tissues, which would be expected to increase the probability of observing abnormally expressed splice forms for the same mutation.

bioRxiv
An earlier version this article is available from bioRxiv: https://doi.org/10.1101/474452

Software availability
Archived code and scripts used as part of this study are available:

Zenodo: Validated Splicing Mutations Beacon API https://doi.org/10.5281/zenodo.1579898

Zenodo: Validated Splicing Mutations Website https://doi.org/10.5281/zenodo.1579822


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Data availability
Underlying data


Zenodo: Dataset 2. Variant Distribution of Majority Molecular Phenotype Classifications Relative to All Classifications. Supplementary figure containing six histograms (generated with GraphPad Prism v6.0). A variant may occur in multiple individuals, tissue types, or splice sites. Histograms present the fraction of molecular phenotype classifications which are consistent among all cases for variants found in at least 2, 3, 5, 10, 15, and 20 different tissue types. DOI: https://doi.org/10.5281/zenodo.3375706

Zenodo: Dataset 3. Mutations which lead to multiple types of aberrant splicing. Representative set of mutations which significantly alter splicing in all evidence types analyzed by Veridical (i.e. cryptic splice site use, exon skipping, intron inclusion). Mutations are linked to their page on https://validsplicemut.cytognomix.com/, which provides additional material such as RNA-Seq images of the regions of interest. DOI: https://doi.org/10.5281/zenodo.1489941

License: CC0 1.0

Consent
Controlled-access TCGA and ICGC sequence data was approved by NCBI at the US National Institutes of Health (dbGaP Project #988: “Predicting common genetic variants that alter the splicing of human gene transcripts”; Approval Number #13930-11; PI: PK Rogan) and by the International Cancer Genome Consortium (ICGC Project #DACK-105647; “Validation of mutations that alter gene expression”).

Acknowledgements
We acknowledge Coby Viner, Stephanie Dorman, William J. Phillippis and Ujani Hazra for their contributions to the early stages of this project.

References

Open Peer Review

Current Peer Review Status: ✔  ❓  ✔

Version 3

Reviewer Report 28 October 2019

https://doi.org/10.5256/f1000research.22439.r53578

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✔ W.W.M. Pim Pijnappel
Department of Pediatrics, Erasmus University Medical Center (Erasmus MC), Rotterdam, The Netherlands

I am happy with the changes, which have improved the manuscript. The only remaining comment is that I find the genome viewer image not so clear: in the example shown the exon skipping is difficult to see, it would help to show at least two or preferably 3 exons in the figure instead of a part of one exon, if possible.

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Splicing

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Version 2

Reviewer Report 30 May 2019

https://doi.org/10.5256/f1000research.20308.r47751

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❓ W.W.M. Pim Pijnappel
Department of Pediatrics, Erasmus University Medical Center (Erasmus MC), Rotterdam, The Netherlands
The manuscript by the Rogan group potentially provides an interesting resource. I would like to suggest to make it more accessible for the user, because in its current form it is very difficult to judge how reliable and strong the splicing predictions and RNAseq data in reality are. Our lab has checked some variants that are know to affect splicing, and it is very difficult to judge how this resource rates these. For example, Ri before and after mutation are difficult to interpret, it is unclear from this manuscript what these values in practice mean. The text in figure 1 that describes the reads under individuals (bottom of A) does not always correspond to the IGV screenshot in B. It is also unclear what figure C actually shows. The term anti-cryptic has not been defined. It is also not clear to me to what extent the predictions were confirmed for the entire set and whether certain types of predictions were more accurate than others. The resource may be very valuable, I would recommend to make it more accessible and understandable for the biological and diagnostic researcher, who could greatly benefit from it.

Is the rationale for creating the dataset(s) clearly described?
Partly

Are the protocols appropriate and is the work technically sound?
Partly

Are sufficient details of methods and materials provided to allow replication by others?
Partly

Are the datasets clearly presented in a useable and accessible format?
Partly

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Splicing

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

Author Response 23 Aug 2019

Peter Rogan, CytoGnomix Inc., London, Canada

Authors: We are grateful to the reviewer for these insightful comments, which we addressed below, in the manuscript and the ValidSpliceMut database.

Reviewer: The manuscript by the Rogan group potentially provides an interesting resource. I would like to suggest to make it more accessible for the user, because in its current form it is very difficult to judge how reliable and strong the splicing predictions and RNAseq data in reality are.

Response: To make this information more accessible to the user, we have made the
following changes to the main text and to the ValidSpliceMut resource:

- Added a sentence which defines $R_i$ (see the next point)
- Added a “fold change” field to ValidSpliceMut, so users can understand how an information ($R_i$) change is related to splice site binding strength (originally defined in reference #9).
- Added a Glossary of terms to ValidSpliceMut containing all information theory-based terms ($R_i$, natural splicing) and Veridical terms (i.e. junction spanning, read abundance, etc) on the website, which were originally defined in the primary publication describing this algorithm and software (Viner et al., 2014).

The publication of the ACMG/AMP guidelines for interpretation of gene variants has accustomed users to simplified categorical characterizations, which has simplified clinical reporting. It should be recognized that a surprising number of variants in these databases have been reclassified recently, due to discordant interpretations and depositing of additional cases in ClinVar and other databases.

We have developed an algorithm which classifies the molecular phenotypes of the variants reported in our database. Note that we do not attempt to perform clinical classification, since the variants we analyzed cannot be definitively causatively linked to the cancer phenotypes of the tumours from which they were derived. Our algorithm performs evidence-based case classification (view additional Methods and the Figure 2 flowchart for details), with thresholds/cutoffs calibrated to the phenotypes reported in ClinVar database shared by ValidSpliceMut. There are three categories: ‘aberrant’, ‘likely aberrant’, or ‘allele-specific alternative splicing’. This system is described in the manuscript in the ‘Methods’ section, and a categorical analysis of these classifications (i.e. proportion of natural site changes categorized as “aberrant”) are reported in the ‘Dataset validation and discussion’ section. We've updated Figure 1 to display the classification results using the same variant example in ValidSpliceMut that was shown in previous versions of this article.

In this version of the manuscript, we also have added an additional analysis of the control population gene expression in which alternate splicing was evident at the site of the mutation (however, these exomes lacked the variant itself). This involved the analysis of variants that did not pass Veridical-based filtering steps (i.e. were not statistically significant). This analysis (presented in the ‘Dataset validation and discussion’ section) was performed completely independently of the previously described, splicing molecular phenotype classification algorithm.

**Reviewer:** Our lab has checked some variants that are know to affect splicing, and it is very difficult to judge how this resource rates these. For example, $R_i$ before and after mutation are difficult to interpret, it is unclear from this manuscript what these values in practice mean.

**Response:** We had not defined $R_i$ here, as it had been described in our prior publications which are cited. However, for completeness, we have added the following statement in the main manuscript:

“$R_i$ is a measure of binding site strength; it is related to affinity through the second law of thermodynamics and is measured in bits (a glossary of IT and Veridical terms can be found
According to Shannon information theory, only sites with $R_i$ values exceeding zero bits can be bound. The minimum fold change in affinity is exponentially related to the difference in $R_i$ values of wild type and mutant binding sites ($\geq 2^{\Delta R_i}$). For example, a 3 bit change would result in at least an 8-fold change in binding affinity.

A glossary of terms to ValidSpliceMut has been included in the current version of the database to assist users in better understand the results of their queries. We also hyperlink to this glossary in the manuscript, where it is first mentioned.

**Reviewer:** The text in figure 1 that describes the reads under individuals (bottom of A) does not always correspond to the IGV screenshot in B.

**Response:** This discrepancy is due to a limitation on the number of reads which IGV can display at one time. This issue is most prevalent in regions of high coverage. While this is described in the main text, this explanation is given in the context of explaining why some IGV images may not display the splicing event at all.

“In some cases, the splicing event observed by Veridical may not be present within the image window as the automated procedure used to create these images does not present all evidential sequence reads due to limitations on the number of reads that are shown.”

This text has been edited to handle both instances where the IGV images is incomplete or absent:

“In some cases, the splicing event observed by Veridical may not be present (or displayed in its entirety) within the image window as the automated procedure used to create these images does not present all evidential sequence reads due to limitations on the number of reads that are shown.”

There are also cases where the user may observe more skipping reads than stated by Veridical. As mentioned in the manuscript, IGV images may include reads that appear to be exon skipping but are, in fact, activating cryptic splice site outside of the window range displayed. Although not explicitly mentioned, these false skipping reads could instead arise either from an alternate splice form that skips multiple exons or could be reads for a completely different gene overlapping the region. In these cases, Veridical will properly count only the reads skipping the exon of interest.

**Reviewer:** It is also unclear what figure C actually shows.

**Response:** The Tissue-specific Expression Histogram (Figure 1C) was included on ValidSpliceMut to allow the user to quickly determine the relative expression of variant-containing gene relative to the overall gene expression corresponding to the normal tissue from which the tumour was derived. Expression levels are binned relative to all genes in the original dataset, and the bin containing the variant-containing gene is highlighted. This figure is relevant because the natural level of expressed at very low levels in a particular tumor type (or control tissue) may limit the number of reads observed, even if no mutation
were present. A low basal level of expression impacts the confidence interval of the p-value reported by Veridical. The diagram in Figure 1C, for example, shows that ATM has a TPM (transcripts per kilobase millions) between 8-9 in mammary tissue (marked in purple; data from GTEx), and that this particular range of expression is low to moderate compared to the rest of the breast transcriptome.

We have made slight adjustments to the Figure Legend:
“C) A dynamically generated histogram presents expression levels of all genes for a selected normal tissue type. Genes are grouped into bins based on expression level, denoted on the x-axis. The number of genes present in each bin is shown on the y-axis (log_{10} scale). The histogram key indicates the expression range which contains the variant-containing gene (purple). Tissue type can be changed via a drop-down list.”

Reviewer: The term anti-cryptic has not been defined.

Response: As this is a Data Note, we did not want to overexplain aspects of the Veridical software that had already been described elsewhere. As mentioned above, the manuscript cites the original Veridical paper (Viner et al., 2014), which defines this term:
○ Cryptic site use/anti cryptic site use: “Equivalent to: cryptic corroborating / non-cryptic corroborating reads”
○ Cryptic corroborating reads: “Reads occurring within the expected region for cryptic splicing to occurring (i.e. spliced-in regions). This region is between the variant splice site location and the adjacent (direction dependent) splice junction.”
○ Non-cryptic corroborating reads: "Also termed "anti-cryptic" reads. Not within the expected region, but still within the portion that would be expected to be spliced out, had cryptic splicing occurred."

We also note that the Information icons (i) found next to each Veridical evidence-type in ValidSpliceMut direct the user to Viner et al. which define these terms. Finally, these terms are indicated in the aforementioned glossary on the ValidSpliceMut website.

Reviewer: It is also not clear to me to what extent the predictions were confirmed for the entire set and whether certain types of predictions were more accurate than others.

Response: In general, lower p-values indicate a high likelihood that the frequency of aberrant splicing reads is caused by the variant. A lower p-value means higher confidence, as would a higher total number of aberrant splicing reads used to compute said p-value. While we provide p-values for all evidence types examined by Veridical, Viner et al. (2014) suggests that junction-spanning cryptic site use, exon skipping, intron inclusion (with mutation) and read-abundance intron inclusion are the most powerful evidence of variant-directed, aberrant splicing:
“Not surprisingly, we have found that junction-spanning reads have the greatest value for corroborating cryptic splicing and exon skipping. Even a single such read is almost always sufficient to merit the validation of a variant, provided that sufficient control samples are used. For intron inclusion, both junction-spanning and read-abundance-based reads are useful, and a variant can readily be validated with either, provided that the variant-
containing experimental sample(s) show a statistically significant increase in the presence of either form of intron inclusion corroborating reads.”

**Reviewer:** The resource may be very valuable, I would recommend to make it more accessible and understandable for the biological and diagnostic researcher, who could greatly benefit from it.

**Response:** Thank you. We hope the improvements that we have incorporated in the article and the website have made the resource easier to understand and use.

**Competing Interests:** PKR cofounded and BCS is an employee of CytoGnomix Inc., which hosts the interactive webpage described in this study. CytoGnomix markets subscriptions to and services based on the software that generated the ValidSpliceMut database. EJM has no conflict of interest.

Reviewer Report 21 March 2019

https://doi.org/10.5256/f1000research.20308.r45974

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**Emanuele Buratti**
International Centre for Genetic Engineering and Biotechnology (ICGEB), Trieste, Italy

I approve the amendments from Version 1.

**Is the rationale for creating the dataset(s) clearly described?**
Partly

**Are the protocols appropriate and is the work technically sound?**
Partly

**Are sufficient details of methods and materials provided to allow replication by others?**
Partly

**Are the datasets clearly presented in a useable and accessible format?**
Partly

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** I have more than twenty years experience in the investigation of pre-mRNA
splicing processes and especially their potential connection with a variety of human diseases, both monogenic (Cystic Fibrosis, Pompe Disease, Neurofibromatosis) and polygenic (Amyotrophic Lateral Sclerosis, Frontotemporal Dementia). I am the author of more than 160 research papers in peer-reviewed publications and of several articles in scientific books on these subjects (orcid.org/0000-0002-1356-9074)

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

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Version 1

Reviewer Report 07 January 2019

https://doi.org/10.5256/f1000research.18813.r41665

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Francesca D. Ciccarelli
1 Cancer Systems Biology Laboratory, Francis Crick Institute, London, UK
2 School of Cancer and Pharmaceutical Sciences, King’s College London, London, UK

The paper entitled "Pan-cancer repository of validated natural and cryptic mRNA splicing mutations" by Shirley, Mucaki and Rogan describes an extensive analysis of pancancer somatic variants in samples of the TCGA dataset to identify mutations that affect splicing. To this aim, the authors combine the methods that they previously developed to predict mutations affecting splicing, with a validation of their effect on matched mRNA data from TCGA.

The study is technically sound and follows a line of investigation that has been a long standing interest of the authors. Despite this, I have a number of comments that hopefully will help strengthen the study:

1. The authors write that their IT-based framework to predict slicing variants "has been validated in multiple studies" and they refer to numerous papers. However, in all of them they act as co-authors, showing that their method is mostly used by themselves and their collaborators. This is not necessarily a problem, but it would certainly strengthen the study if the authors would perform a comparative assessment of their performance with other available methods to predict splicing mutations, for example those in dbNSFP. This will provide a less biased interpretation of the final results.

2. Somehow related to the previous point, the authors mention that their results "contrast with another TCGA study that investigated alternative mRNA splicing". In my opinion this point should be further explored: what are the main differences and what is the extent of overlap in concordant predictions? What are the possible reasons for these differences? This is important because the cited paper in Cancer Cell analysed the same dataset of mutations.

3. The authors notice that the number of variants which activate cryptic splicing exceed the
number reported in a recently published study in Cell Reports. Similarly to before: what is the extent of overlap between the two datasets? Stating that a dataset is bigger than another one is not necessarily an indication that it is better.

4. The authors validate ~27% of predicted splicing variants using the mRNA data (351k validated of the 1.2M predicted). This is a surprisingly low fraction. Later in the manuscript, the authors briefly discuss about the possible reasons of such a discrepancy. One of them is the possible occurrence of nonsense mediated decay which will not confirm the mutations because no or very few reads will be detected. However, as the authors acknowledge, the absence of supporting reads only in mutated individuals as compared to the presence of reads in WT sample would be a strong indication of the effective role of these mutations on splicing. This can be quantified from the same RNAseq data and in my opinion should be done.

5. In general, the authors seem to exclude that their prediction method could lead to false positives. Rather they justify the poor overlap with limitations of mRNA detection. If this is the case, this should be quantified and probably a comparison with other prediction methods could help.

6. Of the >351k mutations with an effect on splicing supported by RNA data, only 35 affect CGC genes. Is this only a subset of mutations affecting driver genes or is it the complete list? In the former case, I would suggest that the authors provide the full list as supplementary data. In the latter case, the authors should discuss the implication of such a low number. Considering that there are >700 CGC genes, does it mean that aberrant slicing is very rarely a driver event? Is the overwhelming majority of splicing variants passenger?

Is the rationale for creating the dataset(s) clearly described?
Yes

Are the protocols appropriate and is the work technically sound?
Partly

Are sufficient details of methods and materials provided to allow replication by others?
Yes

Are the datasets clearly presented in a useable and accessible format?
Yes

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Computational cancer genomics

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

Author Response 07 Mar 2019

Peter Rogan, CytoGnomix Inc., London, Canada

We thank the reviewer for their valuable comments. Our responses follow:
1. The authors write that their IT-based framework to predict slicing variants "has been validated in multiple studies" and they refer to numerous papers. However, in all of them they act as co-authors, showing that their method is mostly used by themselves and their collaborators. This is not necessarily a problem, but it would certainly strengthen the study if the authors would perform a comparative assessment of their performance with other available methods to predict splicing mutations, for example those in dbNSFP. This will provide a less biased interpretation of the final results.

Response: We have previously compared our mutation prediction methods with others. Mucaki et al. Hum. Mut. 34:556-65 (2013) showed that IT-based single mutation and exon definition methods performed as well or better than MaxEntScan and Human Splice Finder websites. MaxEntScan computes relative entropy which is similar to IT, except it applies a correction for local base composition (which does not measure free energy, in contrast with IT: J. Theor. Biol. 201:87-92, 1989). Human Splice Finder does not measure changes in binding affinity and its basis is ad hoc. We also reviewed all articles (300+) which have used IT-based tools to predict changes in splicing (Caminsky et al., F1000Research 3:282, 2014). This reference covers the vast majority of studies that used IT-based bioinformatic tools for mutation analysis and, compared results obtained using these tools with other available software. The cited studies included a large proportion of mutations that we, ourselves, did not coauthor, or were analyzed by others, removing an obvious source of bias.

Regarding potential bias in our results, the IT-based position weight matrices (iPWMs) of splice recognition sites that we derived and use are based on a comprehensive set of splice sites spanning all known coding genes (see appendix of Rogan et al. Pharmacogenetics & Genomics 13(4):207-18, 2003). Other bioinformatic methods for splice site detection are based on many fewer splice sites for PWMs and are much more likely to be subject to bias based on how those sites were chosen. Also, the determination of information content in natural and mutated splice sites obeys the second law of thermodynamics (Schneider, J. Theor. Biol. 189:427-41, 1997); information contents have been formally proven to be related to binding affinities of splice site to splicesomes and splicing factors. MaxEntScan differs from IT because it applies a correction for local base composition, which is energetically and biochemically irrelevant to binding site affinity, and is therefore, biased.

As comparisons of IT with other methods have been covered previously, and the F1000Research Data Note article format is intended to specifically present results using the new resource we describe, in our opinion, reevaluation of other algorithms would not add anything of value to this manuscript.

2. Somewhat related to the previous point, the authors mention that their results "contrast with another TCGA study that investigated alternative mRNA splicing". In my opinion this point should be further explored: what are the main differences and what is the extent of overlap in concordant predictions? What are the possible reasons for these differences? This is important because the cited paper in Cancer Cell analysed the same dataset of mutations.

Response: The paper being referred to in the reviewer's comment is Kahles et al., Cancer Cell., 34(2):211–224.e62018. This paper reports: “In a joint analysis of cis and trans
associations with 50% prior on each type, we identified 32 cis- and seven trans-sQTL (Bonferroni corrected p < 0.05). Information regarding these cis-sQTLs can be found here: https://api.gdc.cancer.gov/data/3920a044-6874-4049-8010-55f0922243b7.

However, the document that provided does not indicate that these are actual splicing mutations. The document contains SNP coordinates, but not sequence changes. An assessment of known rsIDs at these coordinates only accounted for only 7 of the variants in our database. In fact, none of the substitutions (nor the rsIDs) in the article could be related back to changes in mRNA splice strength. It is not possible to comparison the results presented in this paper to those in Kahles et al. (2018).

3. The authors notice that the number of variants which activate cryptic splicing exceed the number reported in a recently published study in Cell Reports. Similarly to before: what is the extent of overlap between the two datasets? Stating that a dataset is bigger than another one is not necessarily an indication that it is better.

Response: Jayasinghe et al. Cell Rep.;23(1):270-281.e3 (2018) identified 2056 variants in TCGA patients which activated cryptic splice sites, of which 1964 were confirmed by manual review of RNAseq data using the Integrated Genome Viewer (IGV). The data provided (Supplementary Table S1; “Passed Manual Review” tab) was sufficient to perform a comparison with our results. We scanned their manually reviewed variants using the Shannon Pipeline (SP). Despite reporting 1964 manually reviewed mutations, 50 mutations are shared between multiple patients (there are 1914 unique mutations total). 1510 of these mutations were found to alter at least one mRNA splice site (natural or cryptic), of which 1176 met our SP filtering criteria (either decreased natural site strength or cryptic splice sites strengthened by ≥2 bits and exceeding the strength of the nearest natural site of the same polarity).

We considered the possibility that splicing mutations in Jayasinghe et al. that were not flagged by SP could have instead altered the strength of splicing regulatory factor binding sites (SRFs). The high-throughput IT-based variant analysis tools needed to address this question were not available at the time the TCGA genomic data were processed. We recently introduced a new version of SP which is capable of analyzing variants impacting both constitutive splice sites and SRFs (including SRSF1, SRSF2, SRSF5, SRSF6, hnrNPA1, ELAVL1, PTB and TIA1). Upon analysis of the variants in Jayasinghe et al. for IT-derived changes in natural, cryptic and SRF binding site strengths, 1746 of the 1914 (91.2%) were found to be significant. It is conceivable that remaining unclassified variants may affect binding by splicing factors for which we have not yet derived iPWMs (i.e. SRSF7).

Of the 1176 variants meeting our filtering criteria, 824 were flagged by Veridical (70.1%). Interestingly, 27 Veridical-flagged mutations alter the same genomic coordinate in different tumours and another 64 affect the adjacent genomic coordinate within the same splice site in other individuals. We further investigated the remaining mutations (those evaluated, but not flagged) to better understand the discrepancy. In approximately 10% of cases, Veridical did not find any alternative validating cryptic splicing events in the region that contrasted with the read distribution in the set of control transcriptomes. For example,
chr9:35389842G>C in TCGA-CN-5370 was expected to abolish the natural acceptor site of *UNC13B* exon 24, however the read counts for intron inclusion in the RNAseq data were too sparse to be deemed significant (p=0.33).

Jayasinghe *et al.* also found mutations in TCGA patients that were not evaluated in our study. This could occur for either because the RNASeq BAM file for a particular TCGA patient failed to download, or the “key” file that associates the BAM file to its TCGA name was incomplete in some tumors (i.e. TCGA-CG-4436; TCGA-STAD), due to both the BAM file name and header lacking this information. This would not impact the accuracy of the data that is present in our database or that we report, only the level of concordance between our results and those of Jayasinghe *et al.*

When comparing this dataset with our own, we discovered an instance where discrepant RNAseq data for the same tumour in the same TCGA patient led to different IT results. Originally, Veridical did not find a significant splicing change in *POLR1B* exon 14 at the +1 position, a G>A mutation g.113331138G>A, present in patient TCGA-Z6-AAPN in the ESCA dataset. When we manually reviewed the RNAseq BAM file used in this analysis, alternate or exon skipped splice forms were not observed, confirming the results reported from Veridical. In fact, TCGA deposited two separate BAM files containing RNAseq data for this same patient: “TCGA-Z6-AAPN-01A-11R-A406-31_rnaseq.bam” and “UNCID_2681450.b17c4505-8a84-4cdd-8782-fbc456deb2a6.sorted_genome_alignments.bam”. The latter BAM file shows evidence of both predicted exon skipping and activation of a cryptic acceptor site 12 nt downstream of exon 14 of *POLR1B*. Comparison of SNPs between these BAM files in other genes did not show any evidence of sample switching or contamination. Although this issue does not appear to be widespread in the TCGA dataset, such discrepancies exceed the scope of our study, and rightfully should be addressed by TCGA.

Nevertheless, this discovery did prompt us to re-analyze the TCGA-ESCA variants through Veridical using the second set of BAM files, and we have now included those results to ValidSpliceMut. In this new set, the previously mentioned *POLR1B* mutation is deemed significant due to increased exon skipping (86 reads showing exon skipping; p=0.0000).

4. The authors validate ~27% of predicted splicing variants using the mRNA data (351k validated of the 1.2M predicted). This is a surprisingly low fraction. Later in the manuscript, the authors briefly discuss about the possible reasons of such a discrepancy. One of them is the possible occurrence of nonsense mediate decay which will not confirm the mutations because no or very few reads will be detected. However, as the authors acknowledge, the absence of supporting reads only in mutated individuals as compared to the presence of reads in WT sample would be a strong indication of the effective role of these mutations on splicing. This can be quantified from the same RNAseq data and in my opinion should be done.

In the revision of this paper, the fraction of validated predicted splicing variants in the ValidSpliceMut database increased from 27% to 31%, as a result of a series of improvements in the software used for processing. SP was significantly upgraded over the course of this project. The previous iteration would incorrectly report information changes at pre-existing
splice sites adjacent to certain mutations. These sites were characterized by genomic coordinates that were altered by insertions or deletions (indel), regardless of whether the site overlapped or included the sequence change. In such instances, some altered natural splice sites could be designated as cryptic sites. SP also reported changes at cryptic acceptor and donor sites in the first and last exons of a gene, respectively, which were not likely to have a meaningful impact on splicing. Therefore, all datasets processed prior to this upgrade were reanalyzed for indel variants. The TCGA ESCA dataset was reprocessed with a second set of RNAseq BAM files (see above response to point 3), which increased the fraction of flagged mutations for that tumor type. Finally, we processed an additional 7 tumor datasets from ICGC and included the validated mutations in our primary beacon database. The statistics of mutations, their distributions and support have been updated in the present version of the manuscript.

From our perspective, the proportion of variants validated is not “a surprisingly low fraction”. The results reported here are consistent with our previous published studies (Dorman et al., Sci. Rep. 4: 7063, 2014). Aside from NMD, as demonstrated below, some mutations that significantly alter splice site strength may not have been flagged by Veridical as a consequence of low levels of expression of the gene in the tumor (or controls) itself. Furthermore, Veridical cannot make an accurate assessment of the region of interest in control samples if these lack sufficient read abundance levels to determine the probability (p-value) of observing expression in the mutation-containing vs control samples. Also, our analysis did not take into account other impacts of the variant on sequences that influence exon definition, such as binding to splicing regulatory factors or mRNA secondary structure. In our response to point 3, we described a discrepancy in BAM file sources, which could also lead to a lower fraction of confirmed variants. Finally, miscalled variants (despite the stringent quality control criteria applied to select variants) could contribute to the fraction of variants not supported by Veridical analysis. Such technical artifacts have been shown to be quite common in exome sequencing in areas of the genome characterized by low mappability (from Shi et al., Cell Rep. 2018 Nov 6;25(6):1446-1457, 2018): “Examination of the genomic locations of mutations revealed that 41.1% of the artifactual somatic mutations occurred in regions of low mappability compared with only 6.4% for the validated somatic heterogeneous mutations.”

As indicated in our response to point 3, the previous version of this paper only evaluated variants for their impact on constitutional mRNA splice sites and cryptic sites, and excluded impacts of mutations at splicing regulatory factor binding sites (SRFs). The scope and time required to assess SRFs precluded the reanalysis of all datasets for such changes. However, to address the issue raised by the reviewer, we evaluated the degree to which ignoring SRFs would affect the overall discovery of splicing-related variants. The updated version of the Shannon Pipeline (SP) with this capability was used to examine 1050 mRNA splicing variants that have been demonstrated to affect exon recognition (Cheung et al., Mol Cell. 2019 Jan 3;73(1):183-194.e8). These splicing variants were experimentally validated using a high throughput, multiplexed splicing minigene reporter assay in that study. SP reported a change in splicing and/or SRF binding strength for 1017 of these 1050 mutations (96.9%; where change in SRF strength was ≥ 3 bits). After accounting for SRF location (e.g. exonic TIA1 sites were eliminated, since these have not been proven to have splicing effects; see Table 1 of Caminsky et al., F1000Res.;3:282, 2014, for a full description of each SRF), the
number of flagged variants was reduced to 940 (89.5%). Based on changes in constitutive splice site strength alone, 447 variants were flagged (435 weaken natural sites, 14 strengthen cryptic sites to a level exceeding the nearest natural site). Therefore, 46.3% of the constitutive mutations at natural or cryptic splice sites were also flagged by SP. This suggests that the lower predictive accuracy of SP in our original submission was, in part, due to the limitations in its ability to detect pathogenic mutations in SRF binding sites.

We addressed the reviewer's suggestion to compare expression of the same gene in tumours with Veridical-validated mutations with other tumors with SP-predicted mutations in the same gene that were not experimentally validated. To perform the analysis, we obtained pre-processed mRNA expression data from the same RNAseq sources of TCGA patients from cBioPortal (www.cbioportal.com; provisional datasets were used, which contained largest number of patients for each tumor type). We extracted these gene expression values with a software program we wrote that determined transcripts per million (TPM) for each gene containing a SP-flagged variant. Expression values for gene present multiple times (due to the presence of multiple splice isoforms) were averaged for the particular tissue from which they were derived.

We separated mRNA expression values for each gene in TCGA patients into the Veridical-flagged vs. non-validated SP-predicted mutation categories, and performed a Student's t-test on the two groups. The expression values of 58.2% of genes were statistically distinct with 90% confidence; >2 patients per category per gene). With at least 10 patients per category, the number of statistically different genes increased to 69.3%. Among these genes, patients with Veridical-flagged variants had higher overall gene expression in 99.7% of cases. These inherent differences in expression suggest that the failure to validate predicted mutations may be related to little or no expression of these genes in tumour and/or control samples, rather than to accuracy of IT prediction methods. Non-sense mediated decay could be responsible for the decreased expression of these mutated genes in the tumour genomes that carry them, or the failure to validate could be related to low levels of expression of these genes in the particular tissues from which the tumours were derived. This analysis is now described in the revised manuscript (‘Dataset validation and discussion,’ para. 8).

Note: The gene expression data from cBioPortal had some limitations: 4196 genes containing variants flagged by SP are not present in the mRNA expression datasets, though the vast majority occurred in non-coding RNAs (i.e. 145 microRNAs, 194 LINC RNAs) or other uncharacterized RNAs (e.g. 324 ‘LOC’ RNAs). Furthermore, certain TCGA patients that we analyzed were not available in cBioPortal among the available expression datasets (2 TCGA-BRCA, 18 TCGA-COAD, 19 TCGA-HNSC, 1 TCGA-LUAD, 119 TCGA-OV, 4 TCGA-READ, 4 TCGA-STAD, 4 TCGA-THCA, and 7 TCGA-UCEC patients). Nevertheless, sufficient data were obtained for the analysis that we carried out.

Furthermore, this analysis has other caveats, especially regarding the accuracy of RNAseq quantification between samples, even upon normalization of the data. The vast majority of TCGA tumor samples do not have a matched normal counterpart, and many TCGA tumor datasets (e.g. TCGA-LAML) do not provide any normal control RNAseq data at all. While we could perform an analysis in which the tumor is compared to a set of normals for the same...
tissue, variation in expression between different individuals would obfuscate evidence of any apparent NMD caused by the mutation. A 50% reduction of expression (or less) may not be observable under these conditions.

5. In general, the authors seem to exclude that their prediction method could lead to false positives. Rather they justify the poor overlap with limitations of mRNA detection. If this is the case, this should be quantified and probably a comparison with other prediction methods could help.

Response: We have quantified RNA expression in tumors for mutations that were validated by Veridical vs those which were not (see response to point 4). Regarding false positives: An extensive comparison of Information Theory to other bioinformatic programs which evaluate variants for splicing impact (MaxEntScan and Human Splice Finder) has been performed (Mucaki et al. 2013; Caminsky et al. 2014; see response to point 1). False positives are extremely rare because strength of binding sites (in bits) is directly related to their binding affinities; we have demonstrated that unused cryptic splice sites in the vicinity of natural splice sites are significantly weaker. Based on our experience and published analyses of a genome-wide site of binding sites (Rogan et al. 2003), such decoy splice sites are nearly always at least 4 bits (2^4 or 16 fold) weaker than sites that are actually recognized by spliceosomes.

The predictive accuracy of the IT methodology for detecting expression-validated mutations was determined to be 87.9% (762 of 867 variants from 122 different publications; changes to SRFs were included in this variant dataset; Caminsky et al. 2014). This value is similar to the predictions made to those of Cheung et al. (2019), where we predicted splice site and/or SRFs changes to 89.5% mutations experimentally validated to cause exon definition events.

The performance of IT-based methods for predicting splicing mutations has been well established over the past two and a half decades. Re-evaluation of its accuracy is not necessary, and this issue is, at best, only tangentially relevant, to the purpose of presenting the resource described in a Data Note article.

6. Of the >351k mutations with an effect on splicing supported by RNA data, only 35 affect CGC genes. Is this only a subset of mutations affecting driver genes or is it the complete list? In the former case, I would suggest that the authors provide the full list as supplementary data. In the latter case, the authors should discuss the implication of such a low number. Considering that there are >700 CGC genes, does it mean that aberrant splicing is very rarely a driver event? Is the overwhelming majority of splicing variants passenger?

Response: There are 25 CGC genes indicated in Table 2, however these were never intended to be interpreted to be a complete list of CGC genes with Veridical-flagged mutations. The table has been renamed to indicate that these are a set of representative mutations. In the previous version of this paper, the number of variants (“n=25”) did not indicate the total number of CGC genes. This has been removed and replaced with the actual number of CGC splicing mutations that were validated:
“In Table 2, we highlight a subset of validated splicing mutations which were identified in known driver genes implicated in the COSMIC (Catalogue Of Somatic Mutations In Cancer) Cancer Gene Census catalog (CGC) \(^27\). In total, 543 “Tier 1” CGC genes have at least one Veridical-flagged variant present in the ValidSpliceMut database.”

**Competing Interests:** Not applicable.

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**Reviewer Report 12 December 2018**

[Link](https://doi.org/10.5256/f1000research.18813.r41666)

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The increasing amount of sequencing data that is being generated in many biological systems has represented a real challenge to researchers in terms of trying to link individual changes to a particular biological process. The attempt, described in this work, to use IT approaches to evaluate the potential biological significance can significantly contribute to fill this gap. The laboratory of Peter Rogan has a long standing and internationally prominent role in addressing the possible consequences of sequence variants on the pre-mRNA splicing process especially with regards to its connection with human disease. The ValidSpliceMut developed in this work presents a user friendly interface that allows users to manually search for a variant (by gene name or genome coordinate range) and obtain information with regards to its possible effect on splicing. This will greatly help to better appreciate the functionality of Variants of Unknown Significance that are currently abundant genomic and transcriptomic Atlases.

**Is the rationale for creating the dataset(s) clearly described?**
Yes

**Are the protocols appropriate and is the work technically sound?**
Yes

**Are sufficient details of methods and materials provided to allow replication by others?**
Yes

**Are the datasets clearly presented in a useable and accessible format?**
Yes

**Competing Interests:** No competing interests were disclosed.
**Reviewer Expertise:** I have more than twenty years experience in the investigation of pre-mRNA splicing processes and especially their potential connection with a variety of human diseases, both monogenic (Cystic Fibrosis, Pompe Disease, Neurofibromatosis) and polygenic (Amyotrophic Lateral Sclerosis, Frontotemporal Dementia). I am the author of more than 160 research papers in peer-reviewed publications and of several articles in scientific books on these subjects (orcid.org/0000-0002-1356-9074)

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

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**Comments on this article**

**Version 3**

Author Response 29 Oct 2019  
**Peter Rogan**, CytoGnomix Inc., London, Canada

An updated version of Dataset 1 from this article is now available (doi: 10.5281/zenodo.3522192). In this version, a subset of called genomic variants - primarily from the TCGA-OV data - have been eliminated from ValidSpliceMut. The accompanying files in the Dataset indicate the affected variants and describe the rationale for this update.

**Competing Interests:** No competing interests were disclosed.