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WASHINGTON UNIVERSITY IN ST. LOUIS Division of Biology and Biomedical Sciences Molecular Genetics and Genomics

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The Splice is Not Right: Splice-site-creating Mutations in Cancer Genomes

A dissertation presented to The Graduate School of Washington University in partial fulfillment of the requirements for the degree of Doctor of Philosophy

> December 2018 St. Louis, Missouri

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Table of Contents

List of Figuresiv
List of Tablesvi
Acknowledgmentsvii
Abstract of the Dissertationxii
Chapter 1: Introduction
1.1 Cancer Cell Fitness1
1.2 Alternative Splicing and Disease6
1.3 References
Chapter 2: Complex patterns of splice site mutations
2.1 Introduction
2.2 Results
SPLICEINATOR – AUTOMATIC PRIORITIZATION OF SPLICE MUTATION EVENTS
DIVERSE SPLICING PHENOTYPES ASSOCIATED WITH SPLICE SITE MUTATIONS
SPLICE SITE MUTATIONS LINKED TO COMPLEX SPLICING EVENTS
LACK OF MEASURABLE SPLICING DEFECT FOR MANY SPLICE SITE MUTATIONS
IMPACT OF SPLICE SITE MUTATION ON GENE EXPRESSION
NON-CANONICAL SPLICE SITE MUTATION DISCOVERY
2.3 Methods
3.4 Discussion
3.5 References
Chapter 3: Discovery of splice-site-creating mutations
3.1 Introduction
3.2 Results
SPLICE-SITE-CREATING MUTATION DISCOVERY
MOLECULAR AND BIOLOGICAL PATTERNS OF SPLICE SITE-CREATING MUTATIONS
EXPRESSIVITY AND PENETRANCE OF SPLICE-SITE-CREATING MUTATIONS
SPLICE SITE-CREATING MUTATIONS ACROSS GENES AND CANCER TYPES
NEO-ANTIGENS INTRODUCED BY SPLICE SITE-CREATING MUTATIONS
3.3 Methods
3.4 Discussion

3.5 References
Chapter 4: Rare germline and somatic splice-site-creating mutations disrupt exon definition in cancer genomes
4.1 Introduction109
4.2 Results
DATASET DESCRIPTION
GERMLINE AND SOMATIC SPLICE-SITE-CREATING MUTATION DISCOVERY
COMPARATIVE SEQUENCE CONTEXTS AND CHARACTERISTICS
TRANSLATIONAL AND TRANSCRIPTIONAL IMPLICATIONS OF SCMS
LANDSCAPE OF SOMATIC AND GERMLINE SCMS GENOME-WIDE
HYPERMUTATOR PHENOTYPE OF RAD54L RGSCM126
4.4 Methods 129
4.5Discussion
4.6 References
Chapter 5: Conclusion and Future Directions
CANCER SPECIFIC EXON DEFINITION 148
5.1 References
Appendix: Additional Projects
5.1 Complex Insertions and Deletions in Cancer Genomes
5.2 Analysis of Somatic Complex Insertions and Deletions in Pediatric Hematopoietic Malignancies
5.3 Children's Discovery Institute: Utilizing CharGer to determine novel variants in undiagnosed pediatric cases
5.4 Discovery of Novel Fusions in Cancer Genomes
5.5 Evaluating Novel Germline Variants in Cancer Genomes
5.6 Effect of Blood Somatic Mutations in PPM1D on TP53 and Cell Cycle Arrest 210
5.7 Discovery of Truncation Mutations Leading to Protein Alterations
5.8 Viral integration
5.9 DROSHA Mutation Analysis 234

List of Figures

Figure 2.1:	29
Figure 2.2	
Figure 2.3:	
Figure 2.4	
Figure 2.5:	41
Figure 2.6	43
Figure 3.1:	66
Figure 3.2	69
Figure 3.3:	73
Figure 3.4	75
Figure S1	77
Figure 3.5:	78
Figure 3.6:	82
Figure 3.7	83
Figure S2:	89
Figure S3:	94
Figure S4:	95
Figure S5:	96
Figure S6:	97
Figure 4.1:	113
Figure 4.2:	117
Figure 4.3:	122
Figure 4.4:	125
Figure 4.5:	127
Figure S7:	140
Figure S8:	141
Figure S9:	142
Figure A.1:	210
Figure A.2:	212
Figure A.3:	221
Figure A.4:	224
Figure A.5:	225
Figure A.6:	227
Figure A.7:	228
Figure A.8:	230
Figure A.9:	234
Figure A.10:	235
Figure A.11:	235

igure A.12:	236
-------------	-----

List of Tables

152
164
175
212
225
236

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Reyka Jayasinghe Washington University December 2018 Dedicated to my mother and grandfather, for always pulling me back to science even

when you weren't around.

Abstract of the Dissertation

The Splice is Not Right: Splice-site-creating Mutations in Cancer Genomes

by

Reyka Jayasinghe

Doctor of Philosophy in Biology and Biomedical Sciences Molecular Genetics and Genomics Washington University in St. Louis, 2018 Professor Li Ding, Chair

Accurate interpretation of cancer mutations in individual tumors is a prerequisite for precision medicine. Large-scale sequencing studies, such as The Cancer Genome Atlas (TCGA) project, have worked to address the functional consequences of genomic mutations, with the larger goal of determining the underlying mechanisms of cancer initiation and progression. Many studies have focused on characterizing non-synonymous somatic mutations that alter amino acid sequence, as well as splice disrupting mutations at splice donors and acceptors. Current annotation methods typically classify mutations as disruptors of splicing if they fall on the consensus intronic dinucleotide splice donor, GT, the splice acceptor, AG. Splice site mutations as a group have been presumed to be invariably deleterious because of their disruption of the conserved sequences that are used to identify exon-intron boundaries. While this classification method has been useful, increasing evidence suggests that splice site mutations can lead to transcriptional

changes beyond disruption and that many exonic mutations that act primarily through alternative splicing are still being overlooked in cancer genomics. My thesis work focuses on developing tools to systematically classify and functionally validate splice site and splice creating mutations using RNA-Seq data, to more accurately understand the functional consequences of mutations on alternative splicing by integrating DNA and RNA-Sequencing data.

First we developed SpliceInator, a semi-automated tool to systematically detect splicing phenotypes using mutation and gene expression data. We interrogated 1,146 conserved splice site mutations across 19 cancer types revealing a wide range of complex splicing phenotypes and emphasize the importance of analyzing patient specific RNA-Sequencing. We further explored beyond the splice site by interogating all mutations in a splicing context using MiSplice for the first large-scale discovery of splice-creating mutations (SCMs) across 8,656 TCGA tumors. We reported 1,964 originally misannotated mutations having clear evidence of creating novel splice junctions. Mutations in a subset of genes including PARP1, BRCA1, and BAP1, were experimentally validated for splice-creating function using a mini-gene splicing assay. Notably, we found neoantigens induced by SCMs are likely several folds more immunogenic compared to missense mutations, exemplified by the recurrent GATA3 SCM. Our work highlights importance of integrating DNA and RNA data for understanding functional and clinical implications of mutations in human diseases. Finally, to further capture the full landscape of SCMs, we explored both somatic and germline mutations for splice-site-creating function using MiSplice. Altogether, we have gathered a set of 2,888 SCMs enabling us to effectively compare the landscape of rare and germline SCMs. This compendium of SCMs has also started to elucidate novel genomic properties of mutations located at the donor and acceptor splice site and SCM containing exons including an overall decrease in the size of the novel exon post mutation, mimicking a natural evolutionary selective pressure but exploited in the cancer genome to maintain proper alternative splicing. To date, this is the first analysis comparing rare germline SCMs and somatic SCMs revealing their comparable dysregulation to the splicing code in cancer. Together my thesis work revealed that splice-site-creating mutants play a much larger role than previously appreciated in contributing to cancer and further expands our understanding of the genetic basis by which mutations can alter the mRNA landscape by dysregulating alternative splicing. More broadly, my work calls for a deeper analysis of seemingly "silent" mutations in any disease as such mutations may alter gene function via alternative splicing and integrating RNA and DNA-Seq can allow for accurate evaluation of mutations in a splicing context.

Chapter 1: Introduction

1.1 Cancer Cell Fitness

Introduction: Cancer, the second leading cause of death in the US, will cause an estimated 609,640 Americans deaths in 2018. The scientific community has come to the conclusion that cancer is a "disease of the genome" but deciphering the origins of cancer is still an evolving field of research. Over time, cells acquire random mutations, many of which have no effect, some with deleterious effects, and a few that confer a selective advantage that allows the cell to grow faster than neighboring cells (Stratton et al., 2009). After a cell acquires a set of genomic alterations that allow it to grow freely, resist cell death, escape normal signaling, and acquire the properties of immortality, metastasis and angiogenesis, the cancer cell can form a malignant mass that is harmful to the host (Hanahan and Weinberg, 2011).

Type of Mutations: Mutations can be germline or somatic in nature. Germline mutations are inherited from parents or acquired *de novo*. Somatic mutations are acquired throughout an organism's lifetime in individual cells due to genetic and environmental factors, such as chemicals and radiation. Most of the damage in the DNA is repaired, but sometimes the alterations are fixed. Acquired mutations include point mutations (single nucleotide variants / SNVs), insertions, deletions, larger copy number aberrations (CNA),

or large scale structural variation (SV). Mutations in genic and regulatory regions can affect gene function in several ways by causing loss or gain of function, altering transcript splicing, and increasing or decreasing gene expression level. Epigenetic changes on histones and DNA, (such as methylation etc.) have been shown to play a central role in turning genes on and off, and are known mechanisms of dysregulation in cancer (Chen et al., 2014; Maunakea et al., 2013; Rajagopal et al., 2014). In combination, such variations can disrupt normal gene function and alter cellular response to regulation giving the cell a selective advantage to proliferate autonomously.

Passenger and Driver Mutations: Genomic changes contributing to cancer can be placed into two categories: driver and passenger events. Driver mutations confer a growth advantage to the cell (under selective pressure) and are positively selected for during the development of the tumor. Novel bioinformatic analyses of large scale sequencing data have helped to infer pathogenic/driver and passenger mutations. Lu *et al.* analyzed allelic imbalance to identify potential initiating germline mutations in cancer (Lu et al.). In a study to identify rare germline truncation variants in 12 cancer types from The Cancer Genome Atlas (TCGA), variants with a higher variant allele fraction (VAF) in the tumor compared to the normal tissue were hypothesized to have undergone positive selection in the tumor. More recently, our lab expanded on this analysis by expanding to a larger cohort of 10,389 samples and expanded the landscape of known pathogenic drivers in a subset of cancer types (Huang et al.). Passenger mutations are those that are acquired in somatic cells but do not increase or inhibit cellular growth potential. These mutations can occur prior to, during, and after tumor initiating mutations but most are present prior to the accumulation of driver mutations (Stratton et al.). Since they are present prior to the driver event, when the driver mutation does confer a selective advantage to a cancer clone, you may observe passenger mutations "hitch-hicking" with the driving clone. Alternatively, initiation mutations are events that confer an advantage to the disease, but require cooperating mutations for developing disease phenotypes. Xie et al. identified initiation mutations in DNMT3A, ASXL1, TET2 and other genes associated with myelodysplasic syndrome (MDS), myeloproliferative neoplasm (MPN), chronic lymphocytic leukemia (CLL) and acute myeloid leukemia (AML) (Xie et al., 2014). These mutations were identified in patients without overt hematological malignancies suggesting that they may initiate clonal expansion but alone are insufficient to result in the development of cancer. Since this publication, there have been several novel studies that have expanded on this analysis by evaluating the presence of initiation mutations in the general population, but at extremely low variant allele fraction. This finding has appropriately initiated an entire cancer genomics field and spurred the development of novel sequencing technologies to accurately classify low level variants relative to background noise (Wong et al.; Young et al.).

Tumor Suppressors and Oncogenes: Cancer drivers can be categorized as tumor suppressors or oncogenes. In 1979, Oppermann and colleagues linked the neoplastic transformation of cells by avian sarcoma virus to a phosphoprotein with kinase activity encoded by *src*(Oppermann et al.). This report identified the first oncogene, a gene where an alteration to one allele could contribute to the malignant tumor phenotype and cellular transformation. In 1984, *RB1* was identified as the first tumor suppressor gene (TSG), a

gene with suppressor or regulatory function(Murphree and Benedict). TSGs commonly require mutation or inhibition of both alleles in order for cellular transformation to occur. According to Knudson's "two hit hypothesis", patients carrying a germline mutation in a tumor suppressor are more likely to develop cancer since a second random somatic mutation affecting the wildtype allele in any cell is much more likely to occur than two random mutations in a non carrier(Knudson). The "two hit hypothesis" is a tale as old as time and is still commonly referred to in the literature as a method to identify putative cancer predisposition genes in large genomic datasets(Park et al.).

Large Scale Cancer Projects: With the first cancer genome sequenced in 2008, the amount of sequencing data in cancer genomics has increased dramatically, supplying the scientific community with a gold mine of data to process, analyze, and refine. Several large scale projects have taken the initiative to integrate genomic data and genome wide annotation tracks allowing multiple teams to work together and pour their efforts into determining key genes and pathways that contribute to cancer. The Cancer Genome Atlas (TCGA) is a project that collected expression, methylation, RNA and DNA sequence data for matched tumor and normal samples across 33 distinct cancer types. The International Cancer Genomics Consortium (ICGC) seeks to provide genomic, transcriptomic, and epigenomic data across 50 different tumor types. The Pediatric Cancer Genome Project (PCGP) is a collaboration between St. Jude Children's Research Hospital and Washington University School of Medicine in St. Louis whose goal is to determine genetic changes that give rise to childhood cancers. The Clinical Proteomic Tumor Analysis Consortium (CPTAC) has expanded off the analyses and projects of the

cancer genome atlas to expand the integration of omics data beyond the transcriptome and to the proteome and phosphoproteome. These projects are just a few of the many large-scale projects that are aimed at identifying the genetic origins that contribute to the development and progression of various forms of cancer.

Pan-Cancer Studies: With the accumulation of large-scale genomic data across tissues types due to the aforementioned large scale-cancer projects, some labs have dedicated their entire research focus on pan-cancer analyses. The goal of these pan-cancer analyses is to glean important and relevant signatures and patterns among many patients to determine and define cancer by molecular subtype in order to expand and properly design personal therapies across cancer types (Cancer Genome Atlas Research et al.; Cooper et al.; Hoadley et al.; Huether et al.; Kandoth et al.; Sveen et al.; Vogelstein et al.). Currently, small sample sizes for individual cancer types limits our statistical power to identify more infrequently mutated driver genes. To address this problem, TCGA lead the effort in emphasizing the importance (statistically and biologically) to compare samples across multiple cancer types to identify infrequently mutated sites that are potentially driving cancer development. Initial findings from pan-cancer studies included: the identification of 127 significantly mutated genes across 12 cancer types (Kandoth et al.), which has more recently been expanded to a list of 299 cancer related genes (Bailey et al.). The identification of rare germline pathogenic variants including an analysis across 12 cancer types and subsequently 33 cancer types(Huang et al.; Lu et al.). While the TCGA phase has wrapped up, the upcoming phases, which will expand into the clinic to guide personalized medicine, is going to be a very exciting new direction for the field and

will bring novel genomic applications directly to the clinic. While these aforementioned pan-cancer analyses have elucidated many differentially mutated regions in the genome, it is important to remember all sequencing projects are limited by the mutational load of the tumor of interest.

1.2 Alternative Splicing and Disease

The potential for phenotypic variability within an individual species is encoded and dictated by the underlying genomic. Understanding how differences in gene architecture can alter a gene's function is essential to understanding many biological questions including disease diagnosis, progression, and treatment. To understand genomic variability, we will briefly discuss the genomic differences between species, evaluate how the genome has evolved in multicellular organisms, and discuss how mutations can lead to disease.

The Splicing Code: One of the characteristic differences between unicellular and multicellular eukaryotic organisms is the difference in genomes. The central dogma of biology postulates that DNA is transcribed to RNA which is then translated to proteins that perform most functions within the cell. This central dogma lead scientists to realize that certain segments of DNA were not transcribed into RNA or translated in the final protein structure. Instead, transcribed regions of the DNA (exons) were often interrupted by longer segments of DNA that were systematically removed from the RNA (introns) to

generate the final sequence that is shuttled for protein translation. In 1978, Walter Gilbert hypothesized different combinations of exons could generate multiple mRNA isoforms from the same pre-mRNA molecule (Berget et al.; Chow et al.; Gilbert; Modrek and Lee, 2002). This hypothesis has come to be known as alternative splicing, the process of joining exons or coding sequences, and splicing out introns.

There is still active debate regarding where in our evolutionary history introns arose. The introns early (IE) hypothesis suggests that introns are ancient structures but were lost in intron-poor species such as prokaryotes (Irimia and Roy; Roy and Gilbert). In opposition, the introns late (IL) hypothesis postulates certain species like eurkaryotes gained introns due to insertion events after the divergence of prokaryotes and eukaryotes. Complicating this question more, intron-rich and intron-poor species are interspersed throughout the eukaryotic branch, suggesting massive intron gain or loss could lead to the diversity of genomic sequences present in eukaryotes.

The average human transcript contains approximately 8 introns. Current studies suggest that between 70-95% of human genes harbor multiple mRNA transcripts(Johnson et al., 2003; Matlin et al., 2005; Modrek and Lee, 2002). Alternative splicing expands the complexity and information content of the eukaryotic genome and allows for tissue, developmental, or temporally expressed isoforms which can perform alternate functions.

The splicing code is made up of cis-acting elements that help the splicing complex distinguish between non-coding and coding regions and facilitates the joining of exons

and exclusion of introns (Wang and Cooper, 2007). The consensus intronic dinucleotide GT splice donor and AG splice acceptor flank spliced exons at the 3' and 5' ends respectively, and can be found in 99% of all introns. Intronic and exonic splicing enhancer (ISE and ESE) and suppressor (ESS and ISS) elements influence the splicing complex to target true splice sites and pseudo splice sites (Supek et al., 2014b) (Ast; Bonomi et al.; Faustino and Cooper; Keren et al.; Matlin et al.). The spliceosome is responsible for joining exons and splicing out introns to form the mature mRNA molecule. The spliceosome is made of five small nuclear ribonucleoproteins (snRNPs), U1, U2, U4, U5 and U6, along with a number of other proteins. The U1 snRNP interacts with the 5' splice site (5' ss), splicing factor 1 (SF1) identifies the branch site, and the U2 auxiliary factory (U2AF) binds the polypyrimidine tract and the 3' splice site (3' ss). Additional proteins that facilitate the splicing process include SR proteins (SR) which bind to ESEs and interact with U2AF. Mutations in splices sites and enhancer or suppressor sequences can ultimately contribute to a pathogenic phenotype by altering the binding affinity for spliceosomal or associated proteins.

The splicing process is dynamic and occurs co-transcriptionally. For this reason, genomic context alone cannot begin to describe the efficiency of the splicing code. But we still need to start somewhere.

RNA polymerase II transcribes DNA to create the nascent mRNA molecule. As the molecule is transcribed, the spliceosomal machinery is recruited to facilitate the removal of introns and the joining together of exons. RNA polymerase transcriptional dynamics

can be influenced by underlying genetic architecture. For example nucleosomes are commonly found positioned on exons which have a high GC content. The median length of exons are coincidently very close in size to the length of genomic DNA that wraps around histones, suggesting an added selective pressure on maintaining an exon size of rougly 140 bp. Nucleosomes are made up of 8 core histone proteins which contain histone tails that have post-translational modifications (PTMs). PTMs can influence both RNA polymerase transcriptional dynamics and RNA processing by attracting necessary splicing factors to facilitate proper splicing. For example the H2A.Z histone 2A variant is known to promote splicing of non-consensus introns in S. cerevisiae (Herzel et al.; Neves et al.).

Splicing alterations and cancer development: Splice alterations have been shown to affect the landscape of mRNA isoforms present in both tumor and normal tissues (Wang and Cooper). Mutations in cis can directly affect the use of a splice site or promote the use of an alternative splice site facilitating exon/intron inclusion. The production of inappropriate transcripts in a particular tissue type can result in disease due to its alternate function. Mutations in canonical splice sites have been linked to several diseases, for instance: A familial study found in hSNF5, when the consensus GT dinucleotide is mutated to an AT, a deletion of exon 7 occurs. This germline mutation was observed in both affected and unaffected family members, but infant brain tumors in affected members showed a loss of the wild-type allele in the tumor(Taylor et al.; Venables). In colorectal and liver metastases, a 3' splice site mutation from AG to AT leads to a loss of the adjacent exon. Furthermore a loss of the wild type allele in the tumor suggests that this isoform is under positive selection in the tumor (Kurahashi et al.). In another familial study, a single intronic base change formed a cryptic splice site resulting in the addition of 11 nucleotides to the tumor suppressor BRCA1 gene creating a truncated protein(Findlay et al.; Hoffman et al.). Less studied are mutations in less conserved intronic regions. In *ATM*, a mutation at the 6th position of an intronic region is linked to breast cancer but only causes a proportion of transcripts to form a truncated protein(Broeks et al.).

All of the above studies are one-off cases where a resulting phenotypic effect was observed in an individual which led to the discovery of the mutation inducing the aberrant splicing pattern. Bioinformatic heavy labs are starting to group similar splicing alterations to evaluate genomic signatures that can predict the resulting phenotypic changes. Below we will describe some of the recent novel findings in the field that have started to broadly characterize the normal and mutation induced splicing landscape across populations.

The global effects of mutations on splicing variation is now being widely studied in largescale RNA-Seq consortia. Rivas et al. predicted the effects of protein-truncating variants on the transcriptome by utilizing data from the Genotype-Tissue Expression (GTEx) and Geuvadis projects (Rivas et al., 2015). Both projects allowed the team to quantitatively evaluate allele specific expression of 13,182 mutations in healthy individuals, primarily focusing on nonsense and splicing mutations. Their findings identified variants within the splicing region with significant splice-disruption events and confirmed their findings in a separate dataset of common variants in Swedish individuals. Using the same dataset, a recent publication evaluated the level of "deleterious-ness" of mutations near the splice-site to determine a more refined definition for the splice-region (Zhang et al., 2018). In addition to the canonical GT and AG sites, the study found one base in the exon and four bases into the intron after the canonical GT (donor) and the first exonic base adjacent to the AG (acceptor) site are mutation intolerant. Additionally, co-occuring mutations were less likely to occur suggesting one mutation in one of the aforementioned sites was enough to perturb the splicing pattern. Additional studies identified synonymous driver mutations in key oncogenes, and found that nearly half of all synonymous driver mutations affected splicing, specifically last base exonic mutations (Supek et al., 2014a). Together, these findings suggest that synonymous driver mutations likely also contribute to cancer through altering alternative splicing.

Shiraishi et al. developed SAVNet to evaluate somatic variants that induced splice altering variants across 31 cancer types in the TCGA cohort (Shiraishi et al., 2018). The authors found that the smoking signature (C>A substitutions) contributed largely to the variants characterizing splicing differences followed by APOBEC.

Cummings et al. used paired transcriptome analyses for patients with undiagnosed muscle disorders and identified recurrent splice-site-creating variants in collagen VI-related dystrophy and predicted functional variants in mutation-rich genes such as *TTN* (Cummings et al., 2017).

Several studies have tried to integrate computational predictions and massively parallel splicing assays to gain new insight into the splicing code. Soemedi et al. evaluated 4,964 single nucleotide variants via a massively parallel splicing assay (MaPSy) (Soemedi et al., 2017). Approximately 10% of variants were confirmed *in vitro* and *in vivo* to disrupt splicing but their assay is limited in only being able to evaluate mutations in exons that were less than 100 nucleotides in length.

Mutations in Trans Elements & Cancer: Mutations in trans-acting elements or splicing factors will affect a larger subset of genes by disrupting the biological machinery that helps to recognize exons and introns. The most well studied splicing factors include serine/arginine-rich (SR) proteins and heterogenous nuclear ribonuclear proteins (hnRNPs)(Kaida et al.; Matlin et al.). SR proteins bind to ESE's and ISE's and promote splicing and inclusion of exons by recruiting spliceosomal proteins. The family of hnRNP proteins function as splicing silencers by binding to ESS's and ISS's. SR proteins in particular are shown to be differentially regulated in cancer and play a role in neoplasia(Fregoso et al.; Kaida et al.). These splicing elements have been shown to function in a position dependent fashion(Lim et al.). SR proteins are usually bound in exons while hnRNPs are found in introns, but modifications that lead to altering the splicing elements binding sites has been linked to their opposite function. Cis mutations that affect binding of hnRNPs and SR proteins can drastically change whether exon usage is promoted or suppressed. An initial study showed that many disease alleles should be classified as splicing mutations even though they were mis-classified as missense mutations because of the motifs that were changed near the splice site(Lim et al.). Using the pan-cancer dataset by TCGA, more motif alterations can be elucidated and linked to splicing alterations.

Mutations in enhancer and silencer sequences can also affect isoform expression because of the proteins that associate with them. Variants in *BRCA1* (Walker et al.), *SMN1/2, PDHA* and *GH* are known to cause exon skipping by disrupting the ESE sequence(Woolfe et al.). This same study showed that splice altering variants (SAVs) are enriched in regions near the splice junctions. Exon skipping SAVs are characterized by a loss of ESE and a gain of ESS while variants with increased exon inclusion are characterized by ESS loss. Alternative isoforms are linked to a number of different diseases which have been shown to contribute to drug resistance, tumor angiogenesis, metastasis, and misregulation of apoptosis(Woolfe et al.). Tools are still needed to identify the functional significance of somatic and germline splice altering mutations within the coding and noncoding regions of a gene.

While the field has made great strides in classifying variants that disrupt alternative splicing, interpreting all variants in a splicing specific context has still been largely ignored in large scale sequencing projects. Variant classification and interpretation is fundamental to understanding the biological consequences of mutations on a gene of interest. If variants are not properly annotated, the biological interpretation is incorrect. Currently, mutations are classified based on their changes to the resulting protein. The degenerate nature of the genetic code whereby 61 codons represent 20 amino acids allows nucleotide changes to affect the sequence of mRNA while leaving the amino acid

sequence of the resulting protein unchanged. "Silent" mutations that do not change the amino acid are still to this day commonly overlooked in cancer genomics. A study analyzing mutations in the CFTR gene showed that synonymous mutations could alter splicing by leading to the use of a cryptic splice site or by altering the 3' splice site of exon 9 which results in a truncated protein product(Niksic et al.; Wang and Cooper; Wilschanski et al.) Another recent study showed that synonymous mutations were enriched in oncogenes and some tumor suppressor genes in a cancer type specific manner (Supek et al., 2014a). Furthermore, this study found many recurrent "synonymous" TP53 mutations are inactivating events that alter canonical splicing of the mRNA. These studies suggest that some seemingly "silent" mutations can affect gene function by modifying splicing, transcription factor binding, or other properties that contribute to mRNA translation(Sauna and Kimchi-Sarfaty, 2011). Using known exonic and intronic mutations that activate cryptic splice sites, Lee et al. developed a machine learning classifier to predict whether a mutation would disrupt alternative splicing (Lee et al., 2017). Interestingly, in analyzing variants in CFTR, 70% of the 47 variants tested, were predicted to be missense variants suggesting many splicing alterations are mis-annotated in wellknown disease-related genes.

The previously mentioned studies all emphasize the importance of evaluating synonymous and nonsynonymous mutations across cancer types in order to uncover potential pseudo or non-canonical splice sites that are currently being overlooked in cancer genomics.

My thesis work focused first on generating bioinformatic tools that systematically classify splice site and splice-site-creating mutations by integrating DNA and RNA sequencing and experimentally validating the predicted consequences of the variants using a minigene splicing assay. In Chapter 2, I helped develop SpliceInator a semi-automated tool to systematically detect splicing phenotypes using mutation and gene expression data. SpliceInator, combines two lines of evidence to assess mutant specific aberrant splicing events and their implications, one based on interpretation of RNA-Seg fragment mapping using TopHat and another based on standard statistical hypothesis testing of RSEM expression values. We interrogated 1,146 conserved splice site mutations across 19 cancer types revealing a wide range of complex splicing phenotypes. 521 variants in our dataset conferred a measurable splicing alteration, with 69.29% associated with only one splicing defect, while the remaining were a combination of two to four different splicing events. Another 624 splice site mutations did not confer any measurable splicing defects but 75.6% were classified as having a low variant allele fraction, low exon expressivity or both while 24.8% were undetermined. Furthermore, synonymous and non-synonymous variants genome-wide were evaluated in a splicing context and we discovered 243 mutations that create and strengthen nearby alternative splice sites, respectively, further justifying the demand for a tool that can systematically evaluate all mutations in a splicing context. To meet this demand, in Chapter 3 we developed MiSplice for the first largescale discovery of splice-creating mutations (SCMs) across 8,656 TCGA tumors. We report 1,964 originally mis-annotated mutations having clear evidence of creating novel splice junctions. TP53 and GATA3 have 26 and 18 SCMs, respectively and ATRX has 5 from low-grade gliomas. Mutations in 11 genes including PARP1, BRCA1, and BAP1,

were experimentally validated for splice-creating function. Notably, we found neoantigens induced by SCMs are likely several folds more immunogenic compared to missense mutations, exemplified by the recurrent GATA3 SCM. Further, high expression of PD-1 and PD-L1 was observed in tumors with SCMs, suggesting candidates for immune blockade therapy. Our work highlights importance of integrating DNA and RNA data for understanding functional and clinical implications of mutations in human diseases.

Finally, to further capture the full landscape of SCMs, in chapter 4 we evaluated both somatic and germline mutations for splice-site-creating function using MiSplice. Altogether, we have gathered a set of 2,888 SCMs enabling us to effectively compare the landscape of rare and germline SCMs. Interestingly, we found mutations overlapping the splice donor site were sufficient to disrupt the canonical splice site but this phenomenon doesn't hold true for acceptor splice site mutations in our dataset. Alternatively acceptor SCMs needed to not only strengthen the novel splice site to facilitate the novel site usage, but also disrupt the canonical splice site. This compendium of SCMs has also started to elucidate novel genomic properties of SCM containing exons including an overall decrease in the size of the novel exon post mutation, mimicking a natural evolutionary selective pressure but exploited in the cancer genome to maintain proper alternative splicing. To date, this is the first analysis comparing rare germline SCMs and somatic SCMs revealing their comparable dysregulation to the splicing code in cancer. Evaluating mutation induced events separately from patient specific de novo events can provide a focused analysis on the genomic features selecting for SCM+ exons relative to leaky splicing or mutation independent cryptic splice site activation. As tissue type specific datasets continue to increase, developing novel tissue specific signatures will help inform the tissue type specific relevance of mutation induced SCMs. Together my thesis work revealed that splice-site-creating mutants play a much larger role than previously appreciated in contributing to cancer and further expands our understanding of the genetic basis by which mutations can alter the mRNA landscape by dysregulating alternative splicing. More broadly, my work calls for a deeper analysis of seemingly "silent" mutations in any disease as such mutations may alter gene function via alternative splicing and integrating RNA and DNA-Seq can allow for accurate evaluation of mutations in a splicing context.

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22

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24

Chapter 2: Complex patterns of splice

site mutations

Contribution: I scripted and developed the full code for the TopHat Module and partial code to the Expression Level Module and Rule Based Classifier of SpliceInator. I manually reviewed all variants validated by our computational tool, performed downstream analyses and created all Figures other than 2.1A.

2.1 Introduction

Large-scale sequence-based studies are increasingly examining the functional consequences of human genomic variants (Dees et al.; Ding et al.; Hoadley et al.; Huether et al.; Kandoth et al.; Wang and Cooper; Xie et al.), with many focusing on characterizing somatic coding mutations and their amino acid sequence alterations in order to investigate the underlying mechanisms of cancer. Splice mutations are of particular interest because of their dramatic modifications of mature RNA products, including possible retention of large intronic segments and loss of whole exons, coupled with significant downstream consequences for associated surviving proteins. With respect to cancer, splicing alterations can affect both tumor and normal tissues (Wang and Cooper), but determining their functional consequences within and outside of the canonical splice site is still understudied in cancer genomics (Barbaux et al.; Broeks et al.; Holmila et al.; Kurahashi et al.; Steffensen et al.; Venables; Woolfe et al.). By one estimate, around 22% of disease causing alleles are mis-classified as missense mutations because they alter

motifs near the canonical site utilized by the splicing machinery. Especially relevant to cancer are synonymous mutations that can be enriched in oncogenes and some tumor suppressor genes in specific cancer types(Supek et al.). Several recurrent mutations in cancer-associated genes, such as *TP53*, were found to be inactivating events, altering canonical mRNA splicing. Silent and missense mutations can also affect gene function by modifying splicing, transcription factor binding, or other aspects of mRNA translation (Sauna and Kimchi-Sarfaty). These studies show that mutations in *cis* can directly affect the use of a splice site, or promote the use of an alternative splice site, thereby facilitating exon skipping or intron inclusion. There is a growing appreciation of the need for reliable splicing analyses to identify alternative biochemical pathways that are causative in contributing to the disease state(Dorman et al.; Lim and Fairbrother; Mort et al.; Rivas et al.; Steffensen et al.). It is likely that many, perhaps most of these pseudo or non-canonical splice sites are currently being overlooked in cancer genomics.

Given the biomedical importance of splice-affecting mutations, a bevy of analysis tools has been developed over the last few decades. Early entrants, including Gene Splicer(Pertea et al.), MaxEntScan(Yeo and Burge), Splice Site Finder(Shapiro and Senapathy), NNSplice(Reese et al.), and Human Splicing Finder(Desmet et al.) sought mainly to identify splice site mutations, but methods have increasingly added new analysis capabilities. For example, tools like Skippy(Woolfe et al.), SpliceTrap(Wu et al.), DEXSeq(Anders et al.) use RNA-Seq data to discern differential exon usage and exonskipping and SpliceSeq(Ryan et al.) added significant visualization aids. Algorithms are becoming progressively more ambitious, for example SpliceR(Vitting-Seerup et al.) and SplicingTypesAnno(Sun et al.) attempt to characterize a broader set of events encompassing intron retention and various exon extensions and contractions. Calculations often depend on tally-based heuristics and ad-hoc rules. The overall problem of comprehensively characterizing splice mutations and their downstream implications remains unsolved.

Research increasingly suggests that splice architectures are complex and their implications for cancer are significant(Ast; Bonomi et al.; Faustino and Cooper; Keren et al.; Matlin et al.; Modrek and Lee; Taylor et al.; Venables; Wang and Cooper) . A number of specific mechanisms have already been observed. For example, variants in *BRCA1*, *SMN1/2*, *PDHA* and *GH* cause exon skipping by disrupting an exonic splicing enhancer sequence(Walker et al.; Woolfe et al.). Mutations near pseudo splice sites can also result in activation, for instance the c.190 mutation in BRCA1(Yang et al.), which appears to be under positive selection in the tumor. This mutation is a clear example of a predicted "missense" event that actually functions as a modifier of splicing efficiency in vivo and points to modified alternative splicing as a more important effect in cancer than has been previously appreciated. Importantly, we found that standard annotation tools did not properly identify this variant as splice altering.

Given the importance of splice effects in cancer and the lack of a purely algorithmic solution for finding and characterizing them, we applied a semi-automated approach to this problem based on large batches of case-control RNA-Seq data that are now

28

available. We apply this method here to examine 1,146 candidate mutations over 19 cancer types and describe several novel findings.



Figure 2.1: SpliceInator Workflow. The SpliceInator workflow can be grouped into two main modules and post processing analyses. In Module 1, case and control RSEM expression values are compared using standard statistical hypothesis testing to classify intron retention and exon skipping events. In Module 2, BAMs are processed with TopHat using RNA-Seq fragment mapping to identify mutant specific alternative splice junctions to classify intron retention, exon skipping, exon shrinkage and exon extension events. In the post processing steps, a rule based classifier is used to derive support from module 1 and module 2 to classify variants with associated splicing defects. Second, quantile analysis is performed to group genes into expression classifications in a tissue specific manner. After processing, users can compare variants to predicted splicing defects to infer the functional consequences of each variant in a splicing context. The right-hand column highlights several thresholds and internal tests that are utilized in SpliceInator that can alter which steps are run for each module. For example, if read count thresholds are not satisfied in the second module, then only module 1 and the post processing steps will be tested and reported.

2.2 Results

SPLICEINATOR – AUTOMATIC PRIORITIZATION OF SPLICE MUTATION EVENTS

Given the difficulty of rigorously characterizing splice mutation events, we instituted a procedure to automatically prioritize candidates from large lists of inputs (Figure 2.1). Called SpliceInator, it identifies the most promising members, assigning each a preliminary assessment of one or more phenotypes. SpliceInator employs 3 forms of analysis. In the first module, expression levels (units of RPKM) are permutation-tested against corresponding controls, for example intron x in the case versus intron x in the control group. If 3' bias of the case gene is sufficiently low (Spearman rank test), this between-samples assessment is augmented by a within-sample permutation test, for instance the subject intron against the other introns in the gene. In this latter scenario, the probability results would then be combined using Fisher's chi-square transform method. The second module uses TopHat(Trapnell et al.) assembly to identify boundaries and establish read counts. Given some minimal read thresholds, a Fisher exact test is applied to assess whether patterns of junction coverage are altered between case and controls due to splice site mutation. A third component of SpliceInator classifies whole gene expression values based on Tukey's quartile analysis, including pronouncement of no expression for values more than 1.5 interquartile ranges below the first quartile. Finally, using a rule based classification we then prioritized variants from module 1 and module 2 using the following rules: 1) Events are classified as intron retention if found to be statistically significant by module 1; 2) Exon extension and exon shrinkage events are reported if found to be statistically significant by module 2; 3) Exon skipping events are classified based on a combination approach. If the exon skipping event is found to be statistically significant in module 1 and module 2, the module 1 pvalue is reported and a designation of M1;M2 is reported. If module 2 reports a statistically significant pvalue, an M2 designation is reported. The codebase is available from GitHub (github.com/ding-lab/SpliceInator).

DIVERSE SPLICING PHENOTYPES ASSOCIATED WITH SPLICE SITE MUTATIONS

We collected high quality mutation calls with a UCSC conservation score greater than 99% from 19 cancer types derived from The Cancer Genome Atlas (TCGA). A total of 1,146 splice site mutations currently defined as substitutions, deletions, or insertions overlapped the 2 bp canonical intronic splice donor or acceptor of 624 cancer associated genes. We predicted mutations located at highly conserved sites in cancer genes would be associated with measurable splicing defects. The TopHat module identified alternative junctions near splice site mutations and each prediction was manually reviewed to produce a high confidence list of mutations and associated splicing defects. Our analysis revealed TopHat accurately classifies exon extension, exon shrinkage, and some exon skipping events, but is unable to identify intron retention, another widespread defect observed in many samples by manual review. The statistical based module using RSEM data properly identified intron retention and exon skipping events that were missed in the TopHat alignment analysis. 45.46% of variants in our dataset conferred measurable splicing alterations. The overall landscape of splicing phenotypes for 521 variants included: exon skipping (230), 5' exon shrinkage (137), intron retention (219), 3' exon shrinkage (49), multi-exon skipping (32), 3' exon extension (27), and 5' exon extension (24) (Figure 2.1a). 69.29% (361) of splice site mutations had only one associated splicing defect, while the remaining were a combination of splicing events including 2 events (24.76%, 129), 3 events (4.99%, 26) and 4 events (0.96%, 5).



Figure 2.2. Splicing defects associated with splice site mutations. (A) An UpSetR plot highlighting the distribution of splicing phenotypes associated with splice site mutations across the dataset. The bar plot on the left hand side reflects the total amount of sites that fall into each splicing phenotype group. The bar plot at the top depicts the total mutations that are classified as having a corresponding splicing defect depicted in the bubble's below. Colors correspond to a separate splicing phenotype. Black lines between bubbles indicate complex splicing events. (B) For each splicing phenotype, we evaluated how the splicing defect would alter the reading frame of the protein.

SPLICE SITE MUTATIONS LINKED TO COMPLEX SPLICING EVENTS

Current tools and analyses focus on the primary splicing defect, but 30.6% of mutations with a measurable splicing phenotype in our dataset have two to four different splicing alterations linked to the same splice site mutation. The most common complex event involves intron retention and single exon skipping in 29 cases. A 2011 study(Ma et al.) in c.elegans provides evidence that intron retention and exon skipping is regulated by the identity of the 3' splice site due this regions interaction with U2AF splicing factors.

Five mutations in KDM6A, KMT2B, CBFB and TP53 presented with 3' exon shrinkage and single exon skipping. Two of the mutations sharing this out of frame complex splicing event are annotated to exon 26 of the H3K27 demethylase, KDM6A, and have lower expression relative to their BLCA cohort (Figure 2.3a). A closer examination of the RNA-Seq revealed equal support of the 3' exon shrinkage and single exon skipping events, and a lack of reads supporting the canonical junction in both cases. Using MaxEntScan we scored the canonical and alternative splice sites using a sliding window approach to evaluate the strongest alternative splice site score with the introduced mutation. Before the mutations the donor site score is 8.07 but with the introduced mutations the score changes to -0.33 and -6.43 for the point mutation and larger deletion, respectively. Although the alternative splice site utilized in the 3' exon shrinkage event in both cases has a lower splice score of -0.88, Human Splicing Finder scored the alternative site as a potential donor site (70.87 out of 100) taking into consideration additional genomic features such as branch point potential and splice factor protein binding sites(Desmet et al.). Altogether, these results suggest the donor splice site mutation weakens the canonical splice site decreasing the overall expression of KDM6A while also utilizing a nearby alternative donor site leading to a 3' exon shrinkage and exon skipping event. The landscape paper on LUAD(Cancer Genome Atlas Research), and two additional studies(Banka et al.; Cheon et al.) of Korean patients with Kabuki syndrome report the

presence of frequent mutations in KDM6A, including mutations overlapping the same splice site, but did not report on the functional consequences of the splice site mutation.

Since the recurrent *KDM6A* splice site mutations had the same splicing phenotype, we next evaluated the landscape of recurrent splicing alterations across the cohort. Our dataset contained 300 recurrent splice site mutations present in two or more samples overlapping the same splice site. Several highly recurrent mutations in the same cancer type showed similar splicing patterns including *GATA3* in BRCA and *MET* in LUAD. The *GATA3* simple indel was identified in 12 breast cancer (BRCA) samples and expressed the same 5' exon shrinkage event across all 13 samples (Arnold et al.; Dorman et al.). Four different mutations overlapping the same splice site in *MET* expressed a similar exon skipping profile that is well supported in the literature(Drilon).

A. Genes with Recurrent Splice Site Mutations







Figure 2.3. Complex Splicing Events. (A) Heatmap showing the distribution of splicing phenotypes associated with select recurrent splice site mutations. Cancer type classification is highlighted above, followed by gene expression classification using quantile based analysis. For select genes, recurrently muatated splice sites in the same gene are grouped by black bands within their gene group. Splicing defects for each site are classified as in frame and out of frame with an empty and filled in square, respectively. (B) Lolliplot with 80 recurrent mutations in TP53 spanning 9 splice sites. Mutations are depicted by a circle, with each circle colored according to the identified splicing defect. If more than one defect is identified to be linked to that mutation, the circle is divided into two or three quadrants, with each quadrant colored according to the splicing defect.

Alternatively, many recurrent sites did not show such consistency across samples and tissue types. *TP53* is highly mutated across cancer types and harbors 80 recurrent splice site mutations across nine splice sites. Evaluating the distribution of splicing phenotypes for each splice site shows a very different effect. Mutations at p.261_splice are derived from BRCA, HNSC, KIRC and UCS and while three cancer types share the same intron retention event, the UCS sample has no measurable splicing alterations. Six of the seven mutations at p.224_splice are derived from four cancer types and all lack a measurable splicing defect, but a single mutation in SKCM expresses 3' exon extension and intron retention which could be explained by the high tumor VAF (94.44%) for this particular mutation. This result highlights the importance of having individualized RNA-Sequencing to complement mutational analysis to evaluate patient specific splicing alterations.

LACK OF MEASURABLE SPLICING DEFECT FOR MANY SPLICE SITE MUTATIONS

We next wanted to explore whether there was a relationship between the presence of an aberrant transcript and the newly formed premature termination codon (PTC). The Nonsense Mediated Decay (NMD) pathway predominantly degrades PTC's and nonsense mutations that differ from the canonical stop codon to reduce expression of potentially damaging transcripts. The general rule of thumb is that PTC's located at least 50-55 bp upstream of the last exon-exon junction drive strong NMD, whereas those out side of this criteria are predicted to escape the degradation process. To evaluate if the presence of a measurable splice defective product is due to the lack of degradation by NMD, we translated the aberrantly spliced product and evaluated the position of the PTC relative to the exon-intron boundary predictions. Our analysis found that X mutations create a PTC that is predicted to escape NMD while X mutations should hypothetically be degraded.



A. Reasons for lack of measurable splicing phenotypes

B. Exon expressivity comparison



Although we identified splicing alterations for 521 variants, 624 splice site mutations were not associated with any measurable splicing defects suggesting one or more of the following: (1) a lack of expression of the transcript; (2) low expression of the variant allele in the tumor; (3) the creation of a highly unstable transcript that is degraded; (4) copy number aberrations; (5) presence of a subclone; or (6) a false positive. To evaluate lack of expression of the case sample relative to the controls, we performed Tukey's quartile analysis, including pronouncement of no expression for values more than 1.5 interguartile ranges below the first quartile (Methods). While 339 sites had high or average expression relative to their cohort, 14 and 272 genes were considered as not expressed or lowly expressed, respectively, suggesting the mutation disrupted overall expression of the transcript. To assess exon expression we evaluated reads supporting the adjacent annotated exon against control samples and normalized by the length of the transcript and total mapped reads (Methods). For 116 of the 624 splice site mutations, the transcript had no expression in the tissue type of interest illuminating the lack of a measurable splicing defect in the presence of a splice site mutation. Furthermore, the group lacking splicing defects had an overall lower exon expressivity and lower variant allele fraction distribution when compared to the group of variants with measurable splicing defects (Figure 2.4). Altogether, our results suggest 75.6% of highly conserved splice site mutations lacking a measurable splicing defect can be classified as having low VAF (<30%), low exon expressivity or both while 24.8% are still undetermined (Figure 2.4).

Gene	Sample	Cancer	VAF %	CN	Purity	Splicing Defects	
TP53	TCGA-N5-A4RS	UCS	88.8	2.02	0.6832		Intron Retention
PTEN	TCGA-EB-A44P	SKCM	75	1.26	0.7913	•	•
TP53	TCGA-22-5482	LUSC	74.07	1.94	0.7887		
KEAP1	TCGA-55-6972	LUAD	72.73	1.38	0.9048		3' Exon Shrinkage
CREBBP	TCGA-FU-A3TX	CESC	63.64	1.97	0.794		
AXIN1	TCGA-CN-A497	HNSC	63.16	2.65	0.7019		5' Exon Shrinkage
ACVR2A	TCGA-53-7624	LUAD	62.3	2.45	0.8436		
KEAP1	TCGA-44-7667	LUAD	60.87	1.78	0.6368		
TSC1	TCGA-GV-A3QF	BLCA	60.18	1.94	0.7913		3' Exon Extension
MGA	TCGA-BT-A20N	BLCA	59.38	1.78	0.7417		
CHD4	TCGA-EA-A4BA	CESC	57.89	1.95	0.9509		5' Evon Extension
WAS	TCGA-AY-6196	COADREAD	55.26	1.77	0.5077		
CDKN2A	TCGA-EE-A20B	SKCM	53.57	1.83	0.691		
PTEN	TCGA-B5-A11R	UCEC	53.02	2.47	0.9212		Multi Exon Skipping
BCORL1	TCGA-CN-5356	HNSC	52.83	2.01	0.6448		
BRIP1	TCGA-EE-A2MD	SKCM	52.43	2.62	0.8366		
MAP4K3	TCGA-GV-A3QK	BLCA	51.52	2.51	0.809		Single Exon Skipping
KMT2C	TCGA-CZ-4857	KIRC	50.7	3.15	0.61		
CBLC	TCGA-EB-A5SF	SKCM	50	1.62	0.8304	۲	
							Out of Frame
							🔵 In Frame

IMPACT OF SPLICE SITE MUTATION ON GENE EXPRESSION

Figure 2.5. Splice altering variants with increased expression. 19 predicted splice eQTLs identified across cohort. Indicated in figure are gene name, sample, cancer, variant allele fraction (VAF), copy number (CN), purity estimate, and the associated splicing defects indicated by color. Color legend on right hand side corresponds to associated splicing defect, and a filled in circle indicates predicted out of frame events.

We next compared cases against cancer specific control cohorts to evaluate the effect of splice site mutations on gene expression. Overall, around 59% of sites had comparable

(high or average) expression within their control cohort, while 41% had lower overall expression (low or not expressed). Interestingly of the 59%, 19% had higher gene expression, selecting for an altered transcript that escapes RNA degradation pathways. In the high expression group, we sought to identify splicing expression quantitative trait loci (eQTLs) where the splice site mutation is associated with higher expression in the gene of interest. We hypothesized splice eQTL's would have a measurable splicing defect, elevated DNA VAF (>50%) and increased overall gene expression of case samples relative to the associated control cohort. Out of 526 mutations with measurable splicing alterations, 19 are predicted to be splice eQTLs in our cohort and they reside in the following cancer associated genes: TP53 (2), PTEN (2), KEAP1 (2), CREBBP, AXIN1, ACVR2A, TSC1, MGA, CHD4, WAS, CDKN2A, BCORL1, BRIP1, MAP4K3, KMT2C, and CBLC (Table 1). Interestingly, 10 of these 19 mutations are complex events, having 2 or more splicing phenotypes associated with the mutation, and 13 are classified as having a retained intron.



Figure 2.6. KEAP1 highly expressed splice eQTLs. (A) Lolliplot depicting location of highly expressed splice site mutations in KEAP1 (p.570). Each domain is highlighted using a different color and each circle on the lolliplot visualizes the location of the mutation on the exonic sequence and highlights the complex splicing patterns identified at each site. The resulting consequences of each splicing defect on the kelch domains is further annotated below the lolliplot revealing crucial structural elements that are absent under each condition. (B) Protein structure of the kelch domains in the KEAP1 structure. Six kelch domains form a propellar structure within KEAP1 (gray) that is responsible for interacting with Nrf2 (yellow). The circles in the upper left hand corner of each structure highlight the splicing defects that will disrupt the associated colored kelch domain in the 3D depiction. The amino acid position where the splice site mutations fall are highlighted in blue and annotated along with the Nrf2 interacting domain (yellow).

Two highly expressed splice eQTLs were identified in the E3 ubiquitin ligase Kelch-like erythroid cell derived protein with CNC homology (ECH) associated protein 1 otherwise known as KEAP1. KEAP1 sequesters and ubiquitinates Nrf2, a nuclear factor erythroid

2-related factor 2 (NFE2L2). In cancer, many mutants disrupt the binding between KEAP1 and Nrf2. Disrupted binding increases the presence of free Nrf2, thereby activating downstream genes that are conducive for cell growth and responding to oxidative stresses. Three LUAD samples in our dataset harbor splice site mutations in KEAP1, two of which were classified as high expressing splice eQTLs. We hypothesized the two mutations in KEAP1 are disrupting the binding between KEAP1 and Nrf2. One mutation was identified at the acceptor site of exon 6 and another at the donor site of exon 5, with a VAF of 61% and 72%, respectively. Copy number data also suggests the mutant with 72% VAF has undergone a copy number loss, providing further evidence as to the selection of the mutant case in the tumor. Furthermore, while one mutation occurs within the last 50 bp of the last exon-exon junction and the other is present in the last exon of KEAP1, this could explain why the splicing defects are tolerated and escape degradation by nonsense mediated decay.

When comparing the KEAP1 mutants against the control group, KEAP1 mutants had higher overall gene expression (wilcox test=0.04) and lower overall NFE2L2 expression (wilcox test = 0.01). Interestingly, one KEAP1 mutant sample with RPPA data had higher Nrf2 protein expression relative to the controls and lower KEAP1 protein expression, providing an answer as to the mechanism by which the splice eQTL could be disrupting this biological pathway. The discordant relationship between protein and gene expression in this case could be explained by a negative feedback loop leading to a decrease in gene expression under conditions of high protein expression. Structurally, the mutations occur within the last of the six kelch repeats that fold into a beta propeller tertiary structure, with each blade compromising four anti-parallel beta sheets (Figure 2.6b). In Figure 2.6b we highlight the kelch domains that are predicted to be disrupted due to associated splicing defects. For example, if exon 5 of KEAP1 was skipped, the fifth kelch domain would be completely absent thus disrupting the overall propeller structure and the binding pocket for Nrf2. These results lead us to believe the KEAP1 high expressing splice eQTLs act in a dominant negative manner by disrupting the binding of KEAP1 to Nrf2 thereby increasing the amount of free Nrf2 in the cell.

NON-CANONICAL SPLICE SITE MUTATION DISCOVERY

Finally, we wanted to perform a genome-wide investigation on the creation and strengthening of de novo splice sites by exonic variants. We used two independent methods to interrogate mutations outside of the splice site in a splicing context. Using the entropy based method, for each variant the sequences of each 9-mer (donor) and 23-mer (acceptor) covering the variant position was extracted from the genome for both the mutant and reference sequence. Their splice score as potential donor or acceptor sites were estimated using MaxEntScan. The largest scores of the 9-mer or 23-mer windows were retained for mutant and reference and their difference (mutant - reference) was calculated. Finally, the scores with largest difference between potential donor site and potential acceptor site was retained as the final score. A novel splice site is predicted to be created if the reference score <3 and the mutant score is >8. We further filtered this list by evaluating nearby junctions in the bam files (reads>10) for predicted novel splice sites. For the second method, we used the TopHat module of SpliceInator to collect all alternative junctions within 14 base pairs of the annotated mutations. We manually reviewed all the predictions to evaluate the efficacy of our method.

For the entropy based method, a total of 243 sites were predicted near a novel junction but after manual review the method correctly called 9 de novo donor sites, 6 strengthened alternative acceptor sites, and 7 strengthened alternative donor sites. Of the 22 sites, 15 were specific to the mutated sample and not present in any control samples. For the TopHat method, a total of 192 alternative acceptor sites and 204 alternative donor sites were predicted to fall within 14 bp of a non-splice site mutation. After manual review, 1 *de novo* acceptor site, 35 *de novo* donor sites, 153 strengthened acceptor splice sites, and 145 strengthened donor splice sites were properly detected. Furthermore, 49 of the 154 acceptor sites had no supporting reads supporting the novel junction in the control and 84 of the 180 donor sites were specific to the mutated sample. The MaxEntScan scoring filtered out many sites that were found using our TopHat method, suggesting using the splice score alone isn't sufficient to properly evaluate mutations in a splicing context. Both methods picked up strengthened acceptor sites in the following genes *CTSH*, *LAMC1* and *NUP98;* and 4 created donor sites in *WDR33*, *NOP14*, *RSRC2* and *PARP1*.

The Poly ADP-ribose polymerase 1, PARP1, is an enzyme involved in recruiting proteins involved in DNA repair pathways. Both methods identified a silent mutation in PARP1 in a LUSC patient that acts as a splice altering variant by creating a de novo donor site in exon 21. RNA-Sequencing data supports the use of the *de novo* donor site leading to an 11 amino acid deletion, which falls within the PARP1 catalytic regulatory domain. 180 reads in the mutated sample supported the *de novo* site while 170 controls contained no supporting reads giving us strong evidence that this "silent" mutation is more appropriately a splice altering variant. PARP1 inhibitors are commonly used in cancer treatment to disrupt DNA repair in tumor cells thereby leading to the accumulation of DNA breaks and ultimately cell death(Malyuchenko et al.).

2.3 Methods

SpliceInator Methods:

The analysis procedures use RPKM values, which were derived for the full-length exons and introns of genes using TCGA RNA-Seq bam files. Briefly, bed files were generated for exon and intron coordinates based on the Ensembl 37.75 database. The SamTools package counted raw reads for each exon and intron. Finally, RPKM was calculated as $109 \cdot R/(N \cdot L)$, where R is the number of raw reads mapped to each exon or intron, N is the total number of mapped reads in the project, and L is the length of the exon or intron. Some analyses also consider ± 50 bps and ± 2 bps around junctions, for which we use RPM (# of mapped reads per million reads) to quantify the expression because the lengths are short and fixed.

The main algorithm, which we call SpliceInator, combines two lines of evidence to assess aberrant splicing events and their implications, one based on standard statistical hypothesis testing of RSEM expression values and the other based on interpretation of RNA-Seq fragment mapping using TopHat. We have found empirically that each of these components excels at a relatively mutually exclusive subset of the various kinds of splicing anomalies that have been observed: intron retention and exon skipping for the statistical method and exon extension and shrinkage for TopHat-based analysis. The latter also can detect some exon skipping events, for which perform ad hoc interpretation in light of the statistical evidence. Consequently, we have not found it necessary to develop any sort of overarching method for combining results of the two calculations.

For the genes of interest, the statistical method takes as input both the gene-wide RSEM and the individual intron and exon RSEM values for each gene over a set of samples. Genes reporting RSEMs below a threshold are considered to have expression too low to properly assess and are skipped. Otherwise, each sample for each gene is processed

sequentially for the different kinds of possible splicing anomalies. Each of these calculations proceeds in essentially the same way, the intron retention calculation being representative, as follows. Preliminary exclusion criteria are evaluated first: the intron's RSEM must be above a threshold and must likewise be greater than the average RSEM over the corresponding introns in the control group, of which there must be at least 50 instances. Assuming these criteria are met, the algorithms then evaluates whether the subject intron's RSEM is significantly higher than the average of those in the control group via simple permutation testing, which returns a "case-control" P-value. The method attempts to maximize power when possible. Specifically, if 3' bias within the gene of the current sample is sufficiently low, a within-sample test is also performed. Potential 3' bias is checked via Spearman's rank test between the introns' RSEM values and their location midpoints, both ordered 5' to 3'. Lack of strong bias is inferred if Spearman's coefficient \leq 0.3, in which case the algorithm runs a second permutation test to check whether the subject intron RSEM is significantly greater than the other intron RSEMs within that gene. This calculation returns the "within-sample" P-value. These tests are essentially independent, by which we finally combine the case-control and within-sample P-values using Fisher's Method. Analysis of exon-skipping is similar. The algorithm is relatively robust against the heuristic parameters quoted above.

For TopHat-based analysis, TopHat Junction output was used for each splice site mutation to identify junction ids where mutations are between the start and stop of the junction feature. Exon intron boundaries were calculated using the overhang value for each junction feature. We enforced a threshold of at least 10 reads aligned to a particular exon-intron boundary. Depths of coverage for each junction for cases and controls were collected using the TopHat junction output. To obtain total unique reads within the junction neighborhood, we performed read counts on the in house bam files and used this value as our total reads for both cases and controls. Reads supporting reference junction versus alternative junction were compared using Fisher's 2x2 exact test between cases and controls to discern altered junction coverage due to presence of the splice site mutation.

Methods Novel splice site mutation discovery:

We propose to use the TopHat Module of SpliceInator to identify all alternative junction predictions across the entire cancer genome atlas RNA-Seq sample set (>7,000 samples) and pediatric cancer dataset (>500 samples). To hone in on mutations that are directly disrupting pseudo splice sites, we will first identify missense and silent variants that are within a threshold distance of an alternative junction prediction. After identifying the alternative junction, we searched for nearby donor and acceptor sites that may have been strengthened or weakened by the introduced mutation. By comparing cases to controls within the same cancer type, we were able to determine if the novel junction is used within the cancer type at a low level or only present in the case with the mutation. Finally, each *de novo* donor and acceptor site is scored using a scoring algorithm. By using entropy and information analysis of the consensus base sequences, we can evaluate the strength of a particular site based on its nucleotide frequencies(Burset et al.). Using a weighted scoring method for each possible acceptor/donor sequence, we multiplied the information of each base by the information weight of the site. The acceptor and donor information and weight is derived from over 25,000 different splice donor/acceptor sites to come up with the information weights and content at each position. To be classified as creating or strengthening a novel acceptor or donor site the mutation fell within 8 bp of the novel predicted donor site, 14 bp of the novel predicted acceptor site, had supporting reads from both the positive and negative strand, and create or strengthen a nearby site that utilizes the canonical GT or AG splice site.

Methods to evaluate exon expression:

Github/Exon-Expressivity

We developed a script which take as inputs a Mutation Annotation File (MAF) and an RNA bamlist to compare total reads supporting an exon against the total mapped reads (normalized by length) to evaluate exon expressivity across cancer types and samples in our dataset.

Exon_Expressivity.pl : Inputs 1) RNA-Seq bam locations 2) Mutation Annotation File (MAF). This script will identify all exons nearest to the splice site mutation of interest designated in the MAF file and collect exon read count data across samples in the RNA-Seq bam list and total mapped reads from the flagstat or alignment stats files in the designated bam directories. The exon expressivity coefficient (Ei) is calculated by dividing the total reads aligned to exon (E) over the total mapped reads of the imported bam (B), normalized by the length of the exon and multiplied by a scaling factor (see equation below). The Ei coefficients are then averaged across all samples for each cancer type to determine the overall exon expressivity of the exon of interest. If multiple transcripts share the same exon, multiple Ei values will be reported for each individual exon.

• Exon Expressivity (Ei) =
$$\frac{\left(\frac{Total Reads Aligned to Exon (E)}{Total Mapped Reads of Imported Bam (B)}\right)$$
Length of Exon (bp) $* 10^9$

To determine a cut off for the exon expressivity (Ei) we evaluated Ei scores across all 1,159 mutations in our dataset and associated control samples. We determined Ei of case samples with detectable splicing alterations to define samples with known "expressed junctions".

<u>Alignment guided junction analysis to identify mutations linked to splicing defects</u>: We propose to identify junctions between two exon-exon boundaries genome wide. A junction is defined by RNA-Seq reads that are split between two exon-intron boundaries, joining two exons. TopHat⁴¹ reports the number of reads supporting the canonical junction shared between two exons, or any deviations including exon skipping, intron retention, 5' exon shrinkage, 3' exon shrinkage, multi exon skipping, 3' exon extension, and 5' exon extension. The number of reads supporting each junction for cases and controls were collected for each sample. Controls were defined for each gene based on tissue type and only include samples lacking mutations in the gene of interest. Reads supporting the reference junction and alternative junctions were compared using Fisher's Exact Test between cases and controls to determine which junctions showed statistically significant altered junction usage. This calculation indicates whether the altered junction is potentially due to the mutation in the case, or if a background level of alternative splicing is occurring at this junction due to a tissue type specific event.

<u>Part 1. SpliceInator</u>: Splice site mutations that lead to aberrant signals have cancer implications. We developed a tool combining TopHat RNA-seq analysis with case-control and within-sample statistical testing of expression data to identify and classify such events.

<u>RNA-Seq Guided Junction Analysis using SpliceInator to identify mutations contributing</u> <u>to splicing defects:</u> SpliceInator is a statistical tool that we are developing to classify variants into the following categories: 1) expression loss, 2) intron retention, 3) exon skipping, 4) exon extension, 5) exon shrinkage, and 6) unknown, using patient specific RNA-seq data. It performs permutation based testing on normalized expression data derived from exons, introns, and splice junctions between mutant and control samples. Additionally, when there is minimal 3' bias, within-sample testing is also performed to add statistical power.

First, reads per kilobase per million (RPKM) are derived for each gene using patient specific RNA-Seq. Next, mutations are tested for intron retention. Within this test, we first check for 3' bias within the given sample according to Spearman's Rank Correlation test of each exon across the gene in a 5' to 3' manner. Absent significant bias, RPKM of each intron is tested against all others within the gene to check for elevated expression. Next, the full intron expression of the mutant sample is tested against the same intron in the control cases. For exon skipping, the same type of calculations are performed, except full exon expression is used. After an initial analysis of a highly recurrent GATA3 acceptor variant, we noted that exon shrinkage and extension events could not be properly identified when comparing the full exon and intron expression between cases and controls. For the exon shrinkage and extension tests, we decided to additionally extract RPKM values associated with +/- 2bp and +/- 50 bp of all defined exon boundaries. This allowed us to test smaller events that could be occurring near a splice site, such as the 7 bp shrinkage identified in the GATA3 variants, or larger events with the +/- 50 bp data.

For exon shrinkage and exon extension, the full exon expression, 2 bp expression and 50 bp expression are all compared between cases and controls. This analysis will allow us to connect our expression analysis with junction-based prediction and identify cases of intron retention, which are outside of TopHat's predictive capability.

Intron retention events are common splicing aberrations: SpliceInator identified 104 intron retention events highlighted in Figure 2.1. An additional 44 exon skipping events were identified by SpliceInator, further supporting the findings of TopHat predicted exon skipping events. Our findings emphasize the importance of using both tools in identifying splicing defects associated with mutations. [add more examples here about interesting intron retention events].

3.4 Discussion

Splice alterations have been shown to affect the landscape of mRNA isoforms present in both tumor and normal tissues. Unlike many missense mutations known to disrupt the protein structure or associated binding sites, an initial review of splice site mutations quickly revealed position and tissue type alone couldn't explain observed splicing patterns. Since looking solely at genomic context could not directly inform whether a splice site mutation would confer a splicing defect we developed SpliceInator, a semiautomated tool to interrogate mutations in a splicing context.

Evaluating splicing defects associated with highly conserved splice site mutations revealed diverse splicing patterns. While some well known splicing patterns were observed, more than 30% of variants with a known splicing defect had a more complex

phenotype consisting of two or more splicing defects. Although some recurrent splice site mutations at the same position conferred similar splicing defects, many exhibited different patterns, suggesting tumor heterogeneity makes predicting the effects of mutations on the transcriptome very difficult without patient specific RNA-Sequencing data. In particular, TP53 harbored many recurrent splice site mutations across cancer types and didn't show consistent splicing patterns while recurrent mutations in KDM6A did.

Even more interesting was the lack of a measurable splicing defect for 54% of highly conserved splice site variants. While we were able to classify approximately 75% of variants from this group into samples having low variant expression and low exon expressivity, approximately 25% were still unclassified. The unclassified variants could be creating a highly unstable transcript that is quickly degraded by the cell, but follow up functional analysis is needed to confirm this prediction revealing the limitation of RNA-Seq data. We were also able to demonstrate mis-annotated variants by evaluating all mutations in a splicing context using SpliceInator. By more accurately classifying variants in genes such as PARP1 and PTEN we can better understand the biological implications of variants disrupting the splicing process and determine patient specific treatments.

Altogether our findings suggest you can't predict what to expect due to the complexity of the splicing process and tumor heterogeneity. While our tool makes a significant contribution by providing an automated method to determine functionally relevant mutations using matched RNA-Seq data there is still much to learn about how cancer can abrogate splicing mechanisms for tumor growth and proliferation. For example, while we focused on defining specific defects such as exon skipping, exon shrinkage, exon extension and intron retention, there were several events that were not easily classified into one or more of the aforementioned categories. Further developing our tool to accommodate more complex splicing defects will be a necessity to better gauge the true landscape of splicing phenotypes in disease.

3.5 References

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61

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Chapter 3: Discovery of splice-site-

creating mutations

Contribution: I developed 1/3 the code for MiSplice with the other parts developed by Qingsong Gao and Song Cao. I manually reviewed all variants validated by our computational tool, performed downstream analyses, created all Figures and performed entire mini-gene splicing assay for 11 variants tested.

3.1 Introduction

Large-scale sequencing studies, such as The Cancer Genome Atlas (TCGA) project, have worked to address the functional consequences of genomic mutations in tumors (Dees et al., 2012; Kandoth et al., 2013; Lawrence et al., 2013; Niu et al., 2016), with the larger goal of determining the underlying mechanisms of cancer initiation and progression. Many studies have focused on characterizing non-synonymous somatic mutations that alter amino acid sequence, as well as splice disrupting mutations at splice donors and acceptors (Jung et al. 2015). Current annotation methods typically classify mutations as disruptors of splicing if they fall on either the consensus intronic dinucleotide splice donor, GT, or the splice acceptor, AG. As a group, splice site mutations have been presumed to be invariably deleterious because of their disruption of the conserved sequences that are used to identify exon-intron boundaries.

While this classification method has been useful, increasing evidence suggests that splice site mutations can lead to transcriptional changes beyond disruption of the canonical junction (Lim and Fairbrother, 2012; Mort et al., 2014; Rivas et al., 2015; Sauna and Kimchi-Sarfaty, 2011; Steffensen et al., 2014). One such example is the c.190 mutation in BRCA1. Conventional annotation had predicted a missense mutation, p.C64G, but our analysis of RNA-seq data in ovarian tumors harboring p.C64G and a published mouse model (Yang et al., 2003) suggested the germline c.190 mutation leads to the creation of an alternative splice junction, resulting in a truncated null protein. There have been several case studies reporting observations of missense and silent mutations activating cryptic splice sites in MLH1 (Nyström-Lahti et al., 1999), LMNA (Woolfe et al., 2010), RB1 (Zhang et al., 2008), RNASEH2A (Rice et al., 2013), MECP2 (Taimoor I Sheikh, 2013), BAP1 (Wadt et al., 2012), KIT (Chen et al., 2005), as well as other studies relating missense and silent mutations with splicing changes (Jung et al., 2015; Kahles et al., 2016; Soemedi et al., 2017; Supek et al., 2014). Despite the broad clinical ramifications of mutation-induced altered splicing, systematic evaluation of their occurrence and resulting effects in cancer has yet to be undertaken, nor have there been significant bioinformatics platforms for doing so.

We applied a newly developed bioinformatic tool called MiSplice, which integrates DNA and RNA-Seq data across thousands of samples to discover mutations that induce splice site creation. In our large-scale analysis across 8,656 tumor samples, we report 1,964 such somatic mutations that had originally been mis-annotated. Splice site-creating mutations (SCMs) are enriched in the new introns, with the highest rate at the -3 nucleotide position of acceptors with two-thirds of such events at aGag and agGag repeats by creating an alternative junction 2 nucleotides away. Partial and full splice creation capabilities across these 1,964 sites are evaluated by measuring the fraction of reads supporting the alternative junction, which we term the Junction Allele Fraction (JAF) and which is found to be negatively correlated with distance to the new splice site. In total, 1,607 genes harbor SCMs, with 248 of them having more than one mutation, including *TP53, GATA3, ATRX,* and *NF1*. Recurrent SCMs were found in *TP53, GATA3, DDX5, KDM6A, PTEN, SETD2, SMAD4, BCOR, SPOP,* and *BAP1,* suggesting association with cancer development. Broadly speaking, integrated DNA and RNA data can furnish a sound basis for discovering SCMs and for accurately understanding functional consequences of mutations in cancer and other human diseases.

3.2 Results

SPLICE-SITE-CREATING MUTATION DISCOVERY

We collected high quality mutation calls from 8,656 tumors across 33 cancer types derived from The Cancer Genome Atlas having available TCGA RNA-Seq data (Methods). For every mutation, we defined a set of control samples in the same cancer cohort that lacked the same mutation in the gene of interest. We sought to assess the landscape of SCMs across cancer genomes by evaluating all mutations already having conventional annotations and their potential splice site-creating effects (Fig. 3.1A). To achieve this goal, we conducted analysis using a bioinformatic tool, MiSplice (Mutation Induced Splicing), which systematically evaluates mutations in a splicing context using RNA-Sequencing data (Fig. 3.1B).



Figure 3.1. Splice site-creating mutation discovery. (A) Examples of splice site-creating mutations for different conventionally annotated mutation types. Splice-in is defined as mutations contained within the newly created exons and splice-out is when the mutation is present in the newly created intron. (B) The MiSplice work flow consists of three steps: alternative junction discovery, filtering, and manual review. First, the user inputs the locations of RNA-Seq BAM files along with a mutation file. MiSplice searches the BAM file to identify any alternative splice junctions near the mutation of interest, while filtering out known splice junctions and calculates the number of alternative junction supporting reads for case and control samples. For the filtering step, the following sites are removed: mutations in HLA genes, low fraction of reads supporting the alternative splice junctions. (C) Breakdown of 2,056 manually validated splice site-creating mutations by conventional annotation.

MiSplice manages large analyses using parallel computation to search for alternative splice junctions within windows of ± 20 bp from the mutation of interest. For example, of

the 1,416,566 candidate mutations examined here, 4,448 had 5 or more unique RNA-Sequencing reads supporting the predicted alternative junction in proximity to the mutation. MiSplice then conducts a series of further evaluations, including Ensemblbased filtering of canonical junctions, establishing observational significance by case comparison to a matched set of controls, and assessing score and depth of each cryptic site using MaxEntScan (Yeo and Burge, 2004) and SamTools (Li et al., 2009). From the resulting subset, MiSplice filters out HLA genes and sites whose junctions have insufficient difference of expression, as judged from the case-control assessment. Here, we evaluated promising alternative junctions with at least 5% of paired end RNA-Sequencing reads at the genomic location supporting the alternative junction of interest. MiSplice processing revealed 2,056 mutations (Table S1) potentially creating an alternative splice site. Manual review indicated a 2.09% false positive rate, suggesting high specificity of the MiSplice algorithm for discovering these types of mutation-induced splicing events. Of these putative splice events, 1.90% and 0.47% are considered complex and are in highly homologous gene regions, respectively, so they were excluded from further analyses (Methods).

Of the final 1,964 splice site-creating mutations (SCMs) passing manual review (Table S1), 52% (1,016) are in annotated splice sites, suggesting disruption of the canonical splice site and selection of a the alternative splice site nearby (Fig. 3.1C). Importantly, 26% (513) and 11% (208) of the SCMs had previously been mis-annotated as missense and silent mutations, respectively. In addition, we found 58 insertions or deletions, 46 nonsense, and 123 non-coding region mutations that likewise create cryptic splicing sites.

MOLECULAR AND BIOLOGICAL PATTERNS OF SPLICE SITE-CREATING MUTATIONS We next characterized the sequence context for the 1,790 SCMs corresponding to single nucleotide mutations. The sequences of each 9-mer (donor) and 23-mer (acceptor) covering the mutation position were extracted for both the mutant and reference sequences. Their splice scores as potential donor or acceptor sites were then estimated using MaxEntScan (Table S1).



Figure 3.2. Sequence contexts and characteristics of splice site-creating mutations. (A)

Frequency distribution of splice site-creating mutations relative to the newly created splice junction, with high frequency shown at the 3rd nucleotide position in the newly created intron. (B) Comparison of splicing scores for the newly created splice site, before (reference) and after the mutation (mutant). Showing larger effect of mutations at the 3rd nucleotide position in the intron (especially for the 3' splice sites). (C) Dominant nucleotide sequence context for splice site-creating mutations at -3 position of the 3' splice site. Mutation position (red dot) is present 3 base pairs away from the newly created exon. (D) Transition and transversion rate at the -3 position of the 3' splice site. Most mutations are G>C transversions, strengthening the consensus sequence of the splicing factor U2AF1. (E) Comparison of splicing scores between the nearest canonical splice junction with and without mutation compared to the newly created splice junction with and without the mutation. Most mutations strengthen the alternative splice junction relative to the canonical splice junction.

Mutations near the alternative splice junctions show higher mutation rates in the introns for both 5' (p<1e-5, binomial test) and 3' splice site (p<1e-6) (Fig. 3.2A). More interestingly, we found an enrichment of mutations at the third nucleotide position in the intron, but depletion at the first and second positions (especially for 3' splice site) (Fig. 3.2A). Comparison of splicing scores between splice site-creating mutants and reference forms shows that most mutants have stronger splice signals than the reference (Fig. 3.2B). Mutations that create a G or T to produce an alternative 5' splice site dramatically increase splice site strength. For 3' splice sites, mutations enriched on the third nucleotide of the newly created intron showed the largest increase of splicing score (Fig. 3.2B). Further examination of the sequence context around mutations at the third nucleotide of 3' splice sites shows that 53% have a mutation on aGag repeats and another 16% are mutated on agGag repeats, all creating alternative junctions 2 nucleotides away from the annotated ones (Fig. 3.2C). Mutations at the -3 position of the alternative acceptor site would potentially enhance U2AF1 recognition of the acceptor splice site. Previous studies have reported S34F U2AF1 mutants preferentially skip exons that contain a T nucleotide at the -3 position (Okeyo-Owuor et al., 2015). Of the 192 mutations that are located at the

-3 position from the alternative junction and contain an AG in the -2 and -1 positions, 56% undergo a G>C transversion (21% G>A, 18% G>T, 3% C>T, 2% A>C, 1% A>T) with C being the preferred base at the -3 position for *U2AF1* binding (Fig. 3.2D).

We also explored the relationship between the alternative and canonical splice junctions. As expected, mutations at splice sites dramatically reduced splice scores of the canonical splice junctions, while strengthening those at the alternative splice junctions in most cases. In contrast, a subset of missense and silent mutations did not drastically alter the canonical junction, but instead preferentially strengthened a nearby alternative splice site (Fig. 3.2E). When analyzing the raw splicing scores (canonical and alternative site before and after mutation), we found that 1,089 out of 1,790 (61%) events showed higher splice score for the alternative splice site than the canonical site, indicating inclination for the alternative sites. Further, while 485 (27%) events saw lower post-mutation alternative splice score, differences between alternative and canonical scores had decreased, suggesting that these mutations are still likely enhancing the preference for the alternative site. Only 214 (12%) events did not show evidence suggesting increased post-mutational preference for using the alternative site. These cases are a good illustration of the fact that many other genomic splicing features are also relevant, including exonic splicing enhancers (ESE), polypyrimidine tract, branch point, and RNA-binding proteins. They are also consistent with the general view that splice score is not definitive (Jian et al., 2014). We emphasize that all 1,790 alternative splice sites demonstrated usage based on patient RNA-Seq data and that 10 out of 11 (>90%) identified splice site-creating mutations were validated experimentally (see below).

EXPRESSIVITY AND PENETRANCE OF SPLICE-SITE-CREATING MUTATIONS

In the presence of the mutation, alternative splice junctions exhibited a wide range of expression. To quantify this effect, we measured alternative junction expression as the fraction of alternatively spliced junction spanning reads over the total number of reads at the genomic location, what we refer to as the Junction Allele Fraction (JAF). Fig. 3.3A shows the distribution of JAF's for all high confidence MiSplice predicted alternative junctions, separated by conventional mutation annotations (Fig. 3.3A). Currently, we use a JAF cut-off of 5% for reporting the final high-confidence sites. However, there are some potential alternative sites excluded by this cut-off. Our analysis revealed alternative junction expression varies widely. As expected, DNA variant allele fraction (VAF) and JAF have a generally positive correlation (Fig. 3.3B), with SCMs in KDM6A and FGFR2 having >75% DNA VAF and JAF. However, a SCM in ARID1A has a DNA VAF of 23% and JAF of 67%. Such large ranges have been noted for mutations outside of the splice site (Broeks et al., 2003; Clarke et al., 2000; Venables, 2004). Both the truncated and normal spliced products can be observed for many variants, either due to the wild type allele or leaky splicing, for example as observed in RNASEH2A, NFU1, SMN1, CFTR, and NF2 (Boerkoel et al., 1995; Caminsky et al., 2014; Ferrer-Cortes et al., 2016; Lohmann and Gallie, 2004; Mautner et al., 1996; Pagani et al., 2003; Rice et al., 2013; Svenson et al., 2001; Vezain et al., 2011).



Figure 3.3. Junction allele fraction of splice site-creating mutations. (A) The junction allele fraction (JAF) is defined as the number of reads supporting the alternative spliced junction relative to total junction spanning reads. Distribution of JAF values separated by conventional annotation type. (B) JAF vs. DNA Variant Allele Fraction (VAF) comparison by conventional annotation type. Most mutation types show a generally positive correlation between JAF and VAF values. (C) Splice site-creating mutations expressed in the newly created exon of the alternative splice junction. Comparison of

mutation position relative to the percent of reads supporting the alternative junction and mutation (Spliced-In JAF). The mean of each position is highlighted by the black point. For all positions, there is a strong correlation between the presence of the splice site-creating mutation and the alternative splice junction.

We next considered the expression of mutations that are spliced-in, i.e. mutations located within the exon of the alternatively spliced product. To this end, we determined the ratio of the number of alternative junction reads containing the mutation versus total number of reads supporting the alternative junction (Fig. 3.3C). Overall, most of the reads supporting the alternative junction also support the mutation, which suggests a strong association between the mutation and alternative splice junction. Regarding the 5' splice site, mutations within the first 6 bp of the new exon junction have a much higher fraction of alternative junction as the distance between them increases. For the 3' splice site, we observe a similar trend, although with a higher variability as a function of the distance from the alternative junction.

SPLICE SITE-CREATING MUTATIONS ACROSS GENES AND CANCER TYPES

A total of 1,607 unique genes harbored SCMs, with 85% (1,359) having one mutation and 15% (248) having two or more. *TP53* contained the greatest number (26), followed by *GATA3* (18). While most SCMs were found outside the current cancer gene compendium (Table S1), Fig. 3.4A shows that a remarkable number of cancer genes harbor splice altering variants, a phenomenon supported in the literature (Sebestyen et al., 2016). A

pan-cancer view reveals that *TP53* was the most mutated across cancer types, while 18 *GATA3* mutations and 6 *ATRX* mutations were specific to breast cancer (BRCA) and low grade glioma (LGG), respectively.



Figure 3.4. Splice site-creating mutations across genes and cancer types. (A) Distribution of splice site-creating mutations in each gene separated by total number of mutations in each gene. *TP53* has the largest number of splice site-creating mutations, followed by *GATA3* and *ATRX*. (B) Genes with the highest number of pancancer splice site-creating mutations. Circle size correlates with total number of mutations for each gene (labeled inside circle), and colored by cancer type. Splice site-creating mutations in *TP53* are present in many cancer types, while mutations in *ATRX* and *GATA3* are specific to LGG and BRCA, respectively. (C) Proteins Timeless (PAB domain) and PARP1 (Chain A) are colored green and pink, respectively. Originally annotated p.S939S mutation (red) and spliced-out sequence (blue) are highlighted on PARP1 (Chain A). (D) 3D protein structure of PARP1 in complex with an inhibitor (PDB ID: 5WRQ). Drug inhibitor and PARP1 (Chain A) are indicated with green and pink, respectively.

We observed 137 mutations nearby to one another (+/- 5 bp) which lead to the creation of the same recurrent splice site-creating events, not only in *TP53*, but also *GATA3*, *DDX5*, *KDM6A*, *SETD2*, *PTEN*, *SPOP*, and *BAP1*. While some mutations did not occur at the same position, 14 mutations creating the same alternative splice junction were found in the same exon, including 2 mutations in the third exon of *BAK1*. While most mutations in close proximity created the same alternative splice junction, two adjacent splice site-creating mutations in *CTNND1* and 2 nearby exonic mutations in *ACP2* and *GMPPB* created different alternative junctions.

SCMs can impact protein structure and have potential therapeutic implications. Poly ADPribose polymerase 1 (PARP1) is an enzyme involved in recruiting protein members of DNA repair pathways including Timeless PAB (PARP1 binding domain) (Fig. 3.4C) (Xie et al., 2015). Since PARP1 is essential to many cellular processes, including DNA repair, it is commonly targeted by antitumor agents (Malyuchenko et al., 2015). PARP1 inhibitors targeting the catalytic domain disrupt DNA repair mechanisms thereby increasing the effectiveness of chemotherapeutic agents (Fig. 3.4D). Identifying mutations that disrupt inhibitor binding are essential to properly evaluate treatment options. MiSplice identified a conventionally annotated silent PARP1 mutation (p.S939S) in a lung squamous cell carcinoma (LUSC) patient that acts as a splice site-creating variant by creating a de novo donor site (Fig. 5A). 82 reads supported the *de novo* donor site, which results in a 10 amino acid deletion (p.940-p.950) that falls within the catalytic domain (Fig. 3.4D). Out of 173 LUSC control samples, none contained reads supporting the alternative junction, providing strong evidence that the annotated "silent" mutation is actually a SCM. Previous reports of missense mutations at p.940 are predicted to reduce PARP1 enzymatic activity by disrupting the binding affinity of *PARP1* to its substrate NAD+ (Alshammari et al., 2014). The in-frame SCM is likely disturbing the local structure of *PARP1* and thereby disrupting the interactions between PARP1, its protein binding partners, and drugs binding within the pocket (Figs. 3.4C and 3.4D).



Figure S1. BAP1 gene and protein expression. Related to Figure 3.5. Violin plot of RSEM and RPPA data for control samples (grey) and novel splice creating mutant samples (red).

We identified two kidney renal clear cell carcinoma (KIRC) samples having the same conventionally annotated missense mutation (c.233A>G, p.N78S) in *BAP1*, a nuclear deubiquitinase, that created the same spliced-out alternative splicing product (Fig. 3.5B). Inactivation of *BAP1* is prevalent among renal cell carcinomas (Pena-Llopis et al., 2012) and an annotated missense mutation (p.L570V) has been reported to create a cryptic splice site in melanoma (Wadt et al., 2012). At the transcriptional level, the expressions of the case and control samples are relatively comparable, but at the translational level, one case with available protein data (RPPA) showed significantly lower expression (p=0.044, permutation test) relative to the controls (Table S2). This result suggests the conventionally annotated missense mutations in *BAP1* are likely creating an alternatively spliced transcript that is not readily expressed at the protein level.



Figure 3.5. Minigene functional assay of splice site-creating mutations. (A) Integrative genomics viewer (IGV) screenshot of the conventionally annotated synonymous mutation in *PARP1* in exon 21. RNA-Seq reads of the candidate splice site-creating mutation reveals the creation of an alternative splice site (red reads) created by the conventionally annotated synonymous mutation. (B) Candidate recurrent splice site-creating mutations in *BAP1*. Conventionally annotated as synonymous variants, *BAP1* mutated region shows alternatively spliced reads (red reads) in the IGV screenshot for each sample with the splice site-creating mutation. (C) IGV screenshot of a conventionally annotated synonymous mutation in *RAD51C* in exon 2. (D) Maximum entropy score of the splice site-creating variant before (purple) and after (red) the introduced mutation for each variant functionally validated in the mini-gene splicing assay. In silico predictions suggest all mutations strengthen the alternative splice site. (E) Candidate splice site-creating mutations validated by the mini-gene splicing assay. Exons of interest were cloned into the pCAS2.1 vector and mutant (red) and wildtype (purple) plasmids were transfected into 293T cells and total RNA was extracted to identify mutation induced alternatively spliced products.

We used a pCAS2.1 splicing reporter mini-gene functional assay, adapted from previous publications (Bonnet et al., 2008; Gaildrat et al., 2010; Malone et al., 2016; Tournier et al., 2008; Vreeswijk and van der Klift, 2012), to validate SCMs in eleven cancer genes, including two originally annotated silent mutations in *PARP1*, *RAD51C*, two splice site mutations in *TP53* and *BRCA1*, and several missense mutations in *ARID2*, *BAP1*, *BCOR*, *CDH1*, *KMT2A*, *PTEN*, and *TSC2*. Wild-type and mutant exons were cloned into a pCAS2.1 vector (Gaildrat et al., 2010) and transiently transfected into HEK293T cells. Total RNA was extracted to evaluate alternatively spliced products by RT-PCR. Examining the change in the MaxEntScan score for the 11 genes revealed mutations in *ARID2*, *BAP1*, *BCOR*, *CDH1*, *BCOR*, *CDH1*, *BCOR*, *CDH1*, *PARP1*, *RAD51C*, *PTEN*, and *TSC2* having dramatically

stronger splice scores in the presence of the mutation, while mutations in *BRCA1*, *KMT2A*, and *TP53* did not (Fig. 3.5D). Except for *PTEN*, variants with stronger splice scores showed higher levels of the alternatively spliced product in the mini-gene assay when compared to the wild-type. Variants with moderate changes in splice score still showed evidence of alternatively spliced transcripts, revealing the importance of utilizing functional assays to evaluate the effect of mutations in a splicing context in addition to *insilico* predictions. The mini-gene assay confirmed 91% (10/11 genes) splicing alterations in all tested genes and sequencing confirmed the alternatively spliced products (Fig. 3.5E, Methods), suggesting a strong concordance between MiSplice predicted splice site-creating mutations and the functional assay.

NEO-ANTIGENS INTRODUCED BY SPLICE SITE-CREATING MUTATIONS

We have further investigated neoantigens produced by splice site-creating mutations. By using the RefSeq transcript database, a total of 2,993 protein sequences were translated for transcripts containing mutation-induced alternative splice forms (Table S3). In the translation, we allowed for different transcripts from each SCM. The HLA types for each sample were adopted from the TCGA pancan immune working group (Synapse ID: syn5974636). NetMHC4 and NetMHCpan-3.0 (Andreatta and Nielsen, 2016) were used to predict the binding affinity between epitopes and the major histocompatibility complex (MHC) and showed a high concordance in total predicted neoantigens (Pearson = 0.94, Supplementary Figure S2). We found that alternative splice forms for some important genes related to tumorigenesis, including *SMARC1*, *KDM6A*, and *NOTCH1*, are highly immunogenic and can contain 40 or more unique neoantigens (Fig. 3.6A) (Dalgliesh et

al., 2010; Papadakis et al., 2015). In addition, the mean number of neoantigens across SCMs from NetMHCpan-4.0 and NetMHCpan-3.0 are 2.0 and 2.5, respectively, which are both higher than the average number of around 1 for non-synonymous mutations. Furthermore, 28 genes contain recurrent neoantigen events (more than or equal to three) across samples (Fig. 3.6B). In particular, GATA3 has the highest recurrence and GATA3 SCMs were mutually exclusive with other mutation types (Fig. 3.6C). The CA deletion at chr8:8111433 disrupts the canonical splice site and an alternative splice site is used for creating the alternative splice form, which results in a frame shifted protein product spanning the Zinc-finger domain (Figs. 3.6D and 3.6E). 19 unique neoantigen peptide sequences were mapped to the frameshifted protein product for the 16 samples (Fig. 3.6F). We were further able to validate one alternative peptide sequence using mass spectrometry data from a recent proteogenomics study on 77 TCGA breast cancer patients (Mertins et al., 2016). For one sample with the highly recurrent and expressed GATA3 SCM, we used MSGF+ to search publicly available mass spectrometry data for evidence of alternative GATA3 peptides. Fig. 3.6G shows one identified mass spectrum supporting one alternative GATA3 peptide, which covers two immunogenic peptides (KPKRRLPG and LIKPKRRLPG) predicted in TCGA-AR-A1AP.





Distribution of neoantigens predicted by NetMHCpan and NetMHC4. Genes with highest number of neoantigens labeled. Mean value for each tool indicated by X and labeled. (B) Genes with the largest recurrence of predicted neoantigens across the dataset. *GATA3* shows the highest recurrence. (C) Mutual exclusivity of protein-affecting mutation (PAM), frame-shifting indel (FS), in-frame indel (IF) and splice site-creating mutations (SCM) in *GATA3*. (D) IGV screenshot of *GATA3* splice site-creating mutation which disrupts the canonical splice site and utilizes a cryptic splice site 7 bp downstream. Mutant reads highlighted in red and normal reads in purple. CA deletion indicated in figure. (E) Predicted functional domains disrupted due to the recurrent splice site-creating mutation in *GATA3*. (F) Predicted neoantigen peptide sequences mapped to the frameshifted protein product for samples with *GATA3* splice site-creating mutations. (G) Mass spectrum of GATA3 peptide in TCGA-AR-A1AP.



Figure 3.7. PD-L1, PD-L2, PD-1, CD8A and CD8B expression. (A) Expression comparison of PD-L1, PD-L2, and T cell markers: PD-1, CD8A, CD8B between samples with (case) and without (control) splice site-creating mutations across 6 cancer types. Symbols: * is p-value less than 0.05, ** is p-value <0.01, *** is p-value <0.001, n.s. is not significant.

High neoantigen burden is associated with an elevated immune response (Turajlic et al., 2017). To test whether SCMs affect immune response, we compared the expression of T-Cell markers PD-1, CD8A and CD8B and PD1 immune checkpoint blockades PD-L1 and PD-L2 (Fig. 3.7). We selected 6 cancer types (BRCA, BLCA, HNSC, LUAD, LUSC and SKCM) with sufficient samples containing SCMs for adequate statistical power. Both T-Cell markers (PD-1, CD8A and CD8B) and immune checkpoint blockade PD-L1 show increased expression in samples with SCMs compared to samples without SCMs (Fig. 3.7), indicating alternative splice forms induced by SCMs increase the overall immunogenicity of these cancers. The highly expressed PD-L1 suggests PD-L1 immunotherapy as potential treatments for samples containing SCMs.

3.3 Methods

DATASET DESCRIPTION

Aligned RNA-Seq bam files were analyzed using the ISB google. These cancer types are Acute Myeloid Leukemia [LAML], Adrenocortical carcinoma [ACC], Bladder Urothelial Carcinoma [BLCA], Brain Lower Grade Glioma [LGG], Breast invasive carcinoma [BRCA], Cervical squamous cell carcinoma and endocervical adenocarcinoma [CESC], Cholangiocarcinoma [CHOL], Colon adenocarcinoma [COAD], Esophageal carcinoma [ESCA], Glioblastoma multiforme [GBM], Head and Neck squamous cell carcinoma [HNSC], Kidney Chromophobe [KICH], Kidney renal clear cell carcinoma [KIRC], Kidney renal papillary cell carcinoma [KIRP], Liver hepatocellular carcinoma [LIHC], Lung adenocarcinoma [LUAD], Lung squamous cell carcinoma [LUSC], Lymphoid Neoplasm Diffuse Large B-cell Lymphoma [DLBC], Mesothelioma [MESO], Ovarian serous cystadenocarcinoma [OV], Pancreatic adenocarcinoma [PAAD], Pheochromocytoma and Paraganglioma [PCPG], Prostate adenocarcinoma [PRAD], Rectum adenocarcinoma [READ], Sarcoma [SARC], Skin Cutaneous Melanoma [SKCM], Stomach adenocarcinoma [STAD], Testicular Germ Cell Tumors [TGCT], Thymoma [THYM], Thyroid carcinoma [THCA], Uterine Carcinosarcoma [UCS], Uterine Corpus Endometrial Carcinoma [UCEC], Uveal Melanoma [UVM]

MISPLICE PIPELINE

The MiSplice pipeline was developed to detect mutation-induced splicing events from RNA-Seq data. It is written in Perl and incorporates two standard tools, samtools and MaxEntScan. The pipeline is fully automated and can run multiple jobs in parallel on LSF cluster. It executes the following steps:

1) Splitting large maf file into multiple smaller files with less mutations (currently, the default setting is 200).

2) Discovering splicing junctions within 20bps of the mutation with at least 5 supporting reads with mapping quality Q20 and then filtering canonical junctions by using the Ensembl 37.75 database. We selected 20bp as a cut-off since it is the farthest distance from the splice junction in a splice region.

3) Computing the number of supporting reads of above cryptic splice sites for control samples without mutations (Table S1).

4) Calculating the splicing scores for the cryptic splice sites via MaxEntScan.

5) Reporting the depth of each cryptic splice site via Samtools.

6) Filtering cryptic sites which fall in HLA loci or less than 5% of reads at the genomic location supporting the alternative junction of interest.

7) Further filtering cryptic sites by comparing the supporting reads in control samples.The final reported cryptic sites must stand as top 5% for the number of supporting reads in the case (with mutation).

SPLICE SITE SCORE ESTIMATION

For each cryptic splice site and nearby canonical splice site, the corresponding nucleotide sequences were first extracted for both the mutant and reference sequences (9-mer and 23-mer for donor and acceptor, respectively). Their splice scores as potential donor or acceptor sites were then estimated using MaxEntScan.

NEOANTIGEN PREDICTION

86

For each predicted SCM, we use a curated RefSeq transcript database (version 20130722) to obtain the translated protein sequences for transcript containing alternative splice forms induced by SCMs. Different length of epitopes (8mer, 9mer, 10mer and 11mer) are constructed from the translated protein sequence. We use NetMHC3pan(Nielsen and Andreatta, 2016) and NetMHC4(Andreatta and Nielsen, 2016) to predict the binding affinity between epitopes and MHC. Epitopes with binding affinity <=500nM which are also not present in the wild-type transcript are extracted from the following neoantigen analysis.

MANUAL REVIEW

All splice site-creating mutations were manually reviewed using the integrative genomics viewer (<u>http://software.broadinstitute.org/software/igv/</u>). Mutations were placed into one of three categories: Pass, Complex, and No Support. Mutations were classified as complex if more than one alternatively spliced product was observed for the mutated sample.

CODE AVAILABILITY

MiSplice is written in Perl and is freely available from GitHub at https://github.com/dinglab/misplice under the GNU general public license. MiSplice uses several independent tools and packages, including SamTools and MaxEntScan, all of which are likewise freely available, but which must be obtained from their respective developers. The MiSplice documentation contains complete instructions for obtaining and linking these applications into MiSplice.

MINI GENE SPLICING ASSAY

Exons of interest and approximately 150 bp of their flanking intron sequences were PCR amplified from HEK293T genomic DNA using primers carrying restriction enzyme sites for BamH1 and Mlul. PCR products were cleaned up using NucleoSpin PCR Cleanup (Macherey-Nagel) or DNA Clean and Concentrator-5 Kit (Zymo Research) and digested with BamHI and MIuI. The digested pCAS2.1 vector and PCR products were ligated using T4 DNA Ligase (NEB). Mutations were introduced via Q5 Site-Directed Mutagenesis (NEB). WT and MUT constructs were confirmed by sequencing of the insert region. The plasmids were transiently transfected into HEK293T cells using Lipofectamine 2000 (ThermoFisher Scientific). 24 hours post transfection, cDNA was synthesized using 2 to 3 ug of total RNA with the Superscript III First-Strand Synthesis System (ThermoFisher Scientific) and priming with Oligo(dT)20. Finally, cDNA was amplified using pCAS-KO1-(5'-TGACGTCGCCGCCCATCAC-3') and pCAS-R (5'-ATTGGTTGTTGAGTTGGTTGTC-3') and the alternative splicing patterns were evaluated on a 2.5% agarose gel with ethidium bromide. Qiaquick Gel Extraction Kit (Qiagen) was used to purify bands for sequencing (Table S3.1-3.2, Figure S3-S6).

88



Figure S2: Mini-gene splicing assay.

Table 3.1: Mutation information for splice site-creating mutations validated in minigene

assay.

Gene	Sample	Position	Refere	Muta	Coding	Amino	Transcript
			nce	nt	Change	Acid	
						Change	
ARID2	TCGA-FS-A1Z3	12:	Т	А	c.1787T	p.Val596	ENST0000334
		46243434			>A	Glu	344

TSC2	TCGA-EE-A17Y	16:209870	С	А	c.89C>A	p.Ser30Ty	ENST0000219
		5				r	476
CDH1	TCGA-GC-A3l6	16:688636	С	G	c.2392C	p.Leu798	ENST0000219
		53			>G	Val	476
TP53	TCGA-FG-A60J	17:757715	Т	G	c.783-	p.X261_s	ENST0000269
		7			2A>C	plice	305
RAD5	TCGA-32-1982	17:567723	А	G	c.234A>	p.Thr78Th	ENST0000337
1C		80			G	r	432
BCOR	TCGA-DM-A1HA	X:399220	А	G	c.4123C	p.Arg1375	ENST0000378
		49			>T	Trp	444.4
BAP1	TCGA-CJ-4637	3:	Т	С	c.233A>	p.Asn78S	ENST0000460
	and TCGA-B0-	52442512			G	er	680
	5107						
PARP	TCGA-66-2791	1:2265508	G	А	c.2817C	p.Ser939	ENST0000366
1		31			>T	Ser	794
BRCA	TCGA-D6-6823	17:412568	А	С	c.301+2	p.X101_s	ENST00000471
1		83			T>G	plice	181
PTEN	TCGA-06-2559	10:896929	G	Т	c.477G>	p.Arg159	ENST0000371
		93			Т	Ser	953
KMT2	TCGA-55-7994	11:118366	G	Т	c.5423G	p.Trp1808	ENST0000534
А		474			>T	Leu	358

Table 3.2: Predicted alternative and wild-type RT-PCR splice products from mini-gene splicing assay.

Туре	RT-PCR Sequence
pCAS2.1	TGACGTCGCCGCCCATCACGCCTCCAGGCTGACCCTGCTGACCCTCCTGCTG
	CTGCTGCTGGCTGG GG ATAGAGCCTCCTCAAATCCAAATGCTACCAGCTCCAG

	CAGCCAAGATCCAGAGAGTTTGCAAGACAGAGGCGAAGGGAAGGTCGCAACA
	ACAGTTATCTCCAAGATGCTATTCGTTGAACCCATCCTGGAGGTTTCCAGCTTG
	CCGACAACCAACTCAACAACCAAT
BAP1 Wild Type	CCCTGTATATGGATTTATCTTCCTGTTCAAATGGATCGAAGAGCGCCGGTCCC
	GGCGAAAGGTCTCTACCTTGGTGGATGATACGTCCGTGATTGAT
	GTGAATAACATGTTCTTTGCCCACCAG
BAP1 Mutant	CCCTGTATATGGATTTATCTTCCTGTTCAAATGGATCGAAGAGCGCCGGTCCC
	GGCGAAAGGTCTCTACCTTGGTGGATGATACGTCCGTGATTGAT
TP53 Wild Type	CTCGCTTAGTGCTCCCTGGGGGGCAGCTCGTGGTGAGGCTCCCCTTTCTTGCG
	GAGATTCTCTTCCTCTGTGCGCCGGTCTCTCCCAGGACAGGCACAAACACGCA
	CCTCAAAGCTGTTCCGTCCCAGTAGATTACCA
TP53 Mutant	CTCGCTTAGTGCTCCCTGGGGGGCAGCTCGTGGTGAGGCTCCCCTTTCTTGCG
	GAGATTCTCTTCCTCTGTGCGCCGGTCTCTCCCAGGACAGGCACAAACACGCA
	CCTCAAAGCTGTTCCGTCCCAGTAGATTACCACGA
BCOR Wild	CTTGCCATCGGCATTCTCCACGTAGTATTCCCCTGTCAGTGGCAATCCCCGCC
Туре	TGGACTCCTGAGGGATCAAGTGTTTGGTTTTGCACAGTCTCTTCCCGGATGGC
	TTCTCGCTGTTGTCGGTGTATTTCTGCAGCAGGGAGGCAGCCTGGCAATCCTC
	TTCTTCGTCTGCACACAGCACATCTGTCTTCTGGTTTTCTTTAATTTTCTGCTGT
	TTGGCAGGCGGCCTGGAGGCTGGTGCGCAGCTTGGCTGAGCCTGCTTTTGC
	CGCCTGCACTGGTGGATGAAAGACTCTTCATGGGCGGAGAGCCGGAGAACAC
	AGGCAAGC
BCOR Mutant	CTGGACTCCTGAGGGATCAAGTGTTTGGTTTTGCACAGTCTCTTCCCGGATGG
	CTTCTCGCTGTTGTCGGTGTATTTCTGCAGCAGGGAGGCAGCCTGGCAATCCT
	CTTCTTCGTCTGCACACAGCACATCTGTCTTCTGGTTTTCTTTAATTTTCTGCTG
	TTTGGCAGGCGGCCTGGAGGCTGGTGCGCAGCTTGGCTGAGCCTGCTTTTTG
	CCGCCTGCACTGGTGGATGAAAGACTCTTCATGGGCGGAGAGCCGGAGAACA
	CAGGCAAGC

RAD51C Wild	AAGTTGGGATATCTAAAGCAGAAGCCTTAGAAACTCTGCAAATTATCAGAAGAG
Туре	AATGTCTCACAAATAAACCAAGATATGCTGGTACATCTGAGTCACACAAGAAGT
	GTACAGCACTGGAACTTCTTGAGCAGGAGCATACCCAGGGCTTCATAATCACC
	TTCTGTTCAGCACTAGATGATATTCTTGGGGGTGGAGTGCCCTTAATGAAAACA
	ACAGAAATTTGTGGTGCACCAGGTGTTGGAAAAACACAATTATG
RAD51C Mutant	AAGTTGGGATATCTAAAGCAGAAGCCTTAGAAACTCTGCAAATTATCAGAAGAG
	AATGTCTCACAAATAAACCAAGATATGCTG
KMT2A Wild	CAGTGGGATGTTACCAAACGCAGTGCTTCCACCTTCACTTGACCATAATTATGC
Туре	TCAGTGGCAGGAGCGAGAGGAAAACAGCCACACTGAGCAGCCTCCTTTAATG
	AAGAAAATCATTCCAGCTCCCAAACCCAAAGGTCCTGGAGAACCAGACTCACC
	AACTCCTCTGCATCCTCCTACACCACCAATTTTGA
KMT2A Mutant	TTGCAGGAGCGAGAGGAAAACAGCCACACTGAGCAGCCTCCTTTAATGAAGAA
	AATCATTCCAGCTCCCAAACCCAAAGGTCCTGGAGAACCAGACTCACCAACTC
	CTCTGCATCCTCCTACACCACCAATTTTGA
PARP1 Wild	CTTTGACACTGTGCTTGCCCTTGGGTAACTTGCTGATATGTGAAGCGTGCTTCA
Туре	GTTCATAC
PARP1 Mutant	TGATATGTGAAGCGTGCTTCAGTTCATAC
BRCA1 Wild	ACTCCAAACCTGTGTCAAGCTGAAAAGCACAAATGATTTTCAATAGCTCTTCAA
Туре	CAAGTTGACTAAATCTCGTACTTTCTTGTAGGCTC
BRCA1 Mutant	CTGTGTCAAGCTGAAAAGCACAAATGATTTTCAATAGCTCTTCAACAAGTTGAC
	TAAATCTCGTACTTTCTTGTAGGCTC
ARID2 Wild	AACGGTCTTTCCAAATCATACAGTGAAGAGAGTGGAGGATTCCAGTAGCAATG
Туре	GGCAGGCACATATTCATGTGGTAGGAGTAAAACGGAGGGCTATACCACTTCCC
	ATTCAGATGTACTATCAGCAGCAACCAGTTTCTACTTCTGTTGTTCGTGTTGATT
	CTGTTCCTGATGTATCTCCTGCTCCTTCACCTGCAG
APID2 Mutant	
	ACGUTCTTTCCAAATCATACAGTGAAGAGAGTGGAGGATTCCAGTAGCAATG
DTEN Wild Type	TTOCACA ATATOCTTTCAACACCATAACCCACACCACCACCACCAACTATCAAAC
----------------	--
	CCTTTTGTGAAGATCTTGACCAATGGCTAAGTGAAGATGACAATCATGTTGCAG
	CAATTCACTGTAAAGCTGGAAAGGGACGAACTGGTGTAATGATATGTGCATATT
	GAAGTAAGGACCAGAGACAAAAAG
PTEN Mutant	TTGCACAATATCCTTTTGAAGACCATAACCCACCACAGCTAGAACTTATCAAAC
	CCTTTTGTGAAGATCTTGACCAATGGCTAAGTGAAGATGACAATCATGTTGCAG
	CAATTCACTGTAAAGCTGGAAAGGGACGAACTGGTGTAATGATATGTGCATATT
	GAA
CDH1 Wild	GACTTTGACTTGAGCCAGCTGCACAGGGGGCCTGGACGCTCGGCCTGAAGTGA
Туре	CTCGTAACGACGTIGCACCAACCCTCATGAGTGTCCCCCGGTATCTTCCCCGC
	CCTGCCAATCCCGATGAAATTGGAAATTTTATTGATGAA
CDH1 Mutant	GACTTTGACTTGAGCCAGCTGCACAGGGGGCCTGGACGCTCGGCCIGAAGIGA
	CTCGTAACGACGTTGCACCAACCCTCATGAGTGTCCCCCG
TSC2 Wild Type	AGGGGTTTTCTGGTGCGTCCTGGTCCACCATGGCCAAACCAACAAGCAAAGAT
	TCAGGCTTGAAGGAGAAGTTTAAGATTCTGTTGGGACTGGGAACACCGAGGCC
	AAATCCCAGGTCTGCAGAGGGTAAACAGACGGAGTTTATCATCACCGCGGAAA
	TACTCACA
	TACTGAGA
TSC2 Mutant	AGGGGTTTTCTGGTGCGTCCTGGTCCACCATGGCCAAACCAACAAGCAAAGAT
	AAATCCCAG



Figure S3. *ARID2, BCOR, BRCA1* **Mini-Gene results. Related to Figure 3.5.** (A,D,G) DNA chromatograms verifying *ARID2, BCOR, and BRCA1* wildtype and mutant sequencing results, respectively. Mutation position is highlighted. (B,E,H) Reverse transcriptase PCR (RT-PCR) with wild type and mutant plasmids, results in triplicate. Numbered bands are sequenced for confirmation. (C,F,I) DNA chromatograms of RT-PCR bands sequenced. Highlighted sequence indicates boundary of pCAS2.1 plasmid.



Figure S4. PTEN, PARP1, KMT2A Mini-Gene results. Related to Figure 3.5. (A,D,G) DNA

chromatograms verifying *PTEN*, *PARP1*, and *KMT2A* wildtype and mutant sequencing results, respectively. Mutation position is highlighted. (B,E,H) Reverse transcriptase PCR (RT-PCR) with wild type and mutant plasmids, results in triplicate. Numbered bands are sequenced for confirmation. (C,F,I) DNA chromatograms of RT-PCR bands sequenced. Highlighted sequence indicates boundary of pCAS2.1 plasmid.



Figure S5. RAD51C, TP53, TSC2 Mini-Gene results. Related to Figure 3.5. (A,D,G) DNA

chromatograms verifying *RAD51C, TP53, and TSC2* wildtype and mutant sequencing results, respectively. Mutation position is highlighted. (B,E,H) Reverse transcriptase PCR (RT-PCR) with wild type and mutant plasmids, results in triplicate. Numbered bands are sequenced for confirmation. (C,F,I) DNA chromatograms of RT-PCR bands sequenced. Highlighted sequence indicates boundary of pCAS2.1 plasmid.



Figure S6. *BAP1 and CDH1* **Mini-Gene results. Related to Figure 3.5.** (A,D) DNA chromatograms verifying *BAP1* and *CDH1* wildtype and mutant sequencing results, respectively. Mutation position is highlighted. (B,E) Reverse transcriptase PCR (RT-PCR) with wild type and mutant plasmids, results in triplicate. Numbered bands are sequenced for confirmation. (C,F) DNA chromatograms of RT-PCR bands sequenced. Highlighted sequence indicates boundary of pCAS2.1 plasmid.

CELL CULTURE

HEK293T cells were cultured in Dulbecco's modified Eagle's medium (DMEM) supplemented with fetal bovine serum (FBS) and penicillin streptomycin.

STATISTICAL ANALYSES

MiSplice assesses the significance of the number of reads supporting the predicted alternative splice junction by comparing to read counts from a control cohort. Specifically, a frequency distribution is constructed from the control cohort, from which threshold values for 5% and 95% tails on the left and right, respectively, are determined. A series of logic tests is then conducted to discern the best explanation of the data. Possible verdicts are low or high expression if the datum is outside the 5% or 95% thresholds, respectively, average expression if no thresholds are exceeded, or no expression in this tissue if the thresholds are zero.

3.4 Discussion

In this study, we applied our newly developed bioinformatics tool called MiSplice (Mutation Induced Splicing) to systematically analyze splice site-creating events that arise from somatic mutations. Our analysis shows MiSplice reliably identifies SCMs across multiple cancer types. Existing studies have largely focused on splice-disrupting events in known splice sites, but the current study substantially extends our knowledge into the realm of splice site-creating mutations in human cancer. For instance, we found 1,016 splice site mutations not only disrupt the canonical splice site, but also create an alternative splice site. We also found that hundreds of mutations that would traditionally be classified as missense, silent, indel, and nonsense are really acting as SCMs. Many important cancer-related genes harbor these mutations, such as *TP53*, *ATRX*, *BAP1*,

CTNNB1, *RB1*, etc. It is noteworthy that we found five splice site-creating mutations in *ATRX* among 288 lower grade glioma cases, likely leading to the disruption of ATRX function. A previous study has shown that loss of wild-type *ATRX* is associated with tumor growth in glioma (Koschmann et al., 2016).

Characterization of these alternative splice events show that most SCMs have a higher splice score, as measured by MaxEntScan, in the post-mutation alternative splice site as compared to the reference. These results are consistent with the preferential selection of these alternative sites as new splicing forms. For the splice-site mutation, the splice score associated with the canonical junction is coincidently decreased after mutation. However, while there is no difference in splice scores of canonical junctions before and after missense and silent mutations, the alternative splice site was often strengthened after mutation. This suggests silent and missense mutations instead act as modifiers of splicing by creating or strengthening cryptic sites within the exon as opposed to disrupting the canonical splice site. In addition, we found a significant enrichment of mutations at the -3 position in the 3' splice site, the two dominant sequence contexts being aGag and agGag, where G is at the -3 position.

In cases where the mutation is retained in the alternative splice junction, we distinguish mutations with two further categories, splice-in and splice-out. For splice-in mutations, we can characterize the association between mutations and cryptic splicing forms. For example, we found high concordance for RNA-seq reads supporting alternatively spliced junctions and mutations, suggesting the association between mutations and cryptic splicing forms.

The current study has greatly extended the insight into the transcriptional ramifications of genomic alterations by identifying nearly 1,964 alternative splice sites introduced by somatic mutations and functionally validating ten of eleven variants in a mini-gene splicing assay. These events were conventionally annotated as missense, silent, splice site, nonsense, or other mutations when, in fact, we have shown that they often create cryptic splice sites. The relative abundance of the alternative and wild-type product suggests varying levels of junction usage, depending on the context of the mutation, and emphasizes the importance of validating predictions using a functional assay to understand the full biological consequence. The alternative products may be therapeutically targetable in some cancer patients. For example, the targeting of neoantigens shows promising result in treating melanoma patients (Carreno et al., 2015). By further evaluating human leukocyte antigen (HLA) genotypes and binding affinities to the major histocompatibility complex (MHC), it is likely that new neoantigens from cryptic splice sites may be discovered. The current study reveals that alternative splice forms induced by splice site-creating mutations are highly immunogenic and correlated with a high T-Cell immune response and an elevated PD-L1 expression, suggesting potential for immunotherapy in these samples. Further investigation of the cryptic splice sites by mass spectra

3.5 References

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<u>Chapter 4: Rare germline and somatic</u> <u>splice-site-creating mutations disrupt</u> <u>exon definition in cancer genomes</u>

Contribution: I developed the entire code for translation of MiSplice variants. I manually reviewed many variants validated by our computational tool, performed all downstream analyses, wrote the entire text, created all Figures, and performed entire mini-gene splicing assay for 7 variants tested.

4.1 Introduction

Mutations can be germline or somatic in nature. Germline mutations are inherited from parents or acquired *de novo*. Somatic mutations are acquired throughout an organism's lifetime in individual cells due to genetic and environmental factors such as chemicals and radiation. Most of the damage in the DNA is repaired, but sometimes the alterations are fixed. Mutations in genic and regulatory regions can affect gene function in several ways by causing loss or gain of function, altering transcript splicing, and increasing or decreasing gene expression level. In combination, such variations can disrupt normal gene function and alter cellular response to regulation giving the cell a selective advantage to proliferate autonomously. The underlying genomic architecture of individual species alters the resulting phenotypic variability. Understanding how differences to gene architecture can alter a genes function is essential to understanding many biological questions including disease diagnosis, progression and treatment. Current studies suggest that between 70-95% of human genes harbor multiple mRNA transcripts (Johnson et al.; Matlin et al.; Modrek and Lee). Different mRNA transcripts are created by the process of alternative splicing which expands the complexity and information content of the eukaryotic genome and allows for tissue, developmental, or temporally expressed isoforms which can perform alternate functions. Understanding how genomic variation in tumors can contribute to alternative splicing defects is an actively expanding field.

For the past decade, cancer genomics studies have focused on identifying and validating germline and somatic mutations by comparing patients tumor samples to their normal tissue. The integration of transcriptomic and proteomic data provides valuable insight as to the biological factors contributing to the cancer genome (Cieslik and Chinnaiyan). A database to explore splice-junction usage across TCGA (Sun et al.) as well as a global analysis of splicing across the TCGA cohort (Kahles et al.) were recently published expanding the current paradigm about the exceptions and rules by which splicing is dysregulated in cancer. While this recent publication analyzed the global landscape of alternative splicing across the TCGA pan-cancer cohort, they limited their analysis to germline mutations that are known to also be somatic in origin, greatly limiting their prediction power (Kahles et al.).

Earlier this year, we published a bioinformatic tool that systematically predicts somatic splice-site-creating mutations across 33 cancer types (Jayasinghe et al.). We identified nearly 2,000 somatic splice-site-creating mutations with 26% and 11% of mutations being initially mis-classified as missense and silent mutations, respectively. This novel finding lead us to question the burden of similar mutations in the germline of cancer patients, as this is an underexplored question in the field. Recently, The Cancer Genome Atlas consortium published a curated mutation profile of germline (Huang et al.) and somatic (Bailey et al.; Ellrott et al.) mutations across cancer genomes. These timely and thorough scientific publications have provided a sound basis for exploring the splice-site-creating landscape in both a somatic and germline context.

Here we developed additional modules for MiSplice to facilitate accurate identification of thousands of rare germline and somatic splice-site-creating events while providing necessary supplemental scripts to infer the resulting transcriptional and translational effects. Using our stringent pipeline we identified 2,888 rare germline and somatic SCMs which were initially inaccurately annotated. We observed interesting patterns in genomic sequence contexts surrounding SCM containing exons (SCM+ exons) including increased nucleosome occupancy and an overall decrease in the novel splice form relative to the original exon size. The discovery of rgSCMs genome wide facilitated the proper annotation of hundreds of variants that could have direct implications in cancer. For example, one recurrent synonymous variant was identified in *RAD54L*, generally characterized as a DNA repair gene, suggesting this variant could be a novel pathogenic germline variant that has been missed in previous studies. Finally, we validated 6 rare

germline SCMs (rgSCMs) in cancer associated genes, including the *RAD54L* silent variant, and one common germline SCM (cgSCM) validating our computational pipeline and workflow for proper and efficient SCM identification.

4.2 Results

DATASET DESCRIPTION

To further explore the germline and somatic splice-site-creating landscape across cancer samples, we collected somatic and germline mutations calls from the MC3 working group (Ellrott et al.) and the Cancer Genome Atlas (TCGA) germline working group (Huang et al.). We sought to evaluate the splice-creating potential of variants conventionally annotated as SpliceDonorSNV, SpliceAcceptorSNV, Missense, Synonymous, IntronicSNV (near the splice site) and Nonsense mutations (Figure 4.1A). With stringent filtering (Methods) we evaluated >3,000,000 somatic mutations, > 10,000,000 rare germline mutations and >20,000,000 common germline variants in a splicing context using the MiSplice pipeline (Jayasinghe et al.) (Figure 4.1C). For rare germline mutations and somatic mutations, we defined a set of control samples in the same cancer type that lacked the same mutation of interest. With respect to common germline mutations, samples were placed in a case group or control group depending on their mutation status for each mutation of interest. We conducted the splice-site-creating mutation analysis using the ISB google cloud using MiSplice (mutation-induced splicing) that systematically evaluates mutations in a splicing context using RNA-seg data. MiSplice manages large analyses using parallel computation to search for alternative splice junctions within

windows of ±20 bp from the mutation of interest. We identified high confidence rare and common germline splice-site-creating mutations (rgSCMs, cgSCMs) and somatic splice-site-creating mutations (sSCMs).



Figure 4.1. Somatic and Germline splice-site-creating variant discovery. (A) Examples of splicesite-creating mutations (SCM) for different annotated mutation types. (B) Breakdown of rare germline SCMs and somatic SCMs, plotted by conventional annotation type. Cancer associated genes are highlighted in black. (C) The updated MiSplice workflow can process millions of somatic and germline variants on the ISB-Google Cloud or on a local compute cluster. Mutation calls and RNA-Sequencing bams are provided as input, and variants in close proximity to a detected splice junction are maintained for additional filtering. Canonical junctions, low coverage, novel junction presence in the controls, large predicted exon sizes and certain genes including HLA are filtered out. Finally an added annotation

module determines allele frequency for each variant in the gnomAD database, predicts the mutation effect using TransVar and predicts the resulting wildtype and novel protein isoforms

GERMLINE AND SOMATIC SPLICE-SITE-CREATING MUTATION DISCOVERY

The expanded discovery of somatic splice-site-creating mutations across 33 cancer types identified 1,782 sSCMs (Figure 4.1B). 237 SCMs are annotated to genes commonly associated with either adult or pediatric cancer. Some of the most highly recurrent events were reported in our previous publication (Jayasinghe et al.) but novel sSCMs were identified by expanding our analysis to this larger MC3 dataset (Table 1). Newly identified sSCMs in cancer associated genes were manually reviewed for accuracy.

In expanding our analysis of splice-site-creating mutations to the germline, we identified mis-annotated germline mutations that show evidence of creating new splice sites. After processing rare variants from 33 cancer types a total of 14,709 cancer specific unique variants were identified as having splice-creating-potential. Additional filters included removing sites with more than 5% of controls containing the alternative splicing event (>2 reads), samples with less than 20 controls, and finally filtered out splicing events identified in highly homologous genes including MUC*, AHNAK* and CRIPAK (Methods) leaving 4,295 potential splice-creating variants. Of the 4,295 variants, 1,121 variants had a maximum allele frequency across populations in the controls subset less than 0.05% as derived from gnomAD browser - version 2.1 (Lek et al.). Finally after peptide translation and additional filtering, we were left with 1,106 high confidence single nucleotide variants

classified as rgSCM events encompassing 995 unique variants from 852 genes, and 63 in cancer associated genes.

Additional filters were incorporated into MiSplice to filter out novel splice isoforms present at a higher frequency in the cancer cohort population. Finally all variants underwent additional annotation post filtering to capture filter out events present at a high frequency in the population, and we incorporated TransVar to provide the most up to date annotation and predict the resulting peptide change. For the entirety of this analysis we will focus on single nucleotide variants.

A majority of SCMs were conventionally annotated as SpliceAcceptorSNVs followed by Missense, Synonymous, SpliceDonorSNVs, Nonsense and a handful IntronicSNVs (Figure 4.1B). This distribution held true for both rare germline and somatic events. Interestingly, our expanded analysis identified 170 novel rare germline SCMs conventionally annotated as synonymous variants and an additional X somatic SCMs not identified in our previous work. A remarkable number of SCMs were annotated to cancer associated genes (Table 1) including 63 rare germline SCMs and 237 somatic SCMs. We will take a closer look at the biological relevance of these events in a later section.

COMPARATIVE SEQUENCE CONTEXTS AND CHARACTERISTICS

With this large and high confidence list of somatic and germline splice-site-creating mutations across thousands of exons, we have the unique opportunity to collect relevant genomic information to characterize SCM positive exons. First we looked to determine

the location of the novel splice junction relative to the canonical splice site (Figure 4.2A). In knowing that the strongest splicing enhancer elements are located near the canonical splice site, we predicted many SCMs would cluster near the canonical site and taper off with increasing distance from the canonical site. As expected splice acceptor and splice donor variants are densely populated near the canonical splice site while synonymous and missense mutations are found deeper within the exonic region. After the mutation is introduced, the largest differences in transcript size post mutation (here on in referred to as effect size) are shared by conventionally annotated missense, synonymous and nonsense mutations, while smaller alterations are observed for mutations in close proximity to the splice site (Supplementary Figure 4.1A). Due to our restricted search window for identifying novel splice sites in proximity to mutations, we anticipated and gladly observed a direct correlation between the effect size and distance from the mutation to the canonical splice site (Supplementary Figure 1B).



Figure 4.2. Sequence contexts and characteristics of splice-site-creating mutations. (A) Cartoon of donor and acceptor splice-site-creating mutations (SCM). Example mutation is plotted in red relative to the novel splice site and canonical splice site. (B) Distribution of rgSCM and sSCM events relative to the distance from the canoncial splice site. Proportion of variants at each position is colored by conventional annotation type. (C) Comparison of the novel splice site score before and after mutation for rgSCM and sSCM broken down by the creation of a novel donor or acceptor splice site. Each point is colored by conventional annotation type. (D) Contrast the overall change in splice score before and after mutation for mutation for the novel and canoncial junctions. Positive values indicate a stronger novel splice score

change post mutation relative to canonical splice site. (E) For variants conventionally annotated as SpliceDonorSNV and SpliceAcceptorSNV, compare the difference in splice score pre and post mutation for the novel and canonical splice site.

To confirm the mutation is indeed a splice-site-creating mutation, we compared the novel splice score of the genomic sequence before and after mutagenesis (Figure 4.2A). Rare germline and somatic SCMs alike globally exhibited an increase in the novel splice score after mutation (deviating in the positive direction above the xy line), with few exceptions (Figure 4.2C). While strengthening a novel or cryptic splice site is necessary to induce alternative isoform usage, we wanted to determine if the canonical splice site was also disrupted due to the mutation. To characterize an overall splice score change, the difference in the splice score was calculated for both the novel (ΔN) and canonical (ΔC) splice site. The difference between ΔN - ΔC would be positive under conditions where the novel splice score change post mutation was stronger than the canonical site in the presence of the mutation, capturing the overall mutations effect on both sites. Figure 4.2D displays the distribution of the overall combined splice score change for each site by conventional annotation type. While all values trend in the positive direction, suggesting a stronger novel site change relative to canonical site change, conventionally annotated splice donor variants are densely clustered together. By taking a step back we can compare ΔN and ΔC separately. In Figure 4.2E, the difference in the novel score (ΔN) increases dramatically, while the canonical site (ΔC) is completely disrupted. On the other hand, for the donor site, the novel splice site doesn't change in the presence of the mutation, but the canonical site is disrupted.

This interesting phenomenon for the donor splice site holds true for the 294 conventionally annotated donor mutations and suggests that by simply disrupting the donor splice site, it's usage can be completely ablated and becomes unrecognizable, placing a stronger emphasis on the donor site relative to the acceptor splice site. Supporting this hypothesis, recently Wong et al. performed a massively parallel splicing assay on the introns of three genes to monitor the effects of all predicted changes to the 9 nucleotide 5' splice site, surveying a total of 32,768 unique donor sites (Wong et al.). From the assay, they determined disruption of the 5' splice site alone was enough to disrupt splicing, despite mutating nearby genomic regions.

The average exon in our dataset containing a splice-site-creating mutation is 289 bp long with a standard deviation of 668 bp. By comparing the relative exon sizes between each of the conventionally annotated mutation types, an increased canonical exon size was observed for missense (p=2.1e-08,1.9e-13) and synonymous sites (2.9e-05,9.8e-11) in comparison to the acceptor and donor splice sites, respectively. Nonsense mutations also showed evidence of increased exon size relative to the donor splice site (p=0.033) (Supplementary Figure 1A).

Next we chose to explore beyond the exon to determine if introns adjacent to the SCM positive exon contained additional information for SCM classification. Interestingly, splice acceptor mutations were more likely to be situated downstream of a larger intron relative to the intron upstream of the synonymous mutation (p=5.4e-05, Figure 4.1C). Similarly

the downstream intron size was larger for splice acceptor SNVs relative to introns downstream of missense (p=3.4e-05), synonymous (p=2.5e-09) and splice donor mutations (Supplementary Figure 1D). These initial observations suggest that while evaluating the splicing score in the presence and absence of the mutation is very informative, with a large enough dataset, we may be able to glean new information from the adjacent genomic sequences to strengthen the identification of putative SCM containing exons.

TRANSLATIONAL AND TRANSCRIPTIONAL IMPLICATIONS OF SCMs

Each variant was annotated according to their resulting translational consequence to interrogate overall trends in splicing dysregulation (Methods). For both somatic and rare germline SCMs, exon shrinkage events were observed at a higher frequency than exon extension. 91% and 89% of translated sites resulted in exon shrinkage trending towards an overall decrease in the novel exon size relative to the canonical exon (Figure 4.3A). The novel exon size decreased by an average of 21 and 33 bp for sSCM and rgSCM, respectively. From an evolutionary standpoint, exon size tends to be far more constrained than intron size. One hypothesis for this constraint is due to the intricate dance between chromatin remodeling and alternative splicing.

Within the nucleus, DNA is wrapped around histones in segments of <150 nucleotides. Nucleosomes are made up of eight core histones and are conveniently positioned on exons. The size of the average exon is ~150 nucleotides long providing strong evidence as to the evolutionary pressure selecting for exons around this length(Amit et al.; Schwartz et al.). Since splicing is a co-transcriptional process, splicing factors and histone marks work together to properly facilitate the splicing process(Jeong). With the aforementioned evidence suggesting a link between nucleosome occupancy and splicing, we performed a nucleosome occupancy enrichment analysis for the rgSCM and sSCMs using NucMap (Zhao et al.). The enrichment scores were much stronger at regions overlapping the exons containing splice-site-creating mutations relative to a 2kb window surrounding the exon of interest (Figure 4.3B). Additional studies have revealed stronger nucleosome enrichment for exons that are adjacent to long introns or have weaker splice sites (Spies et al.). To explore this phenomenon in our dataset, SCMs were binned by the canonical splice site score, and nucleosome enrichment was compared among the four binned groups by quartile, but no correlation was observed in our dataset (Supplemental Figure S7).



Figure 4.3. Novel spliced isoform exon expression and predicted translation. (A) Contrasting overall novel exon size relative to canonical exon size for rgSCM and sSCM events. Each point is colored by the predicted reading frame of the novel spliced isoform. (B) Enrichment analysis of nucleosome occupancy by NucMap on four different cell lines for all 2,888 predicted SCM containing-exons. Each plot is centered on the exon containing the novel splice-site-creating mutation. (C) Junction allele fraction (JAF) distribution by predicted reading frame. P-value reported by wilcox test comparing in frame and off-frame events within each SCM group. (D) Based on the location of the premature termination codon, each predicted protein product was classified as eliciting or escaping nonsense mediated decay based

on the 50 bp rule. P-value reported by wilcox test comparing both groups within each SCM group. Each point is colored by the predicted reading frame of the novel spliced isoform.

The resulting reading frame of the novel splice isoform was annotated as off-frame or inframe based on the size of the exon shrinkage or extension event. As expected, the junction allele fraction, or fraction of reads supporting the alternative splice form, was significantly lower for predicted off-frame SCMs compared to in-frame SCMs (wilcox test, sSCM:p=1.281e-12, rgSCM:p=5.617e-15) (Figure 4.3C). The expression of the resulting off-frame events is predicted to be lower at the RNA level because aberrantly spliced transcripts are often degraded by nonsense mediated decay (NMD), a process that identifies and degrades transcripts containing premature termination codons (PTCs). The general rule of thumb is PTCs located at least 50 bp upstream of the last exon-exon junction drive strong degradation, whereas those outside of this criteria are predicted to escape the degradation process (Brogna and Wen; Lewis et al.; Maguat et al.; Nagy and Maguat; Popp and Maguat; Venables; Weischenfeldt et al.). When applying the 50 bp rule to to our set, 97% of the off-frame spliced products are predicted to elicit-NMD (990 off-frame, 33 in-frame) while 79% of in-frame events are expected to escape-NMD (1465 in-frame, 400 off-frame) (Supplemental Figure S7). For escapee transcripts, we would expect to observe a higher expression of the novel splice form. Indeed, expression as measured by the junction allele fraction supported NMD-escaping transcripts maintaining a higher expression relative to the non-escapee group (wilcox, rgSCM:p=9.078e-15, sSCM:*p*=8.129e-16) (Figure 4.3D).

123

Even though some sites are predicted to undergo degradation, all SCMs were identified due to their expression in the RNA-Sequencing data. Most sites have JAF's lower than 50% suggesting some post transcriptional mechanisms by which the transcript is being degraded, although not to completion. It is important to note that it is very likely we are not capturing the full landscape of SCMs due to efficient degradation of mutation induced alternative transcripts.

LANDSCAPE OF SOMATIC AND GERMLINE SCMS GENOME-WIDE

We next explored the landscape of SCMs genome-wide. Figure 4.4A highlights the most highly recurrent genes in our dataset with *TP53* harboring somatic SCMs spanning several cancer types while *CBWD5* has many SCMs predominantly derived from BRCA. For the subset of highly recurrent SCM containing genes, rgSCM and sSCM almost always do not co-occur in the same gene with the exception of: *LZTR1* and *PTPN13*. Out of 2,163 unique genes harboring at least one SCM, 161 genes encompassing 417 variants had both a germline and somatic SCM reported. Several shared SCM positive genes overlap cancer associated genes including *CHEK2*, *CHD8*, *CASP8*, *FANCL*, *MUTYH*, *RAD51C*, *RPA1*, *BRCA1*, *EML4*, *FANCI*, *PARP3*, and *PARP4*. Interestingly, only *CHEK2* shared a mutation at the same splice donor site, but both events produced different transcriptional effects. Additionally, a conventionally annotated missense variant in *MLLT10* contained both an rgSCM and sSCM resulting in the same 251 bp exon shrinkage, and was validated in our minigene splicing assay (Figure 4.5D).



Figure 4.4. Splice-site-creating variants across genes and cancer types. (A) Overview of most highly recurrent splice-site-creating mutations colored by SCM type (germline or somatic). Values in plot indicate total number of SCMs identified for each gene-cancer type pair. (B) Overall change in gene expression (TPM) between SCMs and their cancer type cohort (wilcox test). Pvalue is indicated at the bottom of each plot and the average of each group are plotted as one value. Grouped by change in expression with a significant increase in expression being on the left and a significant decrease on the right.

Next we wanted to explore changes in gene expression relative to SCM presence. Using gene expression data from the UCSC XenaBrowser, we compared transcripts per million (tpm) values between the SCM mutants, TCGA Cancer Cohort and associated GTEX tissue cohort, when available. By comparing the expression of the SCM mutant to the

cancer type cohort, several SCMs exhibited differential expression including those in cancer associated genes (Figure 4.4B) (Wilcox test). The biological implications of the altered expression will need to formally evaluated in follow up studies.

HYPERMUTATOR PHENOTYPE OF RAD54L RGSCM

RAD54L is involved in DNA repair via homologous recombination and frequently undergoes copy number alterations depending on the tissue type of interest. For example, loss of heterozygosity in *RAD54L* is common in breast cancer and lower-grade gliomas(Nowacka-Zawisza et al.) and associates with longer progression free survival and chemosensitivity(Tang et al.), respectively, suggesting a tumor suppressor phenotype. More recently, the mechanism of cell proliferation and radio-resistance in glioblastoma was linked to CDC7 expression, a gene directly regulated by RAD54L(Li et al.). In contrast, in choroid plexus carcinomas the DNA repair gene was often amplified in tumors and necessary for proliferation (Tong et al.) in murine models, suggesting an oncogenic phenotype.



Signature.13



Figure 4.5. RAD54L and mini-gene splicing assay. (A) Splice junction quantification of te novel and canonical splice sites for two samples containing the RAD54L silent mutation and one normal sample. Genomic sequence of the novel and canonical splice site are highted below the exon pictogram. The novel change in novel and canonical splice score are indicated next to the genomic segments. (B) Mutational signature for RAD54L HNSC Mutant sample. (C) Mutational signature for overall HNSC cohort (D) Candidate splice creating variants validated by the mini-gene splicing assay. Exons of interest were

cloned into the pCAS2.1 vector and mutant (red) and wildtype (purple) plamids were transfected into 293T cells and total RNA was extracted to identify mutation induced alternatively spliced products.

In our sample set two conventionally annotated silent (p.G659G) mutations in *RAD54L* were identified more appropriately as an rgSCM and associated with a slightly higher overall expression relative to the head and neck squamous cell carcinoma (HNSC) cohort (wilcox p=0.099) but not in the testicular germ cell tumor (TGCT) (wilcox p=0.343). The rare germline mutation creates a new splice site (GC>GT) leading to a 58 bp exon shrinkage, frameshifting the remainder of the protein at exon 17 containing a Snf2 specific helical domain (HD2) domain and the C terminal Domain (CTD) (Heyer et al.) (Figure 4.5A). The novel splice junction is located within 55 bp of the last exon-exon junction and is thus predicted to escape NMD. The splicing score post-mutation increases dramatically from 0.98 to 8.73 and the variant allele fractions in the tumor and normal sample are comparable, 49% and 46% for the HNSC sample and 44% and 52% in the TGCT sample, respectively.

We next sought to evaluate whether the predicted rgSCM event in *RAD54L* can induce genomic changes relative to the associated cancer cohort. Both samples exhibit higher mutation rate relative to the rest of their cancer type cohort, with the TGCT sample being the highest mutated sample and the HNSC sample being one of the top 10 mutated samples. To evaluate whether the potentially pathogenic rare germline SCM mimics a hypermutator like phenotype, we evaluated the mutational signatures of each patient. After inspecting the mutational signature of the HNSC sample, signature 2 and signature 13 are the most prevalent followed by 1, 7 and 10 (Methods). In contrast, the HNSC
cancer type cohort did not exhibit a similar mutation signature, suggesting this sample is under a different evolutionary pressure relative to the cancer cohort. The joint signatures of 2, 10 and 13 in the HNSC sample are commonly found in patients undergoing local hypermutator phenotypes and with this sample being one of the 5 top mutated samples in HNSC (lacking mutations in other DNA-repair related genes) (Alexandrov et al.) suggests the *RAD54L* rgSCM variant may be indicative of the observed phenotype.

Finally, to functionally confirm the predicted alternatively spliced product in the lab, we performed a mini-gene splicing assay to validate rgSCMs in 6 cancer-associated genes and 1 common germline SCM (Figure 4.5D). Briefly, we used a pCAS2.1 splicing reporter mini-gene functional assay by cloning mutant exons into the pCAS2.1 vector (Gaildrat et al.) and transiently transfected into HEK293T cells. RNA is extracted after 24 hours and after generating cDNA synthesis, RT-PCR is performed to evaluate the presence of the novel splice isoform relative to the wild-type vector. *RAD54L* rgSCM was one of the seven germline SCMs confirmed in the assay (Figure 4.5D). Taken together, MiSplice was able to predict a recurrent putative rare germline SCM event improperly annotated as a synonymous mutation. While mutations disrupting DNA repair have been identified in another member of the RAD family *RAD51D* (Pelttari et al.), to date no pathogenic mutations have been identified in *RAD54L*.

4.4 Methods

DATASET DESCRIPTION

Aligned RNA-Seg bam files were analyzed using the ISB google. These cancer types are Acute Myeloid Leukemia [LAML], Adrenocortical carcinoma [ACC], Bladder Urothelial Carcinoma [BLCA], Brain Lower Grade Glioma [LGG], Breast invasive carcinoma [BRCA], Cervical squamous cell carcinoma and endocervical adenocarcinoma [CESC], Cholangiocarcinoma [CHOL], Colon adenocarcinoma [COAD], Esophageal carcinoma [ESCA], Glioblastoma multiforme [GBM], Head and Neck squamous cell carcinoma [HNSC], Kidney Chromophobe [KICH], Kidney renal clear cell carcinoma [KIRC], Kidney renal papillary cell carcinoma [KIRP], Liver hepatocellular carcinoma [LIHC], Lung adenocarcinoma [LUAD], Lung squamous cell carcinoma [LUSC], Lymphoid Neoplasm Diffuse Large B-cell Lymphoma [DLBC], Mesothelioma [MESO], Ovarian serous cystadenocarcinoma [OV], Pancreatic adenocarcinoma [PAAD], Pheochromocytoma and Paraganglioma [PCPG], Prostate adenocarcinoma [PRAD], Rectum adenocarcinoma [READ]. Sarcoma [SARC]. Skin Cutaneous Melanoma [SKCM]. Stomach adenocarcinoma [STAD], Testicular Germ Cell Tumors [TGCT], Thymoma [THYM], Thyroid carcinoma [THCA], Uterine Carcinosarcoma [UCS], Uterine Corpus Endometrial Carcinoma [UCEC], Uveal Melanoma [UVM].

MUTATION FILE DESCRIPTION

To evaluate somatic mutations we started with the mc3.v0.2.8.CONTROLLED.maf and filtered out mutations with the following MAF Filter Flags: StrandBias, pcadontuse, common_in_exac, oxog, contest, nonpreferredpair, ndp, badseq, broad_PoN_v2, leaving a total of 10,573,839 variants. With respect to mutation callers, single nucleotide polymorphisms and insertions and deletions were only maintained if there was agreement from two or more callers. Finally mutations with a variant allele fraction of less than 5%

were filtered out. Description of MC3 MAF file and filter flags found here: https://www.synapse.org/#!Synapse:syn7214402/wiki/405297. Final filtered somatic mutation file used for analysis found here: https://www.synapse.org/#!Synapse:syn12074532.

To evaluate germline mutations, individual VCFs were downloaded from Huang et al., 2018 (Huang et al.) encompassing TCGA cancer types listed in the Dataset Description. All variants were annotated using steps 6 and 7 of germline wrapper (https://github.com/ding-lab/germlinewrapper) against hg19 Homo Sapiens fasta file. After annotation, variants were maintained in a common germline MAF if allele frequency is greater than 0.01 and filtered into a rare MAF if allele frequency is less than 0.01 or if not reported. Common variants were only evaluated for BRCA. All rare variants were evaluated for splice-site-creating function using MiSplice. Furthermore, total reads supporting the alternatively spliced product were lowered for genes identified as significant in Huang et al., 2018 (Huang et al.) and re-run through MiSplice. After filtering, any variants with more than 10% of controls samples containing the alternatively spliced product were further filtered out leaving high confidence variants. For the 20,205,168 common variants in BRCA, in an initial screen only 72,245 unique variants were evaluated for splice-creating function. Of the 72,245 unique variants, 287 were shown to have splice-creating function in at least one sample. The variants with splice-creating function were then evaluated in the remaining samples. In total 74,383 variants were evaluated for splice-creating function. 29 variants with less than 10% of control samples exhibiting the alternatively spliced isoform were manually reviewed, resulting in 6 high confidence common germline splice altering variants.

Finally germline mutations were finally filtered as being rare if the allele frequency derived from the Genome Aggregation Database (gnomAD browser; http://gnomad.broadinstitute.org/; release 2.0.2) is less than 0.0005 AF (0.05%).

MISPLICE PIPELINE (SPLICE-SITE-CREATING MUTATION)

The MiSplice pipeline was developed to detect mutation-induced splicing events from RNA-Seq data. It is written in Perl and incorporates two standard tools, samtools and MaxEntScan. The pipeline is fully automated and can run multiple jobs in parallel on LSF cluster. It executes the following steps:

1) Splitting large maf file into multiple smaller files with less mutations (currently, the default setting is 200).

2) Discovering splicing junctions within 20bps of the mutation with at least 5 supporting reads with mapping quality Q20 and then filtering canonical junctions by using the Ensembl 37.75 database. We selected 20bp as a cut-off since it is the farthest distance from the splice junction in a splice region.

3) Computing the number of supporting reads of above cryptic splice sites for control samples without mutations.

4) Calculating the splicing scores for the cryptic splice sites via MaxEntScan.

5) Reporting the depth of each cryptic splice site via Samtools.

6) Filtering cryptic sites which fall in HLA loci or less than 5% of reads at the genomic location supporting the novel junction of interest.

MISPLICE GERMLINE FILTERING

All relevant scripts for filtering germline splice-site-creating mutations can be found: https://github.com/reykajayasinghe/MiSplice_Supplemental/.

1) Gene filter: Large exons with a high affinity for alternatively spliced products were filtered out including: AHNAK, AHNAK2, HLA family, FMN2, CRIPAK, IGH*, MUCIN family, RP11*, orf genes (Table X)

2) Remove events with less than 20 supporting controls.

3) Remove sites with greater than 5% of control samples having the same reported alternative splicing event.

4) Nearby Mutations and combining mutations in the same gene by cancer type: Combing the same splice-site-creating event into one entry when there are multiple mutations nearby one another for the same sample set. Combining the same SCM event across multiple patients into the same entry.

5) All sites are annotated with TransVar (Zhou et al.). Only variants with annotated variants in protein_coding transcripts that were classified by uniprot as TSL=1 were maintained for further analysis.

6) Finally, all single nucleotide variants were annotated with population frequencies derived from the genome aggregation database (gnomAD,http://gnomad.broadinstitute.org/).

SPLICE SITE SCORE ESTIMATION

For each cryptic splice site and nearby canonical splice site, the corresponding nucleotide sequences were first extracted for both the mutant and reference sequences (9-mer and 23-mer for donor and acceptor, respectively). Their splice scores as potential donor or acceptor sites were then estimated using MaxEntScan for single nucleotide polymorphisms.

MUTATIONAL SIGNATURE

For determining the mutation signatures of a the subset of samples, mutations were extracted from the MC3 public mutation file for the HNSC sample while mutations were controlled file extracted from the mutation for the TGCT sample (https://www.synapse.org/#!Synapse:syn7214402). Different mutation files were needed because all the mutations for the TGCT sample were removed from the public mutation file due to filter flags mostly considered "NonExonic." All mutations were run through Mutational Signatures in Cancer (MuSiCa) to evaluate mutation signatures for each patient.

NUCLEOSOME POSITIONING ANALYSIS

For determining the nucleosome positioning of the alternatively spliced exons, all exons containing a rare germline or somatic SCM were collected and converted to bed format.

Coordinates were lifted over to hg38 from hg19 using ucsc hgLiftOver (https://genome.ucsc.edu/cgi-bin/hgLiftOver) and coordinates were modified to match the input for NucMap (http://bigd.big.ac.cn/nucmap/Faq.php#q9) to extract enrichment scores of nucleosome binding sites across several cell lines including: hsNuc0270101, hsnuc0260501, hsNuc0390101, hsNuc0320101, hsNuc0070101, and hsNuc0020101. Using the browser analysis software normalized reads (RPM, reads per million) were extracted from the aforementioned cell lines and an enrichment score was calculated using the online software.

MANUAL REVIEW

All novel splice creating mutations were manually reviewed using the integrative genomics viewer (http://software.broadinstitute.org/software/igv/). Mutations were placed into one of three categories: Pass, Complex, and No Support. Mutations were classified as complex if more than one novel alternatively spliced product was observed for the mutated sample. After annotation, sites with large effect size (>100 bp) were manually reviewed for confirmation. Many events with very large new exons were recharacterized as ultra-short-introns.

TRANSLATION

All MiSplice Supplemental scripts are written in python (https://github.com/reykajayasinghe/MiSplice_Supplemental). Translation.py takes in the TransVar annotated matrix and defines important characteristics surrounding the SCM including: determining size of the upstream and downstream intron, size of the current exon, size of the novel splice isoform, overall change in size of the spliced product,

mutation position relative to the start and end of the exon, determines the resulting wildtype and mutant amino acid predictions for the SCMs, and finally predicts degradation by NMD. Some sites may be lost during this translation script due to improper annotation by transvar to a protein coding transcript or the lack of a known transcriptional start site in the input coding sequence file.

CODE AVAILABILITY

MiSplice is written in Perl and is freely available from GitHub at https://github.com/dinglab/misplice_gsSCM under the GNU general public license. MiSplice uses several independent tools and packages, including SamTools and MaxEntScan, all of which are likewise freely available, but which must be obtained from their respective developers. The MiSplice documentation contains complete instructions for obtaining and linking these applications into MiSplice.

MINI GENE SPLICING ASSAY

Exons of interest and approximately 150 bp of their flanking intron sequences were PCR amplified from HEK293T genomic DNA using primers carrying restriction enzyme sites for BamH1 and Mlul. PCR products were cleaned up using NucleoSpin PCR Cleanup (Macherey-Nagel) or DNA Clean and Concentrator-5 Kit (Zymo Research) and digested with BamHI and Mlul. The digested pCAS2.1 vector and PCR products were ligated using T4 DNA Ligase (NEB). Mutations were introduced via Q5 Site-Directed Mutagenesis (NEB). WT and MUT constructs were confirmed by sequencing of the insert region. The plasmids were transiently transfected into HEK293T cells using Lipofectamine 2000 (ThermoFisher Scientific). 24 hours post transfection, cDNA was synthesized using 2 ug

of total RNA with the Superscript III First-Strand Synthesis System (ThermoFisher Scientific) or igScript Reverse Transcriptase (Intact Genomics) and priming with Oligo(dT)20. Finally, cDNA was amplified using pCAS-KO1-(5'-TGACGTCGCCGCCCATCAC-3') and pCAS-R (5'-ATTGGTTGTTGAGTTGGTTGTC-3') and the alternative splicing patterns were evaluated on a 2.5% agarose gel with ethidium bromide. Qiaquick Gel Extraction Kit (Qiagen) was used to purify bands for sequencing. CELL CULTURE

HEK293T cells were cultured in Dulbecco's modified Eagle's medium (DMEM) supplemented with fetal bovine serum (FBS) and penicillin streptomycin.

SELECTION OF VARIANTS FOR VALIDATION

In selecting variants to validate, we focused on mutations in cancer associated genes and known cancer predisposition genes. After selected gSCMs from cancer predisposition genes, we then filtered sites in 1) low complexity regions to avoid creating non-specific primers; 2) exons containing variants with restriction enzyme cut sites in BamHI and Mlul; 3) exons containing variants in the first or last exon of a gene. In these cases signals from the UTR regions may disrupt the mini-gene splicing assay; 4) complex splicing events that couldn't be properly captured in a mini-gene splicing assay; 5) exons with small introns. For the mini-gene splicing assay, we selected exons of interest that were located far enough away from adjacent exons with specific enough primers to amplify 150 bp of the intronic region on either side of the exon of interest.

4.5 Discussion

Accurate and quick validation of somatic and germline variation is tantamount to personalized medicine. Our MiSplice pipeline has proven to properly classify and characterize splice-site-creating mutations and can help to improve our genomic annotation pipelines when RNA-sequencing and DNA-sequencing data is integrated to predict transcriptional consequences. Our findings also highlight the benefits of taking advantage of the selective mutagenesis that occurs in cancer genomes to evaluate splicing modulation across a given cohort. With our large and high confidence set of 2,888 SCMs we can effectively compare the landscape of rare and germline SCMs while grouping both sets together to evaluate overall trends in SCM+ exons. We were able to accurately determine that mutations overlapping the splice donor and splice acceptor splice site commonly undergo different selective pressures when mutated.

Specifically, mutations overlapping the splice donor site were sufficient to disrupt the canonical splice site usage but this phenomenon doesn't hold true for acceptor splice site mutations in our dataset. Alternatively acceptor SCMs needed to not only strengthen the novel splice site to facilitate the novel site usage, but also disrupt the canonical splice site. Furthermore, exons containing splice acceptor SCMs trended towards having a large upstream intron and downstream intron relative to other SCM+ exons. By surveying additional mutation induced alternative splicing events, we can continue to learn from the surrounding genomic context to tease apart the intricacies of the splicing code as it pertains to SCM+ exons. For example, we observed an enrichment of nucleosome occupancy overlapping SCM+ exons, suggesting active splicing of these exons across at

least the 33 tissue types surveyed in our analysis. Overall, the size of the novel exon tended to decrease post mutation, mimicking a natural evolutionary selective pressure but exploited in the cancer genome to maintain proper alternative splicing. With respect to degradation of the novel isoform, regardless of whether or not the resulting protein product was expected to undergo degradation, we still actively see expression of the novel isoform, although significantly lower for the transcripts predicted to elicit NMD.

To date, this is the first analysis comparing rare germline SCMs and somatic SCMs revealing their comparable dysregulation to the splicing code in cancer. Evaluating mutation induced events separately from patient specific *de novo* events can provide a focused analysis on the genomic features selecting for SCM+ exons relative to leaky splicing or mutation independent cryptic splice site activation. As tissue type specific datasets continue to increase, developing novel tissue specific signatures will help inform the tissue type specific relevance of mutation induced SCMs. For example

Understanding how splice-altering variants can lead to alternative isoforms in tumor samples and determining protein domains that are disrupted or created is still an open area of study. In order to better characterize events specific to a tissue type, developing modules of MiSplice that have known tissue specific transcripts will strengthen annotation and mutation calling. Furthermore, there is room to expand outside of the single nucleotide variant landscape to evaluate insertions, deletions and more complex events that are potentially also contributing to the disease state. Finally, MiSplice is a very stringent algorithm. There are many seemingly tissue type specific events that are present at a low level in some tissue types that have the potential to be exploited in other tissue types especially under direct mutagenesis. While we did not report on these cases in this analysis, the intermediate MiSplice outputs will provide very fruitful datasets for inferring novel biological mechanisms exploited in the cell.



Supplemental Figure S7. (A) Distribution of overall effect size of novel isoform broken down by conventional annotation and SCM type. (B) Comparing change in size of novel

spliced isoform to the distance from the canonical splice site. Both values were generated from different steps in the pipeline, further confirming our annotation pipeline. (C) Distribution of upstream intron sizes by conventional annotation. (D) Distribution of upstream intron sizes by conventional annotation.



Supplemental Figure S8. NMD Prediction Overview . (A) Comparing overall JAF distribution by conventional annotation type and colored by NMD prediction. (B) Based on the location of the premature termination codon, each predicted protein

product was classified as eliciting or escaping nonsense mediated decay based on the 50 bp rule.



Supplemental Figure S9. Nucleosome Occupancy by Canoncical Splice Site

Score. (A) Exons containing SCMs were grouped into bins by strength of canonical splice site and nucleosome occupancy was evaluated by group using NucMap. Low [-17.58,6.39], Medium [6.40,8.21], Higher[8.22,9.77], Highest [9.78,15.07].

4.6 References

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Chapter 5: Conclusion and Future

Directions

CANCER SPECIFIC EXON DEFINITION

From studying differential alternative splicing across species, it is understood the appearance of alternatively spliced exons has manifested through three different mechanisms: (1) the transition of a constitutive exon to an alternatively spliced exon, (2) exon shuffling and (3) exonization of intronic sequences.

When a new or duplicated exon is inserted into an existing gene it is known as exon shuffling. This process is facilitated by the repetitive elements in introns leading to recombination events and thereby exon shuffling. The model of exon creation through exonization was initially thought to occur from nothing. While many exon creation events are due to repetitive elements (Alu elements) which can undergo exonization, there are two well accepted methods of exonization including: intron-mutation induced splice signal creation events and RNA editing. Alu elements belong to the short interspersed element family (SINE), consist of a 300 nucleotide sequence, and are inserted into various regions of the genome via retrotransposition. The regions of an Alu element strongly resemble a canonical splice site in that they contain a poly-pyrimidine tract and splice-site like signals nearby that can be inaccurately identified by the splicing machinery. The use of a new

exon is further encouraged by strong alternatively spliced exons flanking the inserted sequence along with splicing enhancer and silencer elements. Interestingly, Alu exons have been found to be enriched for enhancer elements and depleted of silencer elements, encouraging the exonization event. In addition to the primary sequence, secondary structure can also affect exonization by means of adenosine deaminase acting on RNA (ADAR) editing. Adenosine to Inosine editing is common in Alu elements due to the secondary structure created when two inverted Alu elements are inserted near one another. During this process, the ADAR family of enzymes alters Adenosine residues to Inosine. Since Inosine is recognized by the biological machinery as Guanosine, a functional splice site can be created in an Alu element by altering AA to AG. There have been several cases of exon gain due to Alu insertions in exons (Sela et al.) and introns (unpublished).

Evolutionary conservation or depletion of a splice site provides strong evidence as to the biological function of the alternatively spliced product.

Understanding how exon creation or modification can generate a novel protein isoform is vital to understanding the evolution of canonical gene function. New exon events that arise in cancer or in natural selection in various species allows for the new exon event to be "tested", since the new event is almost always alternatively spliced the wildtype form is also present in the organism as well (Sorek, 2007). If the alternative isoform is detrimental it will be selected against, and if beneficial we should see evidence of selection in the form of expression of the alternatively spliced product. It has been

suggested that exonization events are ways to test out new gene modules with a generally small effect on the overall evolution of the species.

The findings from my thesis work suggests that evolutionary constraints on the tumor has selected for these splice-site-creating mutants and conferred some combination of the following criteria: (1) modify protein structure and/or (2) shift mRNA usage away from the wild type.

Although many of the predicted splice-site-creating events are predicted to escape nonsense mediated decay, about half of the sites should be degraded but are still present in the total RNA population, signifying a novel means of escaping degradation. In this thesis we propose an additional classification of "exonization" through the lens of cancer evolution. We have identified thousands of somatic and germline coding mutations, inducing the use of a novel or cryptic splice site functionally altering the reading frame of the mutant isoform and diversifying the landscape of the cancer genome. In the following sections, I will discuss additional analyses that can be performed on this compendium of splice-site-creating mutants to evaluate selection of SCMs across the TCGA dataset.

Allelic Imbalance of SCMs

Over evolutionary time during the birth of a new exon, while undergoing various selective pressures, newly created exons tend to exhibit increased expression and eventually become fixed over the previous wild-type form (Sorek, 2007). Similarly with our splice-creating mutants, we see varying levels of selection by the tumor, and in some cases the

mutant induced splice form is selected at a higher frequency than the wild type suggesting preference for the alternatively spliced product (Sorek, 2007).

In thinking about cancer from an evolutionary standpoint, we have a unique opportunity to evaluate changes to overall exon definition and determine their potential for purifying selection over a relatively short time scale. When interrogating mutational signatures in cancer, we predict genomic changes will lead to changing expression of the resulting gene product. The clonal heterogeneity of a tumor exemplifies the theory of selective pressures on growing populations of cells in a microenvironment. Selective pressures (including treatment, microenviroment, tissue etc.) alter the clonal architecture and the clone with the most beneficial fitness is maintained until treated (Gerlinger et al.). By looking at the overall variant allele fraction of the DNA and RNA, we can evaluate the transcriptional dynamics that could be exploited by the cancer cell. When exons are newly created, some might fix over time and we can evaluate this by comparing expression of the newly created exons relative to the wildtype sequence (Sorek, 2007). When comparing the different Alu element insertions between mouse and humans, Sela et al., found increased levels of the newly created exons in humans relative to mouse (Sela et al.) suggesting selection for the Alu created exons over evolutionary time.

For approximately 80 rare germline splice-site-creating events in our dataset, we have access to DNA-Sequencing and RNA-Sequencing for both the tumor and normal samples. With these data we can assess the comparative Variant Allele Fractions (VAF) for DNA-seq and Junction Allele Fraction (JAF) in the RNA-Seq between the tumor and

normal data. Understanding allelic imbalance will aid us in understanding the difference in expression between the mutant and wild type allele, a selection of one allele over another in the tumor would suggest either cis or trans factors that affect expression of the associated gene thereby altering the tumor biology.

Protein Support from Mass Spectrometry

Support of the novel splice-site-creating event could be significantly strengthened by peptide support. The Clinical Proteomic Tumor Analysis Consortium (CPTAC) has performed mass spectrometry for a subset of samples from the cancer genome atlas project to further interrogate how mutations can manifest on the translational level. To evaluate the translational potential of the splice-site-creating mutants, for a handful of samples in our dataset we have mass spectrometry data to evaluate if the novel isoform is translated (Table 5.1). In Table 5.1 I have highlighted all the sample and gene pairs for which we have associated mass spectrometry data and can validate the novel splice isoform if it is expressed. Specifically, MSGF+ (https://omics.pnl.gov/software/ms-gf) can be utilized to identify peptides specific to our novel splice isoforms. Although we have evidence of the mutant isoforms in the RNA-Sequencing data, support by mass spec will provide very strong evidence that the mRNA is translated to a protein whose functions could be altered due to the alternatively spliced product.

Table 5.1. rgSCM and sSCM events with mass spectrometry data

Туре	Result	Frame	NMD	Gene	Canc	JAF	Conventional
SCM			Classificati		er		Annotation
			on				
germline	exon-	off-frame	ELICIT-	CBWD5	BRC	8.62069	
	extension		NMD		А		SpliceDonorSN
							V
germline	exon-	off-frame	ESCAPE-	C7	OV	39.2461	
	extension		NMD				SpliceDonorSN
							V
germline	exon-	in-frame	ESCAPE-	SORBS3	BRC	9.52381	Synonymous
	shrinkage		NMD		А		
germline	exon-	in-frame	ELICIT-	TUBGCP	BRC	41.3793	
	shrinkage		NMD	6	А		SpliceAcceptor
							SNV
germline	exon-	off-frame	ELICIT-	CCDC94	OV	9.40171	
	shrinkage		NMD				SpliceAcceptor
							SNV
germline	exon-	off-frame	ELICIT-	MTBP	BRC	22.2222	Missense
	shrinkage		NMD		А		
germline	exon-	in-frame	ESCAPE-	SKA2	OV	21.4286	Missense
	shrinkage		NMD				
germline	exon-	off-frame	ESCAPE-	LMO1	OV	18.75	Synonymous
	shrinkage		NMD				
germline	exon-	in-frame	ESCAPE-	ZCCHC1	BRC	14.8148	Missense
	shrinkage		NMD	7	А		

germline	exon-	in-frame	ESCAPE-	PFKL	BRC	45.1128	
	shrinkage		NMD		А		SpliceAcceptor
							SNV
germline	exon-	in-frame	ESCAPE-	PIAS3	OV	34.9593	Missense
	shrinkage		NMD				
germline	exon-	off-frame	ELICIT-	ELMO3	OV	6.76692	Missense
	shrinkage		NMD				
germline	exon-	off-frame	ESCAPE-	HMGCR	BRC	8.94309	
	shrinkage		NMD		А		SpliceAcceptor
							SNV
germline	exon-	in-frame	ESCAPE-	CHD8	OV	27.3092	Missense
	shrinkage		NMD				
germline	exon-	off-frame	ELICIT-	NEDD1	BRC	5.9322	Missense
	shrinkage		NMD		А		
germline	exon-	off-frame	ESCAPE-	CUEDC1	BRC	36.4985	
	shrinkage		NMD		А		SpliceDonorSN
							V
germline	exon-	in-frame	ESCAPE-	PTPRM	OV	49.1228	
	shrinkage		NMD				SpliceAcceptor
							SNV
somatic	exon-	off-frame	ELICIT-	TP53	OV	46.6667	
	extension		NMD				SpliceDonorSN
							V
somatic	exon-	off-frame	ESCAPE-	ZDHHC1	OV	30.2013	Missense
	extension		NMD	6			

somatic	exon-	off-frame	ESCAPE-	SNAPC1	BRC	21.1538	
	shrinkage		NMD		А		SpliceAcceptor
							SNV
somatic	exon-	in-frame	ESCAPE-	CDC37	OV	16.7965	Synonymous
	shrinkage		NMD				
somatic	exon-	in-frame	ESCAPE-	NFYB	OV	19.5402	
	shrinkage		NMD				SpliceAcceptor
							SNV
somatic	exon-	off-frame	ELICIT-	LANCL2	OV	22.7273	
	shrinkage		NMD				SpliceAcceptor
							SNV
somatic	exon-	in-frame	ESCAPE-	NOP9	BRC	36.3636	Missense
	shrinkage		NMD		А		
somatic	exon-	off-frame	ELICIT-	EDC3	OV	7.5188	
	shrinkage		NMD				SpliceAcceptor
							SNV
somatic	exon-	off-frame	ESCAPE-	C1orf172	BRC	12.8205	Missense
	shrinkage		NMD		А		
somatic	exon-	in-frame	ESCAPE-	GTF2E2	BRC	12.8713	
	shrinkage		NMD		А		SpliceAcceptor
							SNV
somatic	exon-	off-frame	ESCAPE-	ID3	OV	23.3051	
	shrinkage		NMD				SpliceAcceptor
							SNV
somatic	exon-	in-frame	ESCAPE-	EIF1AX	OV	39.0476	
	shrinkage		NMD				SpliceAcceptor
							SNV

somatic	exon-	in-frame	ESCAPE-	FBXO18	OV	8.31889	Missense
	shrinkage		NMD				
somatic	exon-	in-frame	ESCAPE-	CRAMP1	BRC	39.2857	Missense
	shrinkage		NMD	L	А		
somatic	exon-	off-frame	ELICIT-	CDK12	OV	40.7407	
	shrinkage		NMD				SpliceAcceptor
							SNV

Oncogenic Function of SCMs

MiSplice identified two kidney renal clear cell carcinoma (KIRC) samples having the same conventionally annotated missense mutation (c.233A>G, p.N78S) in BAP1, a nuclear deubiquitinase, that created the same novel spliced-out alternative splicing product. Inactivation of BAP1 is prevalent among renal cell carcinomas (Wadt et al., 2012) and an annotated missense mutation (p.L570V) has been reported to create a cryptic splice site in melanoma. At the transcriptional level, the expression of the case and control samples are relatively comparable, but at the translational level, one case with available protein data (RPPA) showed significantly lower expression relative to the controls. This result suggests the conventionally annotated missense mutations in BAP1 likely create an alternatively spliced transcript that is not readily expressed at the protein level. With this evidence it would be interesting to determine the in vitro function of BAP1 splice-creating variant in a cell line of interest. Using the established cell lines with our splice-site-creating variants of interest, downstream assays can be performed to determine if the SCMs have the potential to contribute to oncogenesis.

Mutation Induced Ultra Short Introns

With the birth of a new intron, we see cases of intronization or the alternative splicing of an internal exon (Irimia et al., 2008). With certain events we also see the alternative splicing of internal introns that look to be facilitated by selected mutation induced splicing events. Small intron-creation events in exons were observed in approximately ~150 unique gene-cancer type events. While introns are commonly spliced out by the major and minor spliceosome, several alternative mechanisms can facilitate the splicing of very short introns, typically less than 65 nucleotides in length. For example a U2 snRNP SF3b excises short introns ranging from 43 to 56 bp in NDOR1, HNRNPH1 and ESRP2 (Sasaki-Haraguchi et al.). While introns in humans are quite long, introns in invertebrates are much shorter. With this knowledge, Lim et al. assessed genomic features of short introns across organisms to determine the minimum amount of information required to define an intron (Lim and Burge). While there is ample evidence in the literature supporting ultra short introns (USI), genomic deletions can arise due to repetitive sequences. To filter out common genomic deletions, we overlapped our predicted USI's with computationally predicted USI's in a recent publication by Abebrese et al (Abebrese et al.). In total, 29 USI's were filtered leaving less than 20 putative mutation induced USI's. Furthermore, 3 USI's in KMT2D, CBLN3 and PRRC2C overlapped with Abebrese et al. and PRRC2C had evidence of splice junction usage across several cell lines after screening A549, Bj, H1 hESC, HeLa, HepG2, Hsmm, K562, Mcf7, Nhek, and Nhlf.

To date, mutation induced USI's have not been systematically evaluated in cancer. In general for all the USI's identified, all but one are in frame and generally do not change

the splice site score but are recurrent and span cancer types. While the splicing score doesn't change, it has been reported that short introns generally have non-canonical splice sites such as those observed in IRE1alpha-dependent Xbp1 mRNA splicing known for splicing out 26 nucleotide sequences in various genes including itself (Bai et al.). The predicted mutation induced USIs identified in the TCGA dataset provides additional evidence regarding the mechanisms by which the splicing code can be dysregulated in the human population and potentially exploited in cancer.

Common Germline Splice-Site-Creating Variants

As a proof concept, we wanted to explore common germline splice-site-creating events in one cancer type to determine ancestry specific events. For the 20,205,168 common variants in Breast Invasive Carcinoma (BRCA), in an initial screen only 72,245 unique variants were evaluated for splice-site-creating function. Of the 72,245 unique variants, 287 had evidence of splice-creating function in at least one sample. The variants with splice-site-creating function were then evaluated in the remaining samples. In total 74,383 variants derived from 287 unique sites were evaluated for splice-site-creating function. 29 variants with less than 10% of control samples exhibiting the alternatively spliced isoform were manually reviewed, resulting in 6 high confidence common germline splice-sitecreating variants (Table 5.2).

Table 5.2. Common SCMs

Gene	Mutation	#	% of	# reads in case samples
		Cases	Controls	
			with event	
Neurolysin,	5:6508410	22	0 %	15,14,18,11,8,7,12,13,5,7,11,
mitochondrial	1 A>G			13,6,5,10,13,6,7,18,27,9,10
(NLN)				
Leukocyte	19:550986	20	0.39 %	11,5,7,5,15,6,7,7,5,8,13,10,15
Immunoglobulin	67 G>A			,5,17,9,8,10,9,10
Like Receptor				
A2 (LILRA2)				
Glutaminyl-	19:462062	4	1.07 %	7,5,5,7
Peptide	62 G>A			
Cyclotransferase				
Like (QPCTL)				
TatD DNase	1:2129855	17	2.25 %	6,6,6,6,6,5,8,6,6,7,5,8,5,8,7,6,
Domain	92 G>A			12
Containing 3				
(TATDN3)				
Calpain 13	2:3098597	4	2.64 %	9,11,8,16
(CAPN13)	7 G>A			

NAD kinase 2,	5:3621971	22	6.01 %	13,6,18,9,8,5,17,8,25,18,11,7,
mitochondrial	0 C>T			8,19,12,35,17,34,16,6,6,7
(NADK2)				

The NLN common germline variant has a population frequency of 0.19 in the african american community. 20 of the 22 samples with the germline variant are black or african american in the TCGA dataset and have a slightly higher expression of the NLN gene relative to non-mutated controls in breast cancer, although not significant (wilcox test = 0.09). Our initial findings of the NLN gene and other common variants suggest a subset could be mis-annotated when evaluating both RNA and DNA sequencing. Interestingly a few variants created ultra-short-intron events described in the previous section.

Non-Canonical Splice Sites and Minor Spliceosome Alterations

For both splice-site-creating papers, we only focused on sites creating canonical GT or AG splice sites. Several interesting SCMs created non-canonical splice sites (deviating from GT and AG genomic context) suggesting usage of non-canonical splice sites by the major spliceosome or minor spliceosome.

The splicing code is made up of cis-acting elements that help the splicing complex distinguish between non-coding (intron) and coding (exon) regions. There are two known distinct ribonucleoprotein complexes known as the major and minor spliceosomes that are responsible for joining exons and splicing out two types of introns, U2 and U12. For

U2 type introns, the consensus intronic dinucleotide GT splice donor and AG splice acceptor flank the exons at the 3' and 5' ends respectively, and can be found in 99% of all introns. The remaining 1% are U12 type introns and were initially recognized due to their non-consensus splice site sequences AT and AC at the 3' and 5' splice sites, respectively (Hall and Padgett; Jackson). Interestingly, some introns with the canonical GT-AG splice sites are also spliced by the minor spliceosome. While the consensus sequence alone cannot differentiate between the two groups, several other cissequences including lacking a polyprimidine tract upstream of the 3' splice site, polypyrimidine tract and branch point (Dietrich et al.) can help differentiate between the two groups. The major and minor spliceosome share the U5 snRNP and many spliceosomal proteins.

While only 1% of U12 type introns are present in the genome, they are highly conserved across eukaryotes suggesting a potential functional role in the genome. For example, U12 type introns are shown to be removed at a much slower rate than U2 type introns (Patel et al.), and the minor spliceosomal proteins are present at a much lower abundance than the major spliceosome (Tarn and Steitz). This evidence suggests that minor introns are likely a rate limiting step in the expression of various mRNAs that contain U12 type introns. Despite this evidence, the current compendium of minor introns is still expanding.

The benefit of coupling U12 intron containing genes and U12 spliceosome abundance allows the cell a regulatory mechanism by which only a subset of genes can be differentially regulated in response to stimuli. Younis et al., determined the U6atac snRNA minor spliceosomal gene expression could be increased by cell stress activated kinases leading to increased expression of U12 intron containing genes that regulate cell stress physiology (Younis et al.). RNA-seq analysis revealed a down-regulation of 2,088 genes including 429 minor intron genes. Together, these results show that altering expression of minor spliceosomal genes can alter a distinct subset of genes with overarching changes at the transcriptome level.

With the large dataset provided by The Cancer Genome Atlas mechanisms of U12 dysregulation across cancer genomes can be more effectively evaluated. Furthermore, our analysis of splice-site-creating mutants identified the usage of many non-canonical GT-AG splice sites, suggesting a potential dysregulation of spliceosomal usage.

As with aberrations in major spliceosome related genes, mutations in minor spliceosomal genes lead to different disease phenotypes ranging from brain to skeletal irregularities. Recently, Madan et al., evaluated alternatively spliced junctions in 8 Myelodysplastic syndrome (MDS) patients with *ZRSR2* mutations and identified 689 mis-spliced junctions in all 8 cases (Madan et al.). The mispliced junctions are specific to *ZRSR2* mutants. Additionally, a conditional knockdown of *Rnu11* in mice resulted in microcephaly and upregulated intron retention of minor intron containing genes (Baumgartner et al.). Many of the minor intron containing genes were not differentially expressed but 178 introns were detained at higher levels in the mutant relative to the wildtype while one was downregulated. Finally a family of three sisters were found to have biallelic mutations in *RNPC3* and when downstream U12 intron retention rates were compared, 21 genes out

of 522 tested had decreased U12/U2 ratios in the patient cells(Argente et al.). Interestingly the subset of aberrant U12 introns were relevant to the disease phenotype and some U2 cryptic splice sites were activated instead of the U12 intron. Mutations in U12 spliceosomal proteins have been linked to several other diseases including RNU12 (Elsaid et al.), RNU4ATAC (Edery et al.; He et al.; Heremans et al.; Merico et al.), TRAPPC2 (Shaw et al.), FUS (Reber et al.), and SMN1 (Zhang et al.). The previous studies provide strong evidence that minor splicing factors regulate subsets of minor introns. By evaluating differential expression of related minor introns, novel functional mutations in minor spliceosome genes or U12 minor introns themselves can be identified to determine if minor intron splicing is altered globally in cancer.

As with mutations in spliceosomal genes, mutations in cis can also alter minor intron splicing. *LKB1* encodes a serine threonine protein kinase involved in major cellular processes including cell cycle arrest, p53 mediated apoptosis among others. The second intron of *LKB1* is a minor intron and a mutation that affects the 5' splice site causes Peutz-Jeghers syndrome (Hastings et al.). Interestingly, the 5' splice site mutation changes the splice sites of intron 2 from AT-AC to GT-AC. One would expect that a 5' splice site mutation would cause skipping of the subsequent exon or use of an alternative 5' splice site, but the authors found instead that a cryptic 3' splice site was utilized, thereby altering the frame of the final transcript, which induced a premature termination codon and degraded by nonsense mediated decay.

In a related manner, a mutation in *SCN8A* has been shown to inactivate the 5' splice site thereby resulting in an exon skipping event associated with an X-linked recessive disorder (Shaw et al.). While this exon skipping event doesn't occur in the minor intron itself, it does lead to altering the usage of minor intron contained within the gene, suggesting a strong interplay between splice site usage.

In 2018, our lab published a comprehensive analysis of gene fusions identified in RNA-Seq across the TCGA compendium (Gao et al.). From this publication, several gene fusions were identified involving minor spliceosomal related genes (Table 5.3). All of these resulting fusion products in the associated samples have the potential to dysregulate a large subset of downstream minor introns, but their current consequence is unknown. A case control comparison evaluating overall expression of the fusion genes with wild type counterparts will determine if the expression of the fusion itself or downstream targets are affected due to the predicted fusion product.

Cance	Sample	Fusion	Juncti	Spanni	Breakpoin	Breakpoint2
r			on	ng	t1	
HNSC	TCGA-CV-7090-01A-	DHX15	6	21	chr4:24548	chr4:2638635
	11R-2016-07	RBPJ			855:-	3:+
PRAD	TCGA-G9-6339-01A-	DHX15ETV1	9	35	chr4:24584	chr7:1393589
	12R-A311-07				323:-	6:-
SARC	TCGA-DX-A3LU-01A-	DHX15	1	12	chr4:24541	chr6:4194058
	11R-A21T-07	CCND3			873:-	5:-

Table 5.3. U12 and U12 associated genes with detected Fusions in TCGA
COAD	TCGA-F4-6809-01A-	FUS	2	116	chr16:3118	chr16:312022
	11R-1839-07	PYCARD			2422:+	03:-
ESCA	TCGA-IG-A51D-01A-	FUSIL9R	6	7	chr16:3118	chrY:5718942
	11R-A36D-31				2664:+	6:+
HNSC	TCGA-CV-6941-01A-	FUS	1	11	chr16:3118	chr16:311497
	11R-1915-07	PRSS36			4396:+	31:-
OV	TCGA-13-1481-01A-	FUSKAT8	22	22	chr16:3118	chr16:311270
	01R-1565-13				0227:+	35:+
STAD	TCGA-HU-A4HB-01A-	FUS	71	35	chr16:3118	chr16:857222
	12R-A251-31	TMEM114			5179:+	4:-
BRCA	TCGA-LD-A7W5-01A-	PDCD7	3	16	chr15:6512	chr17:358177
	22R-A352-07	TAF15			9032:-	16:+
BRCA	TCGA-E2-A14Z-01A-	RNPC3	3	5	chr1:10352	chr5:9441738
	11R-A115-07	KIAA0825			6262:+	0:-
BRCA	TCGA-E2-A10A-01A-	SNRNP48	2	1	chr6:75951	chr3:5248451
	21R-A115-07	NISCH			01:+	3:+
SARC	TCGA-IW-A3M4-01A-	ZCRB1	1	8	chr12:4232	chr12:422106
	11R-A21T-07	YAF2			5924:-	19:-
SKCM	TCGA-D9-A6E9-06A-	ZCRB1	18	64	chr12:4232	chr12:502224
	12R-A311-07	LIMA1			4019:-	85:-
BLCA	TCGA-G2-AA3B-01A-	ZMAT5	4	15	chr22:2976	chr22:297936
	11R-A39I-07	ASCC2			6949:-	76:-
GBM	TCGA-41-2572-01A-	ZMAT5	110	32	chr22:2976	chr22:297905
	01R-1850-01	ASCC2			6872:-	48:-
HNSC	TCGA-T3-A92M-01A-	ZMAT5	5	11	chr22:2976	chr22:295618
	31R-A39I-07	NIPSNAP1			6872:-	62:-

LUAD	TCGA-05-4426-01A-	ZMAT5	1	3	chr22:2976	chr22:250144
	01R-1206-07	KIAA1671			6949:-	30:+
LUSC	TCGA-85-8048-01A-	ZMAT5HIRA	2	7	chr22:2973	chr22:193315
	11R-2247-07				8330:-	56:-
SARC	TCGA-Z4-A9VC-01A-	DDIT3FUS	3	23	chr12:5752	chr16:311896
	11R-A37L-07				0418:-	65:+
BRCA	TCGA-E2-A1IO-01A-	CLPX	1000	1000	chr15:6516	chr15:651331
	11R-A144-07	PDCD7			6642:-	83:-
LUAD	TCGA-97-A4M7-01A-	MPG	32	116	chr16:8325	chr16:57086:+
	11R-A24X-07	SNRNP25			6:+	
LUSC	TCGA-21-1070-01A-	ARHGAP17	1	7	chr16:2496	chr16:56539:+
	01R-0692-07	SNRNP25			4197:-	
SKCM	TCGA-D3-A8GC-06A-	FAM129B	6	35	chr9:12751	chr16:55777:+
	11R-A37K-07	SNRNP25			6857:-	
SKCM	TCGA-EB-A44Q-06A-	KDM2B	5	24	chr12:1215	chr12:123458
	11R-A266-07	SNRNP35			32806:-	021:+
BRCA	TCGA-E2-A14T-01A-	TMEM117	18	26	chr12:4384	chr12:423241
	11R-A115-07	ZCRB1			4928:+	04:-
STAD	TCGA-BR-8077-01A-	YAF2	9	8	chr12:4221	chr12:423139
	11R-2343-13	ZCRB1			0355:-	86:-
OV	TCGA-24-1425-01A-	ST6GALNAC	7	8	chr17:7664	chr22:297424
	02R-1566-13	1ZMAT5			3508:-	80:-
PAAD	TCGA-3A-A9IH-01A-	NF2ZMAT5	3	5	chr22:2967	chr22:297424
	12R-A39D-07				8323:+	80:-

In evaluating disruptions of minor introns, several experiments can validate the *in silico* predictions. If the minor intron is retained due to mutation or disruption of U12 or U12-

associated splicing factor we would expect to see either expression of the retained intron in RNA-Seq or decreased expression of the associated gene. If the minor intron has an associated premature termination codon, one would predict the resulting transcript would be degraded by nonsense mediated decay. If effectively degraded, no reads supporting the transcript would be identified in the RNA-Seq. To confirm this, the transcript containing the mutation can be transfected into the cell line of interest and puromycin treated to inactivate the NMD pathway thereby allowing us to evaluate the full repertoire of transcripts produced by the mutant transcript. If the transcript is alternatively spliced due to the mutation in the minor intron and is normally degraded by NMD, in this experiment we should see evidence of the minor intron retained due to the knockdown of NMD factors by puromycin. Alternatively, if we do not see the transcript even after NMD knockdown, then the mutation in the minor intron is likely not disrupting this particular intron retention event.

If the minor intron is retained as suggested by evidence in the RNA-Sequencing data and it isn't degraded efficiently, we can next try and determine if the major and minor spilceosomal machinery is responsible for removal of the minor intron of interest. Since the U2 spliceosome is known to be degenerate and can identify multiple sequences, it is possible that the U12 spliceosome isn't playing a role in removing the minor intron of interest. To test this experimentally(Hastings et al.), a mini-gene derived with our exon, intron and exon of interest can be can be incubated in nuclear extract. When incubated in the presence of ATP, we can evaluate mRNA products that are derived from ATP dependent splicing reactions. Then in this same experimental condition, U2 and U12 dependent spliceosomes can be inactivated using antisense oligonucleotides directed at the U2 small nuclear RNA, U12 snRNA or both. Finally, RNA is extracted from all conditions and alternatively spliced products can be evaluated via RT-PCR. Outcome 1: If U12 spliceosome is important for this particular minor intron splicing then you will see a decrease or absence in the spliced product when associated spliceosomal RNA is knocked down. And when U2 spliceosome is knocked down there should be no change in spliced product. Outcome 2: If the U2 spliceosome is important for the splicing of this transcript then you will see a decrease or absence in the spliced product when associated spliceosomal RNA is knocked down. And when U12 is knocked down there should be no change in spliced product. Outcome 3: Both spliceosomes may be important for this process thereby resulting in differences in the spliced products under both knockdown conditions.

5.1 References

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170

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171

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172

Appendix: Additional Projects

5.1 Complex Insertions and Deletions in Cancer Genomes

Systematic Discovery of Complex Indels in Human Cancers (Nature Medicine, 2016)

(Ye et al., 2016)

Contribution: I lead the development and scripted several scripts that were used to develop a variant quality control pipeline for filtering complex insertions and deletions. I manually reviewed many complex indels and helped with writing and submission. I helped to validate complex insertions and deletions identified in a COLO829 cell line by sanger sequencing.

Refer to Supplementary Note 2: Sanger Sequencing of Validated Complex Indels from COLO829.

5.2 Analysis of Somatic Complex Insertions and Deletions in Pediatric Hematopoietic Malignancies

Identification of Complex Indels in Pediatric Cancer Genomes

Complex indels are formed by deleting and inserting DNA fragments of different sizes at a common genomic location. Application of the publicly available Pindel-C suite (https://github.com/genome/pindel) to 545 whole genome Pediatric Cancer Genome Project (PCGP) samples uncovered 21 somatic complex insertions and deletions among 20 pediatric hematopoietic cancer samples listed in Table below. Our analysis was restricted to the following genes commonly mutated in hematopoietic diseases including: *TP53, CBL, CREBBP, FLT3, KMT2D (MLL2), PAX5, SETD2, IKZF1, PMS2, RAD51, NCOR1, TBL1XR1 and KMT2C (MLL3).* Nine of the 21 complex insertions and deletions identified differed from those reported in the original studies as follows:

- 2 complex events in TBL1XR1 and SETD2 were not reported;
- 6 complex events in *MLL2*, *SETD2*, *IKZF1*, and *CREBBP* were classified as simple indels;

• 1 complex event in *TP53* was classified as both a SNV and an INDEL separately. The remaining 12 events occurring among several key genes are found among samples that are currently not publicly available. These samples are among the following cancer types: Hyperdiploid Acute Lymphoblastic Leukemia (SJHYPER), E2A-PBX Acute Lymphoblastic Leukemia (SJE2A), and ETS-Related Gene Associated Acute Lymphoblastic Leukemia (SJERG). A handful of somatic and germline complex indels have also been identified in non-coding regions of the genome, but our initial analysis focuses on coding somatic mutations in cancer genes.

This preliminary analysis demonstrates the importance of using the most updated form of Pindel-C to identify complex events that may be absent or classified as other variant types in current cancer genomics studies.

Sample	Gene	Chromosome	Start	End
SJE2A043	PAX5	9	37002647	37002649
SJERG031	MLL2	12	49444452	49444454
SJERG020052	MLL3	7	151842335	151842365
SJERG020306	SETD2	3	47059133	47059166
SJERG020307	TBL1XR1	3	176756153	176756179
SJERG020309	MLL2	12	49444958	49444960
SJETV001	TBL1XR1	3	176769294	176769299
SJETV024	SETD2	3	47059194	47059204
SJHYPER005	NCOR1	17	15952238	15952243
SJHYPER007	FLT3	13	28608280	28608286

Table A.1. Complex indels identified for PCGP Samples

SJHYPER051	TP53	17	7577595	7577601
SJHYPER097	SETD2	3	47162194	47162195
SJHYPER104	PAX5	9	37002675	37002676
SJHYPER108	MLL2	12	49443861	49443863
SJHYPER111	CBL	11	119148875	119148900
SJHYPER111	FLT3	13	28608305	28608309
SJHYPER227	IKZF1	7	50367310	50367314
SJHYPO002	CREBBP	16	3843416	3843439
SJHYPO040	CREBBP	16	3799606	3799632
SJINF013	TP53	17	7577093	7577111
SJPHALL020043	SETD2	3	47147487	47147488

Complex indel discovery

- 1. Read Extraction
- 2. Pattern Growth-Based Alignment
- 3. Distinguishing Complex Indels from Simple Indels
- 4. Remove False Predictions and Variant Allele Frequency Analysis

Filtering procedure

- 1. Coverage: All sites with at least 20 reads supporting the site of interest in both the tumor and normal sample were maintained for further analysis.
- Coding Region Selection: Identified all complex indels overlapping an exon as defined by ensembl 75 including flanking 2 bp splice sites.
- Repetitive Regions: Removed any variants falling in repetitive regions as defined by MSI Sensor. Reptitive regions are defined as areas having more than 6 or more repeat unit bases in a segment of the genome.
- 4. Manual Review: Finally all sites were manually reviewed in the integrative genomics viewer to finally confirm all complex indels.

The screenshots below from the Integrative Genomics Viewer show the events described in the table above (tumor sample on top, normal sample on bottom).

Sample	Gene	Chromosome	Start	End
SJE2A043	PAX5	9	37002647	37002649



Sample	Gene	Chromosome	Start	End
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Sample	Gene	Chromosome	Start	End
SJERG020052	MLL3	7	151842335	151842365



Sample	Gene	Chromosome	Start	End
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Sample	Gene	Chromosome	Start	End
SJERG020307	TBL1XR1	3	176756153	176756179



Sample	Gene	Chromosome	Start	End





Sample	Gene	Chromosome	Start	End
SJETV024	SETD2	3	47059194	47059204

	chr3 p26.2 p25.3	p24.3 p24.2 p23	p22.2 p21.33 p21.2 p14	3 p14.1	p13 p12.3 p12.1	q11.1 q12.1 q13.11	q13.2 q13.32 q21.2	q22.1 q22.3	q24 q25.1 q25.31	q26.1 q26.2	q26.32 q27.1 q28 q29
	47,059,194 bp	1	47,059,196 bp	1	47.059.198 bp	11 bp	47,059,200 bp		47.059.202 bp	1	47,059,204 bp I
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Sample	Gene	Chromosome	Start	End

	SJHYPER005	NCOR1	17	15952238	15952243	
	chr17 p13.3 p13.2 p13.1 p12	p11.2 p11.1	q11.2 q12 q21.1 q21	.31 q21.32 q21.33 q22	q23.1 q23.3 q24.2 q2	4.3 q25.1 q25.2 q25.3
	≠ 39 1	15.952,230 bp			15,952,250 bp	15,952,260 bp
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134044309.bam						
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135582585.bam Junctions						
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135582585.bam				-		
Sequence RefSeq Genes		GAACCT	G G G G T G G G	GAGATG	G G C T C G T P E Y	A G G G C T P E
			NCOR1			

Sample	Gene	Chromosome	Start	End
SJHYPER007	FLT3	13	28608280	28608286



Sample	Gene	Chromosome	Start	End
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Sample	Gene	Chromosome	Start	End
SJHYPER108	MLL2	12	49443861	49443863



Sample	Gene	Chromosome	Start	End



Sample	Gene	Chromosome	Start	End
SJHYPER111	FLT3	13	28608305	28608309



Sample	Gene	Chromosome	Start	End


Sample	Gene	Chromosome	Start	End
SJHYPO002	CREBBP	16	3843416	3843439



Sample	Gene	Chromosome	Start	End







5.3 Children's Discovery Institute: Utilizing CharGer to determine novel variants in undiagnosed pediatric cases

There is an urgent need for improved and quick diagnosis of birth defects. To address this concern our lab has developed CharGer (unpublished) to assess the pathogenicity of variants using and automated query of public data sources (example: Clinvar). Using patient mutation files (example: VCFs) derived from the mother, father and child, we can derive both de novo and compound heterozygous mutations.

When focusing on de novo variants we are interested in identifying spontaneous mutations in developmental genes compared to inherited mutations that are derived from both parents. We also require that mutations have a low frequency in the population (PM2). In this partial blind study we ran mutation files through the CharGer pipeline without knowing the associated phenotype. Variants of interest reported through the pipeline were reported to a physician at Washington University to determine the overlap between mutations called by an orthogonal gene sequencing service, GeneDx, and those that were missed.

To validate our automatic classification, we applied CharGer on a data set of 7 families with various birth defects in infants and children. For 6 families, CharGer confirmed a predicted "likely pathogenic" variant reported from GeneDx or followup functional studies from collaborators. Using variant call format (VCF) from GeneDx we derived mutations inherited from the mother, father and de novo variants in the child. Variants were filtered by high population allele frequency (>0.05) derived from 1000 genomes (1KG), Exome Aggregation Consortium (ExAC) and Exome Sequencing Project (ESP) and reported variants in our predefined gene list made up of 597 developmental genes (Saunders et al.,), 625 cancer genes and 11 additional genes identified in GeneDx positive analyses. CharGer identified 3/6 families as having 'Likely Pathogenic 'candidate variants and the remaining 3/6 as having 'Unknown Significance'. Furthermore, CharGer identified additional candidate variants for all 6 families, along with candidate variants for the 7th GeneDx negative family.

5.4 Discovery of Novel Fusions in Cancer Genomes

Contribution: I helped oversee the overall development of this project including figure development and writing.

Cell Rep. 2018 Apr 3;23(1):227-238.e3. doi: 10.1016/j.celrep.2018.03.050.

Driver Fusions and Their Implications in the Development and Treatment of Human Cancers {(Gao et al., 2018)}.

5.5 Evaluating Novel Germline Variants in Cancer Genomes

Contribution: I helped to manually review many germline variants and helped significantly with the BRCA1 homologous recombination assay for the below publication.

Patterns and functional implications of rare germline variants across 12 cancer types (Lu et al., 2015).

5.6 Effect of Blood Somatic Mutations in PPM1D on TP53 and Cell Cycle Arrest

In 2014, our lab was one of the first to identify blood specific somatic mutations in elderly patients without any adverse hematopoietic diseases (Xie et al., 2014). After, performing a similar analysis on additional samples, we identified one novel recurrently mutated gene with a significant enrichment of blood-somatic mutations. PPM1D is a phosphatase involved in the dephosphorylation of DNA damage response genes including TP53. While this gene has been associated with breast and ovarian cancer predisposition, it has still not yet been reported in hematological malignancies. C-terminal truncating mutations are predicted to hinder the activation of DNA damage response proteins by enhancing the dephosphorylation activity of *PPM1D*. Figure X highlights blood specific mutations identified across 2,278 patients.



Figure A.1: Blood specific somatic mutations identified in *PPM1D*

We hypothesized blood specific mutations in *PPM1D* disrupt *TP53* phosphorylation. To evaluate this hypothesis, we determined the level of p53 Ser15 phosphorylation of PPM1D Mutants after radiation treatment (Methods). Under conditions of DNA damage (ex. Radiation) TP53 is phosphorylated and will activate downstream genes to initiate the DNA damage response pathway. Normally PPM1D will dephosphorylate TP53, thereby inactivating the DNA damage response pathway. Without UV treatment you would expect to see low levels of TP53 phosphorylation. As shown previously in our lab, PPM1D mutants further decreased Ser15 phosphorylation levels (stronger than WT), suggesting PPM1D mutants are more "active" than the wild type. Under wild type conditions, we expect to see a larger number of cells undergoing apoptosis after UV treatment. Under normal conditions TP53 should be phosphorylated and activated thereby halting the cell cycle process in response to DNA damage. To evaluate this, we can use propidium iodide (PI) staining to evaluate cells in various stages of the cell cycle. PI dyes DNA and binds to DNA in proportion to the amount of DNA present in the cell. Cells in S phase will therefore have more DNA than cells in G1 and will take up proportionally more dye (more fluorescence) with a higher amount of DNA content. Cells in G2 should be twice as bright as cells in G1 phase. Using FACS and the PI dye, we can conditionally sort these cells using fluorescence.



Figure A.2. FACS sorted cells from mutant and wildtype treatments after UV

induction.

Table A.2. Predicted results for assay

Strain	TP53 Activity	Cell cycle changes after
		UV treatment
BAF3/Null	ON	More cells undergoing
		apoptosis
BAF3/WT-PPM1D	On-ish/Off-ish	Less cells undergoing
		apoptosis (less dividing
		cells)
BAF3/PPM1D-D314A	ON	More cells undergoing
(phosphate dead)		apoptosis

BAF3/PPM1D-Q520*	More OFF-ish	Less cells undergoing
(activating)		apoptosis

The results from Figure A.2 suggest that blood-specific PPM1D mutants tend to have higher phosphatase activity against TP53, and prevented the cells from apoptosis after DNA damage, suggesting that PPM1D mutations were potentially involved in cell proliferation regulation. Under conditions of BAF3/Null and the D314 phosphatase dead cell line, we would expect to always see active TP53 which should successfully halt the cell cycle process therefore more cells in S/G1. With the Q520 mutant, we expect to see higher expression of PPM1D thereby inactivating TP53 and more cells in G2 phase (mitotic cells-dividing). TP53 is unable to stop cells from entering the cell cycle even after DNA damage.

Protocol for Cell Cycle Analysis

Day 1

Prior to starting make sure cells are confluent. Move media to water bath.

- Transfer cells from flasks to 15 mL tubes. BAF3 cells are a cell suspension cell line (so most cells are in the media floating around).
- 2. Spin down 200 rpm for 5 minutes
- 3. Remove supernatant.
- 4. Resuspend pellet in 1 mL of appropriate media.
- 5. Count Cells
 - a. 1:5 dilution

- b. Add 80 ul of Trypan blue to small tube.
- c. Add 20 ul of sample to small tube.
- d. Calculate # of cells using hematocytometer. Need to have 10^6 cells
- 6. Spin down resuspension one more time.
- 7. Remove supernatant.
- 8. Add 1000 ul media.
- Label 6 well plates. One should be UV treated and the other should be no UV.
 Set up experiments in duplicates or triplicates if possible.
 - a. BAF3
 - b. WT
 - c. D314A
 - d. Q520

10. Add 2 mL of media to each well.

- 11. Add calculated amount of each sample to the appropriate well.
- 12. Transfer remaining cells back to cell culture flask with 6 mL of media.
- 13. Incubate Errbody!

Day 2

Wait 24 hours and irradiate the cells with 6 Gy UV

Day 4

Wait 48 hours and harvest cells for fixation

Fixation

- 1. Remove cells from 6 well plate and move to 15 mL tubes.
- 2. Centrifuge at 500 rpm for 5 min

- 3. Remove supernatant and resuspend cells in 1 mL of media.
- 4. Count cells.
- 5. Resuspend at 2x10⁶ cells in 1 mL ice cold buffer.
- Vortex gently, slowly adding the cell suspension dropwise to 9 mL of 70% ethanol in a 15 mL polypropylene centrifuge tube.
- 7. Store at 4 degrees C for 24 hours.

Day 5

- 8. Centrifuge cells at 200 x g, 10 min, 4 degrees Celsius
- 9. Resuspend pellet in 3 mL cold PBS and transfer to tubes.
- 10. Wash cells with cold PBS.
- 11. Resuspend cells in 300-500 ul PI triton staining solution to 10 mL of 0.1 % (v/v) Triton X-100 (Sigma) in PBS add 2 mg DNAse-free RNAse A.
- 12. Incubate 37 degrees Celsius for 15 min or for 30 min at 20 degrees Celsius.
- 13. Transfer tubes to ice or store at 4 degrees Celsius protected from light.
- 14. Acquire data on flow cytometer within 48 hours.
- 15. Perform FACS on department of pathology and immunology

5.7 Discovery of Truncation Mutations Leading to Protein Alterations

Studying the allele specific expression of nonsense mutations across genes genome wide will broaden our understanding of rules by which nonsense mediated decay can effectively degrade a transcript. To study mutation enrichment and degradation efficiency across cancer types, we propose a novel scoring method using RNA-Sequencing and whole exome sequencing variant allele fraction ratio as a proxy for allele specific expression. By comparing case expression to a control dataset in each cancer type, our analysis has uncovered an enrichment of nonsense mutations predicted to escape nonsense mediated decay (NMD) and a subset showing evidence of N and C terminally truncated proteins. Identifying variants that target a transcript for degradation via NMD or produce a transcript that could be translated to a truncated protein are both clinically pathogenic. We further expanded our investigation to consider enriched mutations across mutation types, including missense, silent and splice site variants, to identify activating and inactivating truncation mutations.

Of note our analysis identified a subset of mutations in a tumor suppressor, STK11, utilize a downstream start codon to create N terminally truncated proteins lacking the N terminal localization sequence in lung cancer samples. We propose a novel scoring method using RNA and DNA variant allele fraction ratio as a proxy for allele specific expression to study mutation enrichment and degradation efficiency across cancer types. Recurrent protein isoforms are shown to be present in prostate, lung, hepatocellular and tumor samples, but only a small percentage of samples in a few cancer types are shown to harbor mutations in splicing factors(Brooks et al.; Rajan et al.; Zhang et al.). Since the dominant isoform present in many tumors are not attributed to mutations in splicing factors, this directs us to focus on introduced genomic variants that are contributing to altering splicing patterns. Furthermore, since one-third of alternative splicing events in human genes are thought to cause NMD(Brogna and Wen; Lewis et al.; Ni et al.; Pan et al.; Venables; Weischenfeldt et al.; Weischenfeldt et al.), it is vital that we can predict nonsense mutations that will and won't be degraded by the introduced mutation.

One-third of SAVs are estimated to introduce a premature termination codon (PTC), which could lead to dominant negative or gain of function effects. Annotated nonsense mutations are predicted to create premature termination codons (PTC) in the resulting transcripts, which are predominantly degraded by the Nonsense Mediated Decay (NMD) pathway. The general rule of thumb is that PTC's located at least 50-55 bp upstream of the last exon-exon junction drive strong NMD, whereas those outside of this criteria are predicted to escape the degradation process. Finally, as noted in previous studies, N terminally truncated proteins can lead to translation re-initiation at a downstream codon, thereby bypassing NMD and creating a product with residual function. This exception to NMD is known for few genes, but has yet to be characterized in cancer and genome wide.

Understanding how SAVs can lead to alternative isoforms in tumor samples and determining if the new transcripts become a target of NMD is still an open area of study.

217

Identifying variants that target a transcript for degradation via NMD or produce a splicing product that could be translated to a truncated protein are both clinically pathogenic. A pan-cancer analysis of SAVs can expand the current paradigm about the exceptions and rules by which NMD and splicing are altered in cancer.

Dataset and mutation distribution

Stringent filters (Supplementary Methods) were used to create high quality mutation calls for 18 cancer types: breast adenocarcinoma (BRCA), bladder urothelial carcinoma (BLCA), cervical squamous cell carcinoma and endocervical adenocarcinoma (CESC), colon and rectum adenocarcinoma (COADREAD), glioblastoma multiforme (GBM), head and neck squamous cell carcinoma (HNSC), kidney chromophobe (KICH), kidney renal clear cell carcinoma (KIRC), kidney renal papillary cell carcinoma (KIRP), acute myeloid leukemia (LAML), low grade glioma (LGG), liver hepatocellular carcinoma (LIHC), lung adenocarcinoma (STAD), thyroid carcinoma (THCA), uterine carcinosarcoma (UCS) and uterine corpus endometrioid carcinoma (UCEC).

Filters: We collected nonsense mutations predicted to introduce termination codons into a transcript of interest. We performed RNA-Seq and DNA-Seq readcount analysis to determine sites that had at least 20X coverage. From the high coverage nonsense mutation gene set, we collected silent and missense mutations, performed readcount analysis and filtered out sites with less than 20X coverage. Finally sites with both copy number and gene expression data available through the Broad GDAC Firehose were collected for further analysis.

218

Our final mutation dataset consisted of 88,845 silent, nonsense and missense mutations in 5,023 genes across 16 cancer types from 3,129 samples.

Dataset and mutation distribution

Stringent filters (Supplementary Methods) were used to create high quality mutation calls for 18 cancer types: breast adenocarcinoma (BRCA), bladder urothelial carcinoma (BLCA), cervical squamous cell carcinoma and endocervical adenocarcinoma (CESC), colon and rectum adenocarcinoma (COADREAD), glioblastoma multiforme (GBM), head and neck squamous cell carcinoma (HNSC), kidney chromophobe (KICH), kidney renal clear cell carcinoma (KIRC), kidney renal papillary cell carcinoma (KIRP), acute myeloid leukemia (LAML), low grade glioma (LGG), liver hepatocellular carcinoma (LIHC), lung adenocarcinoma (LUAD), lung squamous cell carcinoma (LUSC), Stomach adenocarcinoma (STAD), thyroid carcinoma (THCA), uterine carcinosarcoma (UCS) and uterine corpus endometrioid carcinoma (UCEC). We collected 11,952 nonsense mutations predicted to introduce a premature termination codon into the transcript of interest. We additionally collected silent and missense mutations in all genes with nonsense mutations for comparison. Our final mutation dataset consisted of silent, nonsense and missense mutations in 6,107 genes across 18 cancer types in 3,863 samples. All mutations were collected in copy number neutral sites with at least 20 reads spanning the site of interest in RNA-Sequencing data and DNA-Sequencing.

Identification of activating nonsense mutations in copy number neutral data

219

To evaluate the effect of nonsense mutations on stability, we used a novel method to differentiate between nonsense mutations degraded and escaping degradation as measured by next generation sequencing data. We chose to leverage variant allele fractions (VAFs) measured by DNA and RNA sequencing of the tumor samples to define the relationship between the genomic position and transcriptome effect. One would expect a comparative relationship between RNA and DNA VAF if allelic content and transcription were directly proportional. But in practice there are a number of variables that can cause a deviation from this proportional relationship including post transcriptional modifications such as RNA degradation.

By comparing the RNA to DNA VAF ratio across exons for nonsense, missense and silent mutations in copy number neutral regions we can evaluate the positional effect of nonsense mutations. This comparison between mutation types across exons and genes will highlight exons of interest that have a higher than expected RNA VAF when compared to DNA VAF and can potentially be used as a proxy to measure sites that escape nonsense mediated decay.



Figure A.3. Bar plot of total sites with RNA/DNA VAF Ratio greater than (top) and less than one (bottom) separated by gene.

Figure A.3 highlights the top 14 exons found to have the largest number of nonsense mutations and their RNA/DNA VAF Ratios. Many exons predicted to contain nonsense mutations that escape NMD are present in the last exon of the gene including ARID1A, AHNAK, APC, SMAD2, as expected. Of the 2,538 exons from 2,194 genes with at least one nonsense mutation with an RNA/DNA VAF greater than one, 1,069 of the exons are classified as the last exon of the gene of interest. This confirmed our first main finding that many nonsense mutations in cancer act as expected and are degraded effectively by the NMD pathway. Interestingly, there are still a large number of nonsense mutations that are present at a high variant frequency in RNA and DNA, suggesting evasion of the degradation pathway. We focused on a set of tumor suppressor genes that contained a large number of nonsense mutations with high RNA VAF including CDKN2A, TP53, SMAD2, CREBBP, ARID1A, PTEN, MAP3K1, KMT2C and KMT2D.

A subset of our mutations had available normalized RSEM gene expression data to compare the gene expression of the case to associated controls. For each of the genes mentioned above, we compared the case expression of the gene of interest to control samples in the same cancer type, and also compared genes within the same pathway to identify downstream genes with altered expression due to the activating truncation mutation. After identifying potential activating mutations, each mutation was annotated using TransVar(Zhou et al.), a multilevel variant annotator.

One sample had a mutation in exon 1 of CDKN2A in HNSC and was found to have much higher expression than control samples. CDKN2A expression leads to activation of INK4A and ARF which in turn inhibits MDM2, CDK4 and CDK6. Gene expression of the downstream genes showed significantly decreased expression of MDM2 and CDK6 but lower expression of TP53 and RB1 which should have increased expression with decreased expression of MDM2 and CDK6. A closer look at the mutations present in this sample revealed a frame shift insertion in TP53 and upon inspection of the RNA-Seq showed very few reads spanning TP53 exons and present in a copy number neutral region and similar findings for RB1 (but without a mutation).

For each gene of interest we identified missense, silent, nonsense and splice site variants deemed to be activating mutations by our gene expression permutation test. We used a lolliplot to visualize the predicted activating mutations within each functional domain. Our findings identified a number of "hotspots" where mutations cluster that create similar truncating activating mutations.

We defined genes as expressed if they had a RSEM value greater than the lower quartile of the distribution of RSEM control values. Furthermore, we focused on sites that have RNA and DNA VAF ratio greater than 1 and RSEM value greater than the lower quartile for the distributed RSEM control values.

Measuring allele specific expression by integrating RNA and DNA variant allele fractions



Figure A.4. Comparison of RNA Variant Allele Fraction and DNA allele fraction by conventional annotation type. Top panels indicate all sites and bottom panels are only expressed.



Figure A.5. Normalized gene expression (RSEM) and variant allele fraction for all sites.

Table A.3. Total number of	of variants	in each	RNA/DNA	VAF	ratio	category
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	No F	ilter		Expre					
Mutation Type	Ratio >=	Ratio < 1	Fisher	Ratio >= 1	Ratio < 1	Fisher			
	1		(Silent)			(Silent)			
Missense	33304	24727	8.863e-	24632	17556	2.143e-			
Mutation			15			12			

Nonsense	2219	6329	<2.2e-16	1556	3623	<2.2e-16
Mutation						
Silent Mutation	11960	10049	-	8793	7150	-

76,501 mutations fall into copy number neutral sites and 63,489 mutations have case RSEM greater than the lower quartile of matched cancer control samples lacking mutations in the gene of interest. We classify these mutations as expressed relative to their control samples.

We chose to leverage variant allele fractions (VAFs) measured by DNA and RNA sequencing of the tumor samples to define the relationship between the genomic position and transcriptome effect. One would expect a comparative relationship between RNA and DNA VAF if allelic content and transcription were directly proportional. But in practice there are a number of variables that can cause a deviation from this proportional relationship including post-transcriptional modifications such as RNA degradation. Figure A.4 highlights the relationship between DNA and RNA VAF across missense, nonsense and silent mutations while table 1 gives a numerical representation of the number of mutations falling above a slope of 1. Nonsense mutations show an increased number of mutations with higher DNA VAF than RNA VAF, suggesting a higher level of degradation efficiency compared to missense and silent mutations. Overall we can see a higher level of RNA VAF enrichment of missense mutations (positive selection) compared to silent mutations while nonsense mutations show on overall negative selection.

Using the silent mutation RNA and DNA VAF distribution, we can make the assumption silent mutations follow a null distribution. We used fishers exact test to compare the proportion of mutations greater than and less than a RNA DNA VAF Ratio of 1 to determine if the difference in proportions is significant between both missense and nonsense with silent mutations as the null distribution.



Novel inactivation of STK11

Figure A.6. Lolliplot of novel inactivating mutations in STK11. Statistics of STK11 variants below.

STK11 Ensembl Transcripts





Figure A.7. All Alternative isoforms of indicated STK11 mutants.

We identified 7 mutations in the canonical transcript of exon in ensembl transcript ENST00000326873. We used TransVar¹ to perform equivalence annotation to map the mutation of interest to all other transcripts defined by Ensembl based on the genomic coordinates of the variant. To determine which transcript is readily expressed in the tissue type, we determined the total number of reads supporting the first and last base of the exon unique to each Ensembl transcripts. For the example of STK11 we determined transcript X is expressed in LUAD by assessing number of reads supporting the unique exon for that transcript.

Studies found that the subcellular localization of kinase deficient mutants were found in the nucleus while mutants with disrupted NLS were localized in the cytoplasm. Furthermore, the mutant with the disrupted NLS could still induce G1 cell cycle arrest comparable to the wild type protein. We hypothesize a subset of nonsense mutations in the first exon of STK11 utilize a downstream start codon to create N terminally truncated proteins lacking the N terminal localization sequence.

Identification of activating mutations using allele specific expression

Our analysis was able to pick up well known activating mutations in key cancer genes including EGFR. Of the 28 EGFR mutations tested in our gene expression test comparing case to control gene expression, 16 had RNA VAF greater than 10% suggesting some level of transcription supporting the variant allele. 9 of the 16 mutations are functionally characterized activating mutations in EGFR including: L858R⁴ (4 - LUAD), L861Q (1 - LUSC), G719A⁴ (1 - LUAD), G598A,V (2 –LGG), R108K⁵ (1 - GBM). Furthermore, a tool using protein structure guided discovery identified R252C/P (2 – LGG), S768I (1 – LUAD) and L833V (1 – LUAD) as potential activating mutations due to their proximity to known activating mutations⁶.

Kelch-like erythroid cell derived protein with CNC homology (ECH)-associated protein 1 (Keap1) is essential in the regulation of cytoprotective and detoxifying defense systems. Keap1 is responsible for sequestering nuclear factor erythroid 2-related factor 2 (Nrf2 or NFE2L2) in the cytoplasm and interacts with Cul3-E3 ubiquitin ligase complex to target Nrf2 for ubiquitination and degradation by the proteosome. Many mutations in lung cancer disrupt the binding of Keap1 to Nrf2, thereby increasing the presence of free active Nrf2, or the degradation efficiency of Nrf2 by disrupting binding to Cul3⁷. In 2014, several "superbinder" mutants were found to increase the levels of Nrf2 within the nucleus without disrupting the interaction between KEAP1 and Nrf2⁸. To evaluate how this could be the

case, the authors looked at protein turnover of Nrf2 and found the superbinder mutants R320Q and R470C dramatically stabilized Nrf2 more so than the wild-type Keap1 but maintained successful ubiquitination. This surprising result could suggest that the increased affinity for Keap1 for Nrf2 suppresses substrate turnover by disrupting a degradation step post ubiquitination.

Assessing degradation efficiency of nonsense mutations using allele specific expression



Figure A.8. (A) Allele specific expression denoted by RNA/DNA VAF Ratio of nonsense mutations across exons. Red corresponds to mutations found in the first exon, green in middle exon and blue in last exon. Wilcox test was used to compare groups of nonsense mutations located in the first, middle and last exons of a gene. (B) RNA/DNA VAF Ratio distribution across all cancer types for mutations found in the first, middle and last exons. (C) RSEM distribution across all cancer types for mutations found in first, middle and last exons.

Nonsense mutations in the last exon tend to have higher allele specific expression than mutations in another exon, as expected. Interestingly, nonsense mutations in the first exon show a similar trend as the last exon mutations (higher allele specific expression). N terminally truncated proteins can lose their localization sequences, thereby altering the cellular localization of the protein. Studies of β -globin show a very distinct transition between nonsense mutations in the first exon that are able to bypass NMD and those that are subject to degradation. The potential of the transcript to evade degradation is dependent upon the presence of a downstream start codon to reinitiate translation.

All nonsense mutations from the first and last exon were annotated to all possible transcripts using TransVar to determine mapping potential to alternative isoforms. 143 of the 207 nonsense mutations annotated to the first exon maintained the nonsense mutation prediction on alternative isoforms. Of the 143 variants annotated solely to the first exon, 58 variants had an RNA VAF less than 10% suggesting selection against the variant allele in the tumor or the presence of a subclonal mutation. The presence of the remaining 85 variants with greater than 10% RNA VAF supporting the N-terminal nonsense mutation could be explained by translation readthrough or reinitiation at a downstream start codon. Two studies reported leak expression when nonsense mutations were introduced up to residue 70 of the Shaker voltage-gated potassium (Kv) channel¹⁰ and residue 26 of B-globin⁹, suggesting a sharp divide between mutations susceptible

and resistant to NMD. Both studies came to the conclusion that the likely mechanism is due to reinitiation of translation at a downstream translation start site, and in the case of the Kv channel, non-canonical start codons were used (AAG, AGG). Additionally, translation isn't limited to the next start codon, but alternative start sites farther downstream can be used by a fraction of ribosomes.

5.8 Viral integration

Contribution: I helped with gene expression figures related to discordant pair analysis. All Figures and supplemental documentation can be seen in the following publication.

Divergent viral presentation among human tumors and adjacent normal tissues, Scientific Reports, 2016 Jun. doi:10.1038/srep28294

Cao S., Wendl M., Wyczalkowski M.A., Wylie K., Ye K., Jayasinghe R.G. et al.

5.9 **DROSHA Mutation Analysis**

Through this collaboration we identified novel mutations in DROSHA and Dicer and evaluated RNA-Seq expression for associated downstream products that could be altered. Of note, for one of the mutated samples, MDM2 expression was increased, matching our collaborators findings in a patient of interest.



Figure A.9. DICER1_14_95556886_T_G_TCGA-EL-A3GO

Human hg19		chr14			chr1	4:95,556,	866-95,5	56,906	Go	1 -	 ¢ 🗆	x F	1							IGV																			=		
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	÷				8.84.87	1 hp								95.	556,890 hp						411	əp —			95.5	556,890 hp									95	,556,900 bp					
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Figure A.10. DICER1_14_95556886_T_G_TCGA-EM-A2CT



Figure A.11. DICER1_14_95557629_T_C_TCGA-EL-AD35



Figure A.12. Distribution of normalized expression for downstream genes and DICER1 for mutants vs. control samples

Table A.4: Samples with DROSHA Mutations
Ca	Diag	С	star	R	V	amino_a	samp	Normal	Tumor	Tumo	Selecti
nc	nosi	h	t	е	а	cid_cha	le	wxsv	wxsv	r RNA	on
er	S	r		f	r	nge		AF	AF	VAF	
	(Ag										
	e)										
LA	45	5	314	A	-	p.L879fs	TCGA	0.57	0.48	0.24	Not
ML			516				-AB-				under
			86				2983				selectio
											n
ST	75	5	315	G	A	p.R16*	TCGA	0.53	0.37	0.15	Not
AD			269				-HU-				under
			94				8610				selectio
											n
ΤН	77	5	315	С	Т	e2-1	TCGA	0.42	0.54	NA	Under
CA			270				-EM-				selectio
			20				A2CO				n
UC	65	5	314	Т	С	e23-2	TCGA	0.39	0.38	NA	Not
EC			317				-D1-				under
			87				A2G7				selectio
											n
UC	59	5	315	G	А	p.R287*	TCGA	0.54	0.43	NA	Not
EC			213				-AX-				under
			18				A1CJ				

				selectio
				n

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