Specific Aims: Splicing is a process where introns are removed from pre-mRNAs and the exons are stitched together. This offers the potential to introduce tremendous diversity into the proteome by changing which sequences from the pre-mRNA are included as exons. Accordingly, more than 90% of human genes are alternatively spliced. Errors in splicing are involved in a wide variety of human diseases including familial dysautonomia, early onset Parkinson disease, and cancer.² Recent efforts have produced drugs that modulate splicing outcomes to treat spinal muscular atrophy, Huntington's disease, Duchenne muscular dystrophy, and cancer.³ Improving our understanding of splicing will deepen our knowledge of development and differentiation and our ability to target these processes with therapeutics. Much progress has been made using massively parallel splicing assays (MPSAs) to probe the mechanisms of splicing.^{4–17} These efforts have utilized Illumina technology which cannot produce reads capable of covering the whole length of most genes, greatly affecting the isoforms that can be detected. This issue has been overcome by studying systems shrunk to fit an Illumina read or by ignoring unexpected isoforms. Yet the median human intron is 1.7 kb in length 18 far longer than an Illumina read and perturbations can cause complicated changes in splicing outcomes.^{8,19,20} Further, MPSAs have so far focused on the decision to include or exclude a single exon. However, recent work has begun to show that correlated splicing decisions across exons are far more common than was previously understood.^{21–24} A particularly interesting case of correlated splicing is mutually exclusive exons (MXEs). MXEs are clusters of exons that are spliced such that every isoform includes exactly one exon from the cluster. A recent analysis of human RNA-seg data identified 629 MXE clusters, of of which less than 25% had an identifiable mechanism for maintaining mutual exclusivity (MMX).²³ In order to overcome these issues, I propose to develop a long read massively parallel splicing assay (LR-MPSA) capable of handling correlated splicing, full size introns, and complicated isoform distributions. I will apply this assay to investigating MXEs with a focus on Pyruvate Kinase M (PKM). Aim 1: Develop an MPSA protocol capable of handling the complexity of natural splicing. Sub-aim 1.1: Create an MPSA protocol utilizing nanopore sequencing technology. I will develop an LR-MPSA optimized for characterizing the mechanisms of complex splicing decisions in the context of full native introns and all isoform outputs. I will adapt MPSA techniques honed by the Kinney lab 17,25 by creating new sample and library preparation pipelines optimized for unbiased isoform enrichment and nanopore sequencing. Sub-aim 1.2: Develop an analysis pipeline for quantifying isoform abundances from LR-MPSA data. Calling low abundance RNA isoforms from nanopore reads faces challenges from the low depth and high noise. I will develop an analysis pipeline for identifying and quantifying isoforms that optimizes for power to detect low abundance isoforms and minimizes incorrect isoform assignment. I will simulate nanopore reads given a known ground truth isoform distribution for each variant to test and optimize the new pipeline at various

Aim 2: Characterize the mechanism for mutual exclusivity in Pyruvate Kinase M (PKM) exons 9 and 10. PKM catalyzes the rate limiting step in glycolysis and has two primary isoforms defined by the inclusion of either exon 9 or exon 10. The conversion between these two isoforms is a primary driver of the Warburg effect and is important in a wide variety of cancers. However, the mechanism for MMX between exons 9 and 10 is unknown.

Sub-aim 2.1: Identify sequence elements necessary for mutual exclusivity in PKM splicing. I will use PKM minigene constructs designed by the Krainer lab¹⁹ to screen for mutants that loose MMX. I will use mutant scans over exons 9 and 10 to find MMX elements because previous work has shown that regulatory sequences are located in these exons. ¹⁹ Broader random mutagenesis will map elements in introns. MMX regions will be mapped at higher resolution with deep mutagenesis. If the nanopore screening technique fails, I will use short-read or low throughput methods for screening.

Sub-aim 2.2: Mechanistically characterize sequence elements enforcing mutual exclusivity. I will follow up identification of MMX elements with mechanistic studies. One hypothesis for the identity of new MMX elements is RNA secondary structural elements. I will use mutant cycle analysis to show dependence on structure for any identified secondary structural elements. Another hypothesis is that MMX elements are motifs for RNA binding proteins (RBPs). I will use pull downs as well as motif mutants coupled with RNAi knockdown and over-expression of RBPs to demonstrate the importance of RBP binding motifs.

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depths and error profiles.

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