

## Abstract

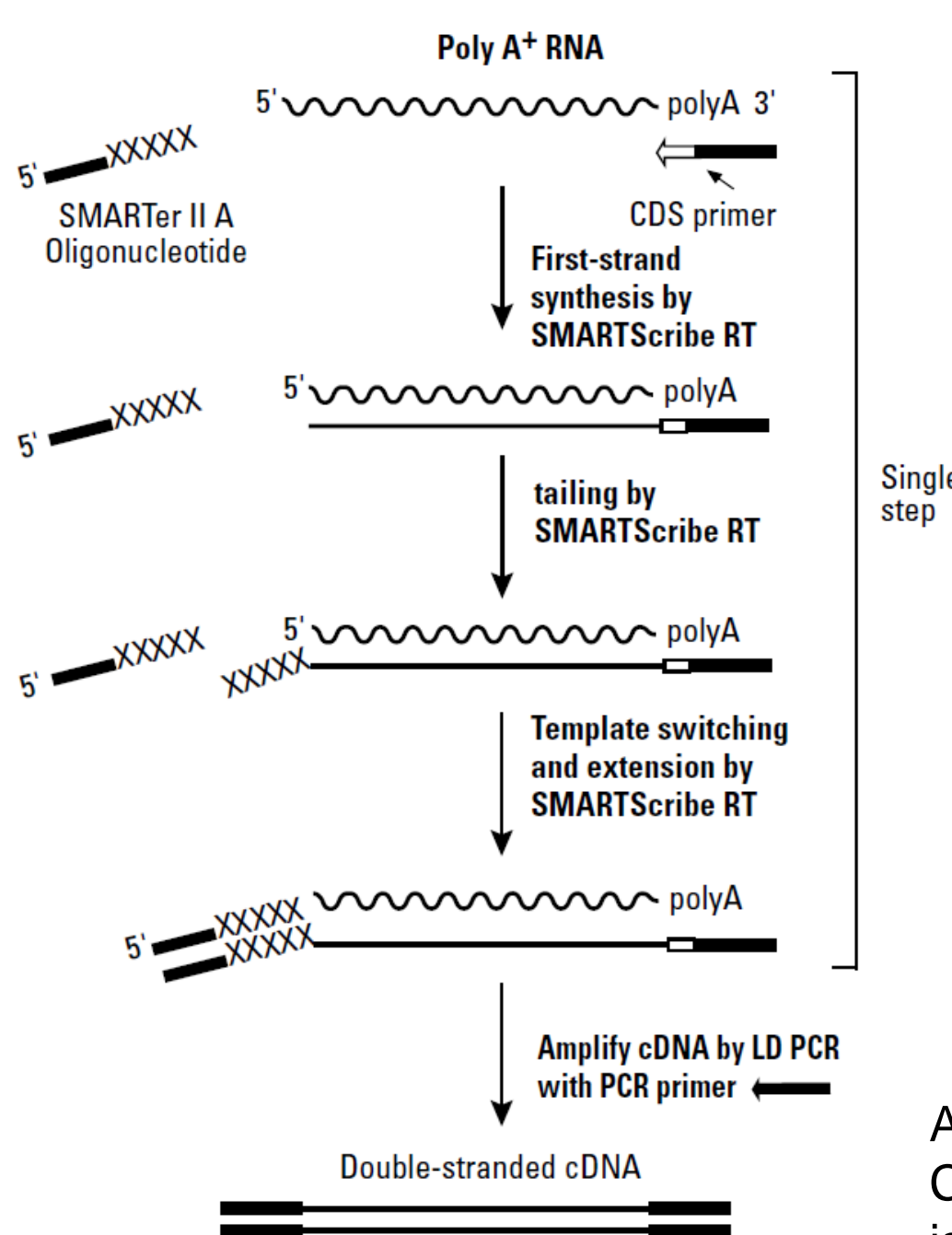
The aberrant transcription and expression of alternative RNA isoforms has been observed in multiple types of cancer, and is hypothesized to contribute to oncogenesis in certain cancer subtypes. Identification and annotation of cancer-specific mRNA isoforms is critical to understanding how mutations in the genome affect the biology of cancer cells. While microarrays and other NGS-based methods have become useful for studying transcriptomes, these technologies yield short, fragmented transcripts that remain a challenge for accurate, complete reconstruction of splice variants. In cancer proteomics studies, the identification of biomarkers from mass spectroscopy data is often limited by incomplete gene isoform expression information to support protein to transcript mapping.

The Iso-Seq protocol developed at PacBio offers the only solution for direct sequencing of full-length, single-molecule cDNA sequences needed to discover biomarkers for early detection and cancer stratification, to fully characterize gene fusion events, and to elucidate drug resistance mechanisms. Knowledge of the complete isoform repertoire is key for accurate quantification of isoform abundance. As most transcript sizes range from 1 – 10 kb, fully intact RNA molecules can be sequenced using SMRT Sequencing without requiring fragmentation or post-sequencing assembly. However, some cancer research applications have presented a challenge for the Iso-Seq protocol, due to the combination of limited sample input and the need to deeply sequence heterogeneous samples.

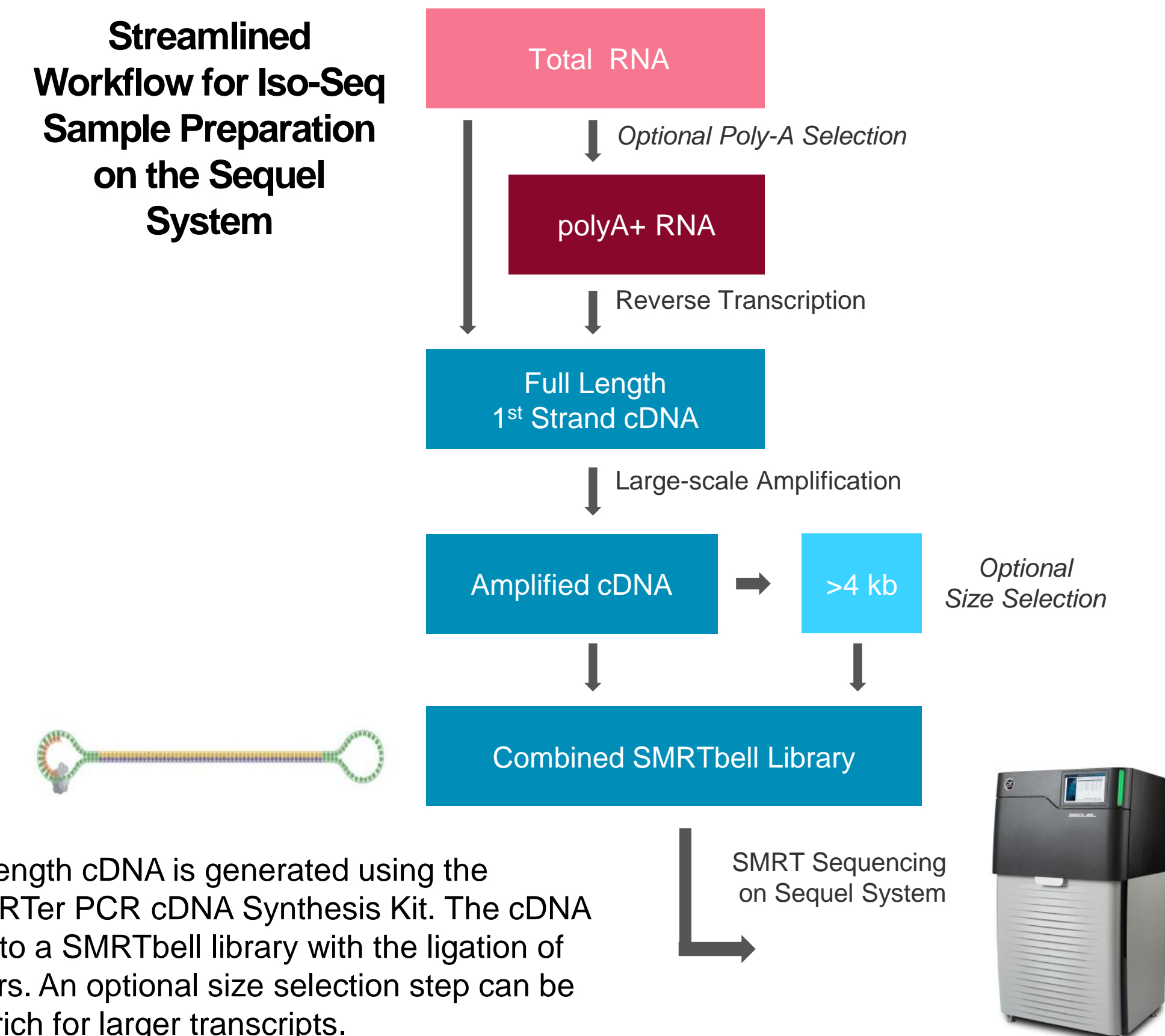
Here, we report the optimization of the Iso-Seq library preparation protocol for the PacBio Sequel platform and its application to cancer cell lines and tumor samples. We demonstrate how loading enhancements on the higher-throughput Sequel instrument have decreased the need for size-fractionation steps, reducing sample input requirements while simultaneously simplifying the sample preparation workflow and increasing the number of full-length transcripts per SMRT Cell. The results highlight the potential for broader application of the Iso-Seq method to more comprehensively characterize alternative splicing in cancer.

## Iso-Seq Sample Preparation Methods

### Clontech SMARTer PCR cDNA Synthesis Kit



### Streamlined Workflow for Iso-Seq Sample Preparation on the Sequel System

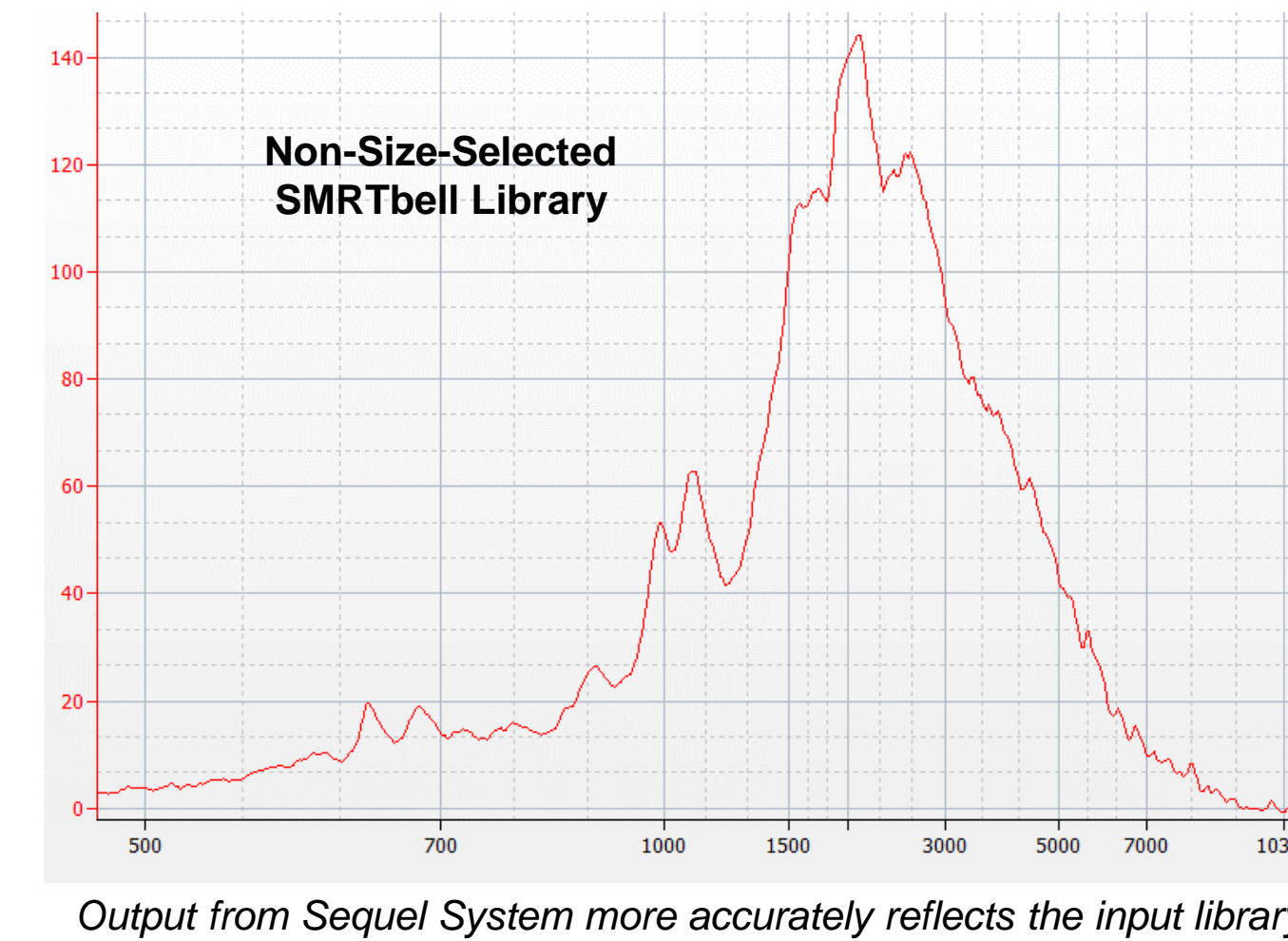
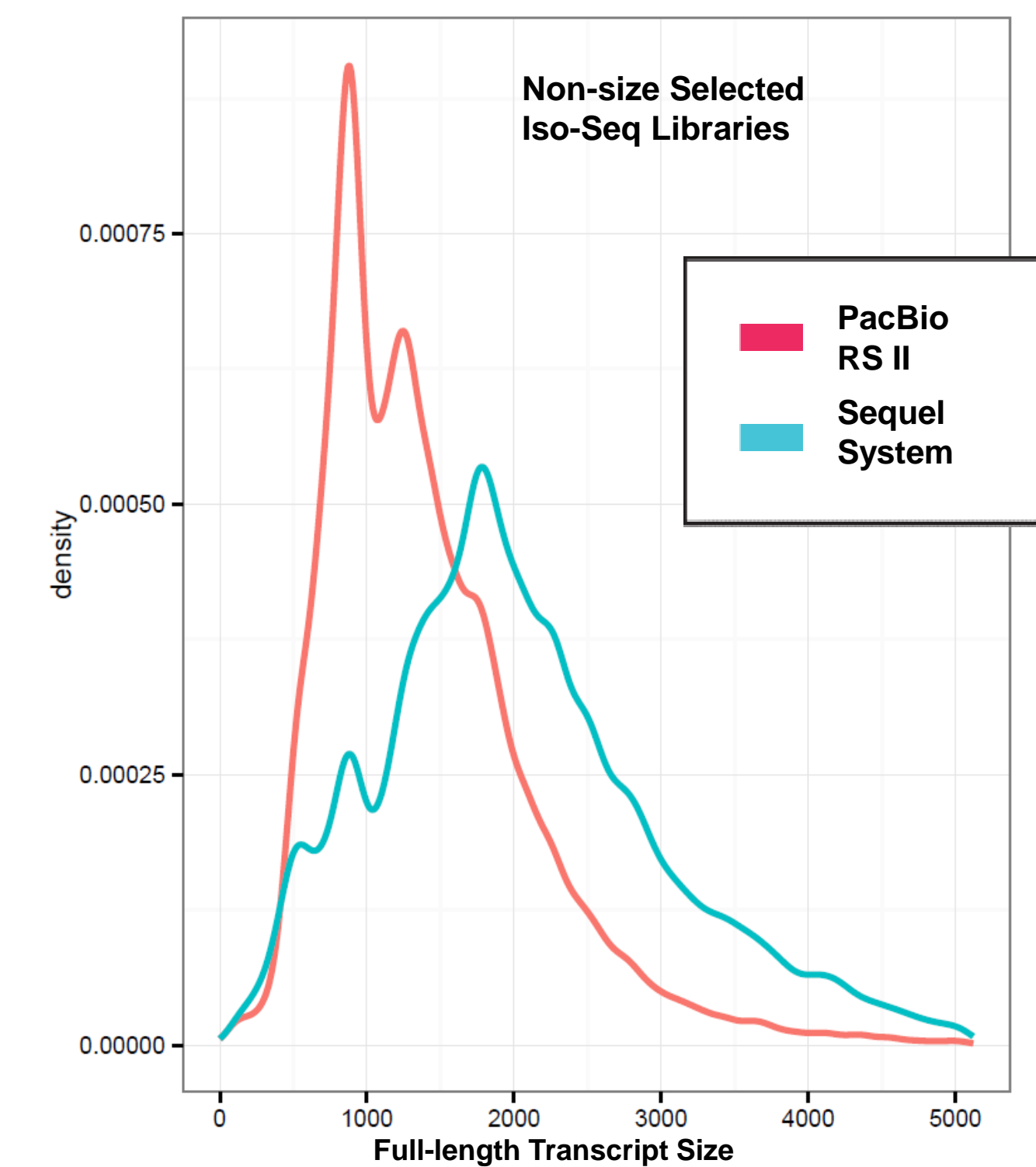


Amplified full-length cDNA is generated using the Clontech SMARTer PCR cDNA Synthesis Kit. The cDNA is converted into a SMRTbell library with the ligation of hairpin adapters. An optional size selection step can be included to enrich for larger transcripts.



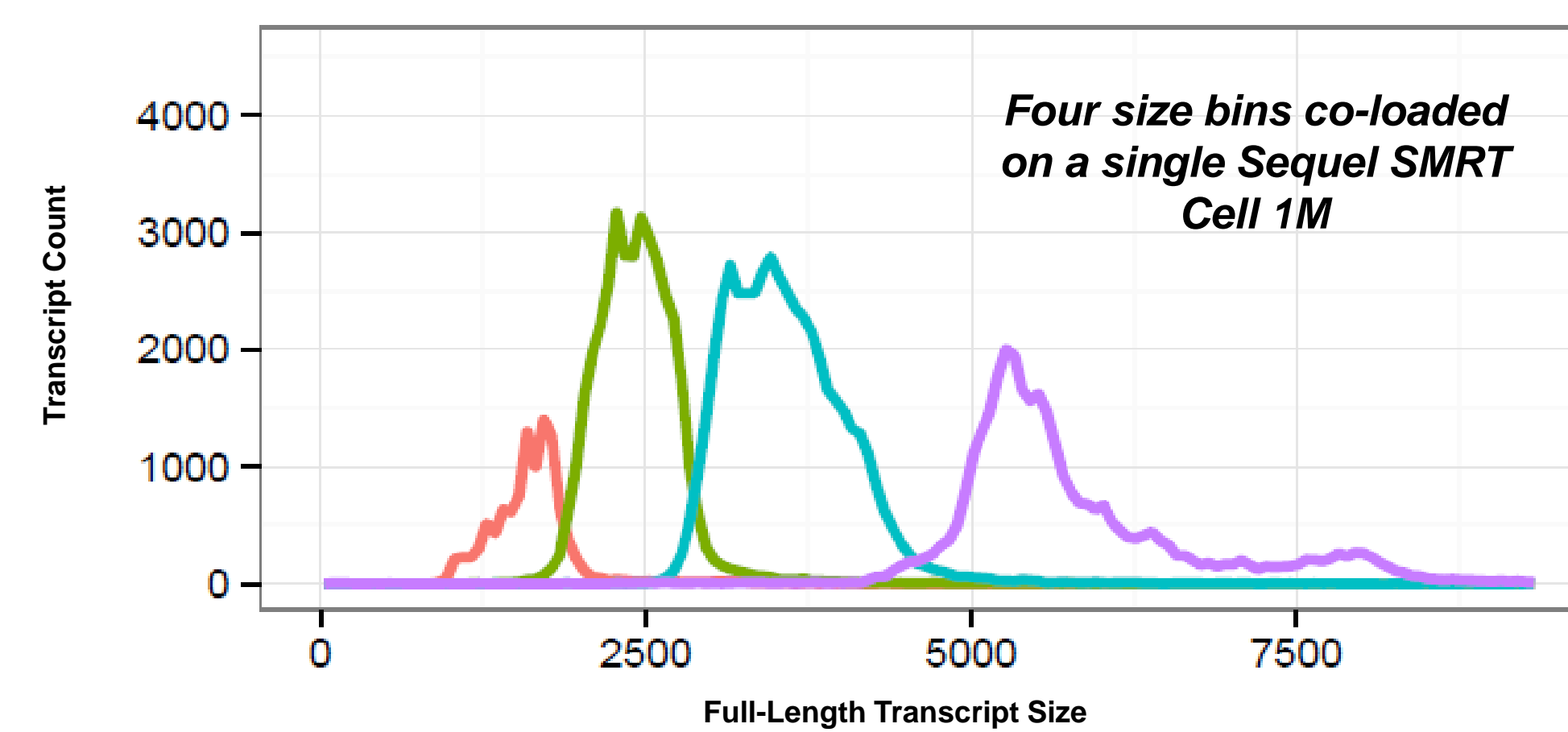
## Optimizing the Iso-Seq Application on the Sequel Platform

### Magbead-loaded Samples on Sequel System Have Decreased Loading Bias



A non-size selected Iso-Seq library was run on both the PacBio RS II and Sequel System. A histogram of number of full-length sequences by transcript length is plotted on the left. The full-length cDNA sequences run on the Sequel System closely resembles the size distribution of the input SMRTbell library, as shown in the bioanalyzer trace of the non-size selected library above.

### Multiple Size-selected Libraries Can Be Co-loaded on the Sequel System



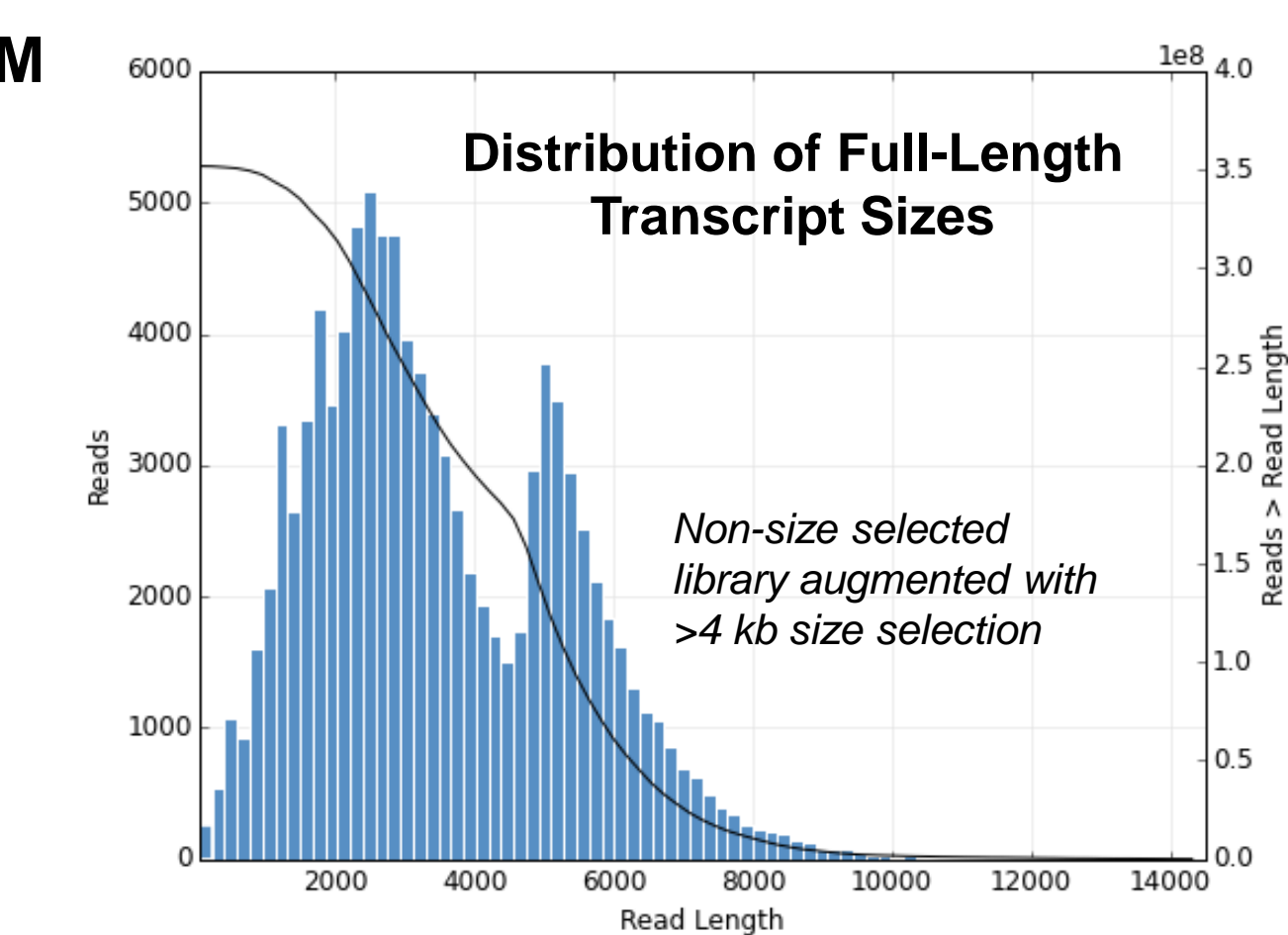
Size-selected Iso-Seq libraries were combined and run together on a single Sequel SMRT Cell 1M. Amplified cDNA had been size fractionated into four size bins (1-2 kb, 2-3 kb, 3-6 kb, and 5-10 kb) using the Sage Science BluePippin system, per the RSII Iso-Seq protocol. Barcoded SMRTbell libraries were made from each of the four size bins, to enable deconvolution post sequencing. A histogram of the full-length transcript lengths from each of the four size bins is shown at left, demonstrating that all four size bins are well represented when co-loaded. To boost representation of the longest isoforms, customers may optionally size select cDNA > 4 kb and pool at a non 1:1 ratio.

## Sequencing of NALM6 Precursor B-ALL Cell Line on Sequel

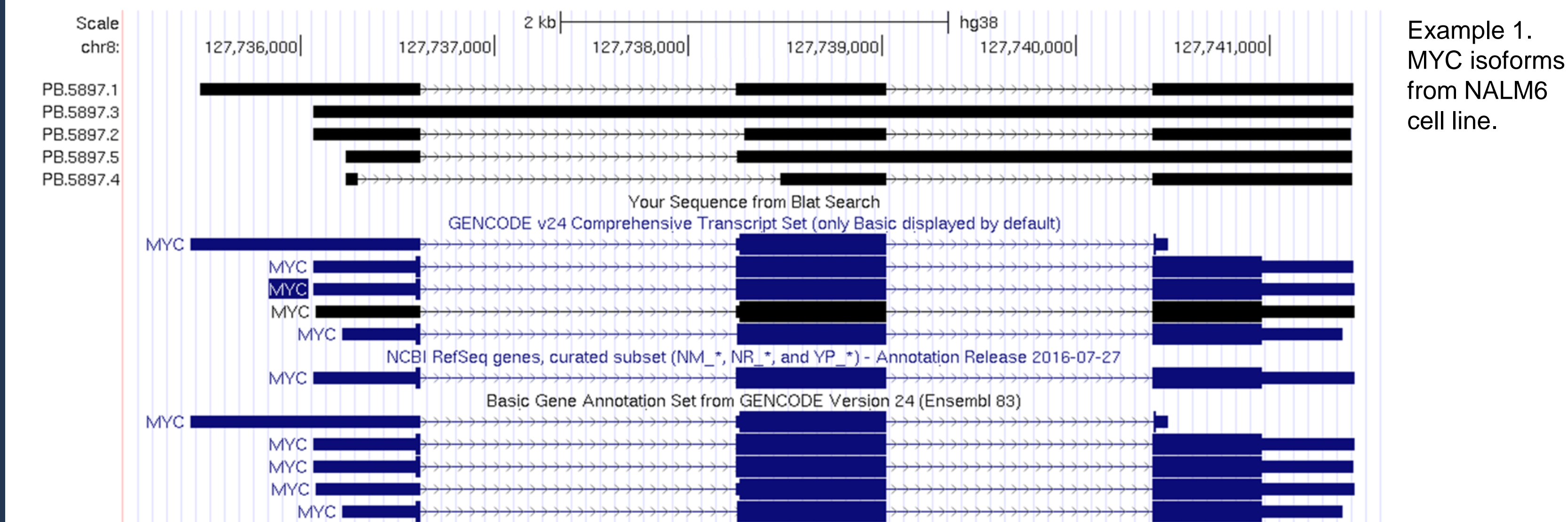
### Typical Sequencing Statistics From a Single Sequel SMRT Cell 1M

Transcript Classification		Transcript Clustering	
Value	Analysis Metric	Value	Analysis Metric
595,533	Number of consensus reads	143,392	Number of unpolished consensus isoforms
294,535	Number of five prime reads	17,765	Number of polished high-quality isoforms
338,803	Number of three prime reads	125,467	Number of polished low-quality isoforms
323,415	Number of poly-A reads	3,196	Mean unpolished consensus isoforms read length
230	Number of filtered short reads		
348,362	Number of non-full-length reads		
246,941	Number of full-length reads		
244,521	Number of full-length non-chimeric reads		
738,399,613	Number of full-length non-chimeric bases		
3,019	Mean full-length non-chimeric read length		

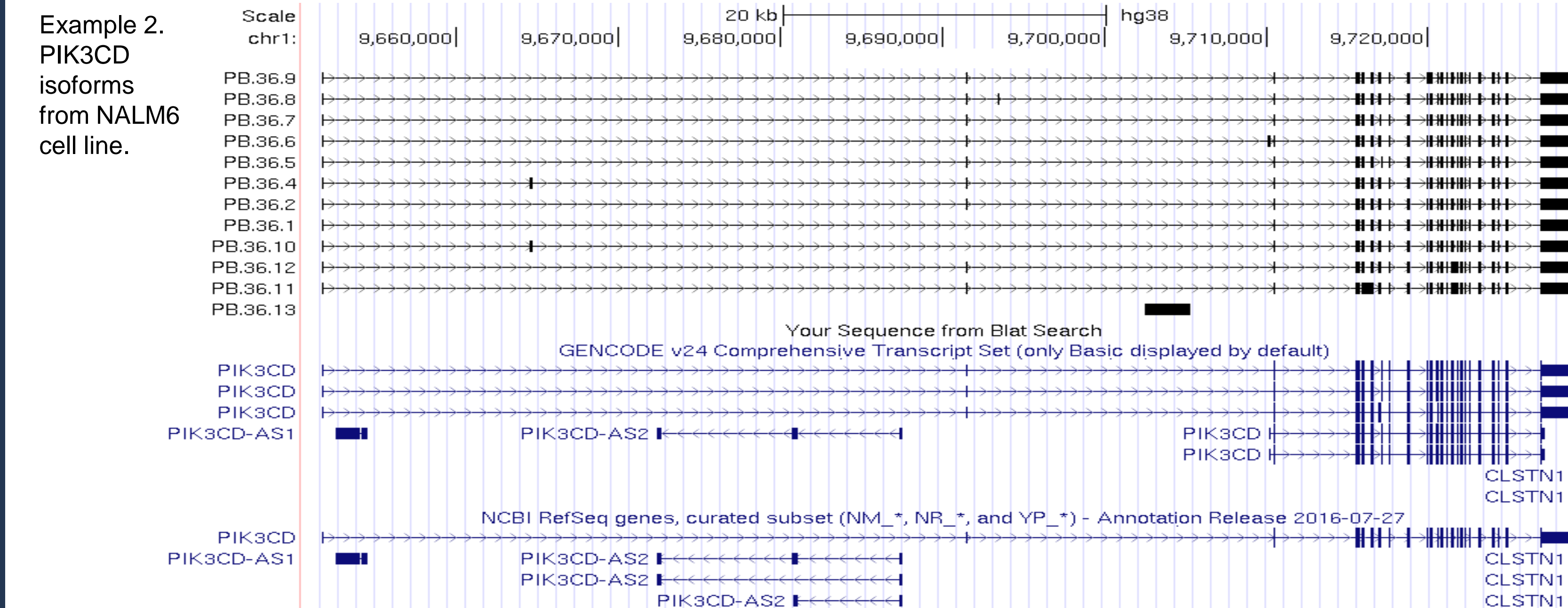
Data from a single, optimally loaded SMRT Cell processed in SMRT Analysis using the Iso-Seq analysis pipeline.



## Alternative Splicing in Cancer



Example 1. MYC isoforms from NALM6 cell line.



Example 2. PIK3CD isoforms from NALM6 cell line.

## Summary and Resources

- PacBio Iso-Seq method generates full-length transcript sequences without the need for assembly of short fragments
- Decreased loading bias on the Sequel System allows for a simplified Iso-Seq sample prep workflow that does not require multiple size-selection steps
- Barcoding during the cDNA generation allows multiplexing samples in a single SMRTbell library
- Improved throughput on the Sequel instrument increases the number of full-length transcripts per SMRT Cell.
- The Iso-Seq method is a powerful tool in the study of cancer providing full-length isoforms, alternative splicing information, and the capability to identify fusion genes.

More information on full-length transcript sequencing (Iso-Seq Application) can be found on the PacBio website: <http://pacb.com/iseq>