# Unlock insights from RNA-Seq studies

Explore transcriptomics data and gain a deeper understanding of gene expression with Partek™ Flow™ software

**Partek Flow software** offers a user-friendly bioinformatics solution for analyzing and visualizing RNA-Seq data, empowering researchers of all skill levels to maximize insights from transcriptomics data.



#### **Advanced visualization tools**

Create informative, publication-ready visualizations with just a few clicks



#### **Powerful statistics**

Access industry standard statistical algorithms for results you can trust



#### Point-and-click analysis

Analyze data easily with an intuitive, context-sensitive interface



#### **Customizable workflows**

Build custom analysis pipelines visually and share with collaborators

### Partek Flow tools for RNA-Seq analysis

- Principal component analysis (PCA)
- Differential expression analysis (LIMMA, ANOVA, DESeq2, and more)
- Interactive dot plot, volcano plot, heat map, scatter plot, chromosome view, and Venn Diagram
- Gene ontology and KEGG pathway enrichment
- K-means, hierarchical, and graph-based clustering



### **Powerful analysis** and visualization tools

Whether looking for patterns of gene expression changes in time series data, building a classification model from published data, or discovering biomarkers by identifying correlation between, Partek Flow software has the tools for easily analyzing and visualizing your RNA-Seq data.

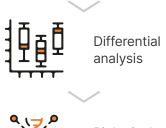
### **Cross-platform integration** for multiomic insights

Analyzing multiple omics data sets in parallel provides a more complete view of the molecular mechanisms underlying phenotypes. Partek Flow software enables integration and side-by-side visualization of data from multiple assay types, including RNA-Seq, ChIP-Seq, ATAC-Seq, microRNA expression, and microarray analysis.

### Steps in the RNA-Seq analysis workflow









Biological interpretation

## Data to **biological insights** in a single solution

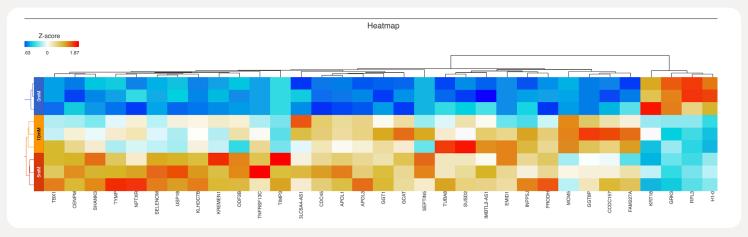
- Start the RNA-Seq analysis pipeline with either unaligned, aligned, or quantified reads, followed by trimming adapter sequences and low-quality bases
- Align reads to any reference genome or transcriptome using popular aligners, including STAR, GSNAP, Bowtie 2, TMap, HISAT-2, and more
- Use Salmon for ultrafast pseudoalignment and transcript quantification
- Map aligned reads to any public or user-defined transcriptome database
- Normalize data using total count, RPKM, and TMM
- Detect differentially expressed genes or transcripts
- Identify alternatively spliced transcripts and fusion genes
- Remove technical noise with flexible feature filtering options

## Multiple ways to explore RNA-Seq data

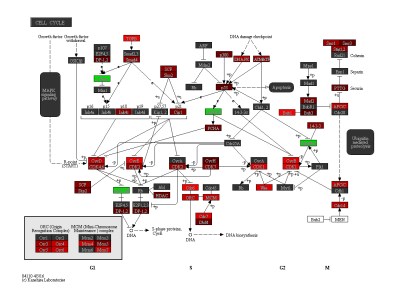
Transform raw RNA-Seq data into interactive figures and maximize biological insights from your transcriptomics research. The integrated genome browser in Partek Flow software makes visualizing isoforms and differential expression fast and easy.







Visualize large amounts of multi dimensional data, compare gene expression patterns between experimental groups, and identify clusters of rows with similar values using heat maps.





Analyze enriched pathways visually by color coding KEGG pathway maps with fold-changes or p-values.

Abbreviations: ANOVA, analysis of variance; API, application programming interface; ATAC-Seq, assay for transposase-accessible chromatin with sequencing; ChIP-Seq, chromatin immunoprecipitation with sequencing; GSNAP, gene set analysis; KEGG, Kyoto encyclopedia of genes and genomes; QA, quality assurance; QC, quality control; REST, representational state transfer; RPKM, reads per kilobase of transcript per million mapped reads; STAR, spliced transcripts alignment to a reference; TMap, Torrent mapping alignment program; TMM, trimmed mean of M-value.

#### illumına<sup>®</sup>

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