



JMP® Genomics

Discover the biological patterns in genomics data.
From the two most trusted names in analytic software: SAS and JMP.

What is JMP® Genomics?

JMP Genomics is a statistical discovery software tool from the two most trusted names in analytic software: SAS and JMP. Research organizations use JMP Genomics to uncover meaningful patterns in high-throughput genetics, expression, copy number and proteomics data. Dynamically interactive graphics make it easy to explore data relationships using a comprehensive set of traditional and advanced statistical algorithms.

Why is it important?

Research organizations want to maximize their return on investment of the time, money and resources required to generate high-quality genomic data sets. Specialized statistical analyses can help identify the “nuggets of gold” hidden in long lists of candidate genes or biomarkers. Whether it’s used to identify potential drug targets, to explore the biology of a model organism or to develop a predictive disease model, JMP Genomics helps researchers gain a competitive advantage by quickly identifying key genes or proteins.

Who should use it?

JMP Genomics is designed for biologists, biostatisticians, statistical geneticists and students engaged in analyzing the vast stores of data that are common in genomic research. It delivers a comprehensive set of analysis methods in a single desktop software package. Adopting new software across a large organization can be challenging. That’s why JMP Genomics offers analytics for many data types in the same package, making it easy for you to move into new areas of genomics as the scope of your research expands.

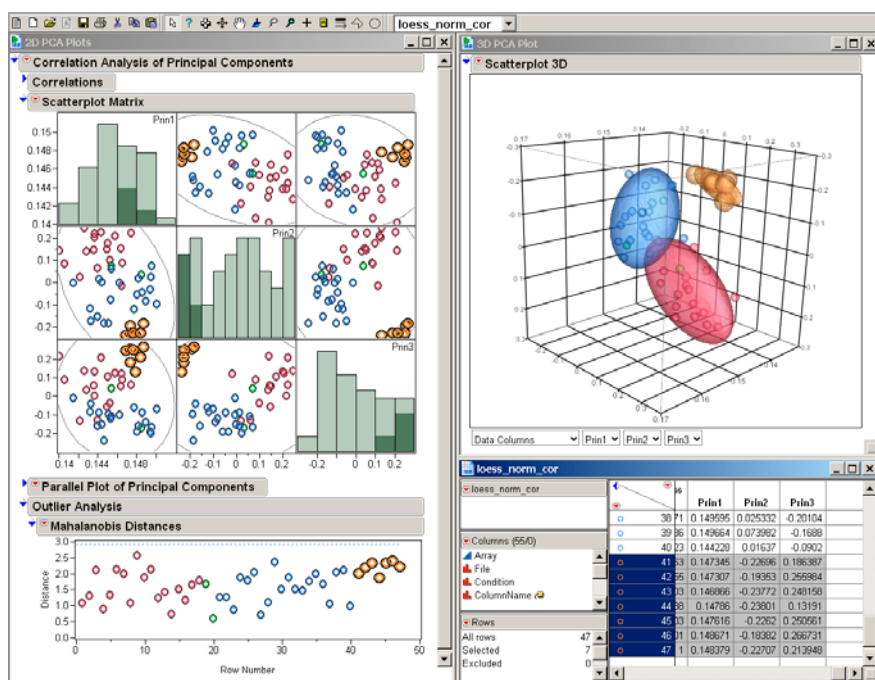
JMP® Genomics software from SAS brings high-powered, sophisticated genomic data exploration and analysis to the desktop. Its unique pedigree integrates the full power of the JMP statistical discovery platform with industry-leading SAS® Analytics tailored for heavy-duty processing of genomics data sets.

This all-in-one solution helps biologists, biostatisticians and statistical geneticists understand data generated from genetics, expression, exon, copy-number and proteomics studies. JMP Genomics dynamically links

“I think people are starved for software with this level of statistical power and flexibility.”

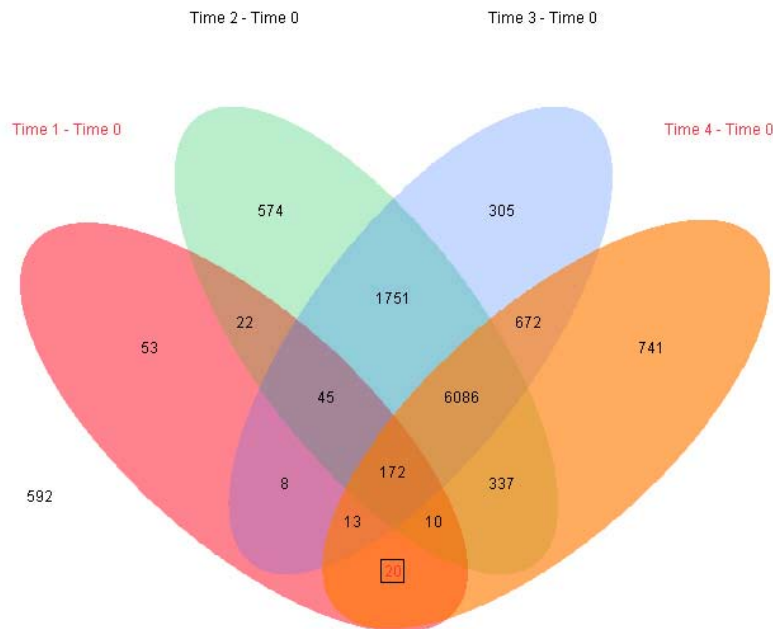
Erik Sulman, MD, PhD
Cancer Researcher
Houston

advanced statistics with graphics to provide a complete and comprehensive picture of research results, whether the data comes from traditional microarray studies or data summarized from next-generation technologies.



Visual representations of genomics data make it simple to see patterns in your data. JMP Genomics links graphics and data tables for easy browsing of significant results. Output of principal components analysis in JMP Genomics includes both 2-D and 3-D plots, both shown here, where enlarged markers in the graphics correspond to selected rows in the data table.

**STATISTICAL
DISCOVERY.™
FROM SAS.**



Visualize shared patterns with multiway Venn diagrams. Overlay statistical findings and annotation categories to drill down on the most important gene sets.

With JMP Genomics, experienced biologists, biostatisticians and statistical geneticists enjoy a menu-driven system that simplifies the workflow throughout the research process and interactive data visualization capabilities that let them see and explore their data from every angle, then easily share findings with colleagues. Even students new to genomic analysis quickly begin

“We take the graphical features for granted. But to be able to visualize that separation [in high-dimensional data] is so wonderful. Important differences just pop right out.”

Faye Schilkey
National Center for Genome Resources

to discover important trends and outliers in their data, thanks to point-and-click menus and customized workflows that eliminate the need for extensive programming skills or advanced statistical training.

Just point, click and you’re on your way. It’s powerful analysis made easy.

JMP Genomics brings the power of SAS to the study of genomics, whether you’re screening a genome for significant genetic markers, looking for meaningful patterns from expression studies or examining spectral data in a proteomics lab. As your genomic studies expand to new areas, you can explore new data in a familiar environment — without wasting time and money learning multiple software packages and manipulating data sets to move between them.

Beyond the comprehensive genomics menu, JMP Genomics includes full access to the extensive analysis and graphical features of the JMP platform, a powerful statistical tool for the rest of your laboratory needs. You can design experiments that are large yet efficient, and construct a variety of dynamically interactive graphics driven by a host of generic statistical methods. With JMP Genomics, you are never restricted to a small subset of specialized analysis tools.

JMP Genomics 4, our latest version, includes more new capabilities than any prior release. Based on new functionality from the cutting-edge SAS 9.2 and JMP 8 platforms, JMP Genomics 4 incorporates more tools for data exploration, enhanced visualization capabilities and a streamlined installation process. It provides new processes aimed at identifying cross-correlations between different types of measurements on the same samples (e.g., Copy Number and Expression). Additional graphical features, like Graph Builder, appeal very much to scientists with limited backgrounds in statistical analysis.

SAS developers worked with members of the global JMP Genomics user community to incorporate the most sought-after new features in JMP Genomics 4. These new and enhanced capabilities appeal to genomics scientists with a wide variety of research interests.

“You don’t have to be a whiz-bang programmer to get your answers. It’s very visual, but also very statistically accurate.”

Susan Dorsey, PhD, RN
University of Maryland, Baltimore

Expression

New Intermediate Workflows in JMP Genomics 4 build on Basic Workflows for exon and expression analysis, providing additional options requested by power users. These easy-to-use, pre-built workflows simplify quality control and statistical analysis of transcript and exon expression data sets, with options that include groupwise intensity filtering and point-and-click selection of normalization, analysis and pattern discovery methods. JMP Genomics includes plenty of flexibility for customized analysis by statisticians with sophisticated modeling requirements. The software uses file-based SAS Analytics to power popular normalization methods such as RMA and GCRMA, so users can apply these methods to much larger data sets than is possible with freeware or most other commercial tools.

Genetics

Huge data sets common to genomic research pose no problem for JMP Genomics. It provides an exceptional toolkit for statistical geneticists, with analysis options for data sets as large as 1.5 million SNPs for 15,000 individuals on a PC workstation. Enhanced workflows offer streamlined SNP and individual filtering capabilities. Sophisticated linear modeling capabilities for multiple trait types expand analysis options and are even available for analysis of genotypes from complex survey designs.

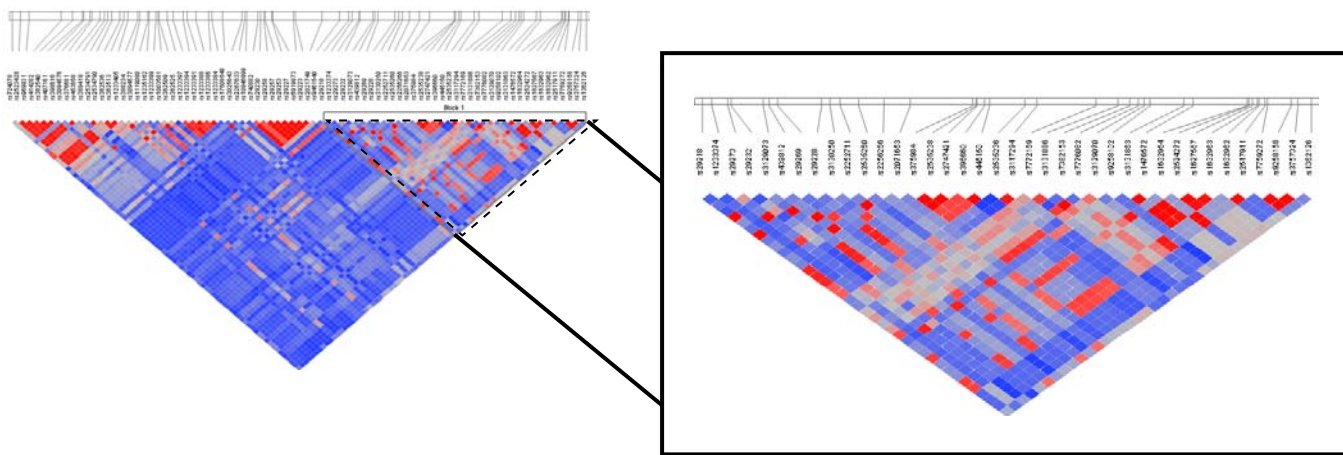
Copy Number

Easily explore copy number differences between groups or individuals with JMP Genomics. New visuals make it easy to pinpoint common regions of variation across multiple samples when compared to a reference group, and ANOVA-based methods offer statistical

verification to results. Users can assess the quality of copy number data using distribution analysis and principal components analysis to identify potential outliers and filter raw data based on statistical criteria.

Predictive Modeling

The breadth and depth of predictive modeling methods, as well as options for predictor filtering and cross-validation, distinguish JMP Genomics 4 from other genomics solutions. The software identifies key predictors within data sets incorporating multiple data types – SNP, expression, copy number – so users can build models using only the most significant biomarkers, and perform cross-validation with a choice of holdout methods and size options. Users of both JMP Genomics and SAS Grid Manager can even take advantage of experimental grid enablement of predictive modeling functions, saving time by parallelizing the analysis.



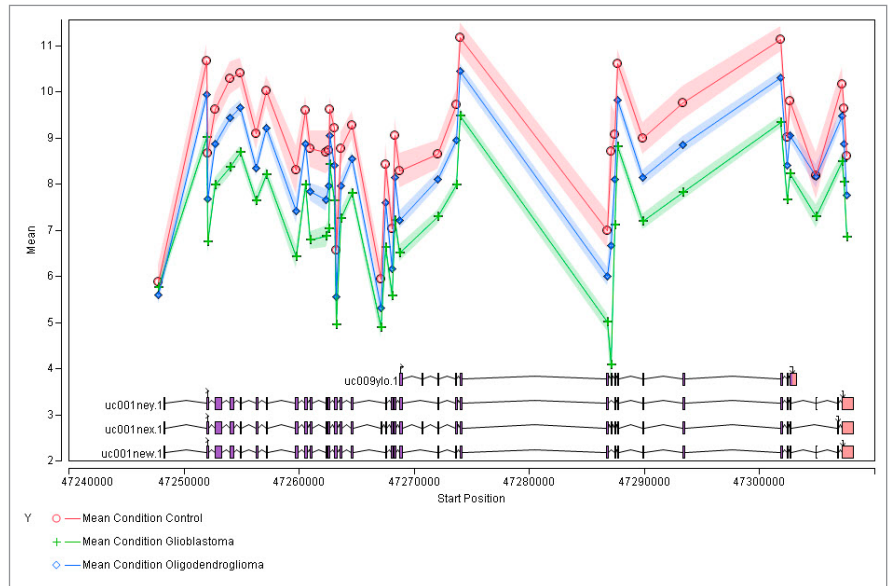
Examine patterns of linkage disequilibrium by position to identify genomic regions of greatest interest, then drill down by highlighting blocks.

Next-Generation Sequencing

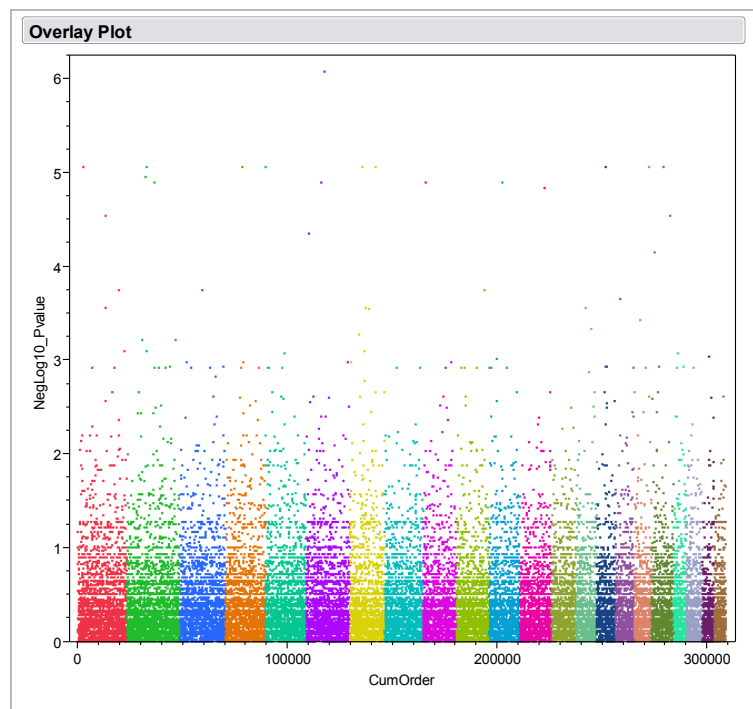
JMP Genomics frees you to follow your data wherever it leads instead of forcing your analysis down predefined paths. Whether you're working with data from traditional microarrays or transitioning to statistical analysis of data summarized from next-generation technologies, JMP Genomics sets itself apart with flexible statistical tools for quality control, normalization, pattern discovery and analysis. By partnering with leaders in next-generation sequencing, JMP Genomics provides sophisticated downstream statistical analysis capabilities to users of state-of-the-art sequence analysis pipelines from Illumina, GenoLogics and the National Center for Genome Resources.

JMP Genomics provides a cost-effective solution for flexible analysis of multiple data types from a variety of platforms, so you spend more time analyzing your data and less time reformatting it. As your analysis expertise grows, so does your ability to take full advantage of its powerful capabilities.

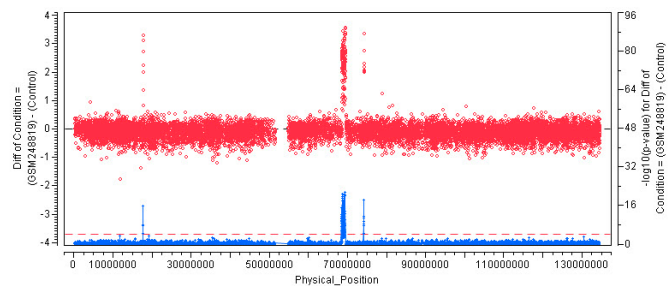
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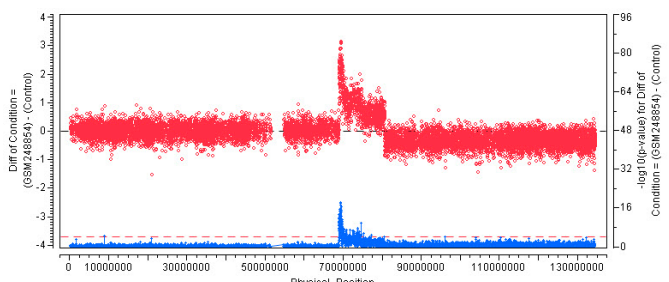
View statistical results from exon analysis in genomic context with overlay of new gene tracks. Here, mean intensity values for probe sets from an Affymetrix exon array are displayed for control samples and two different tumor types.



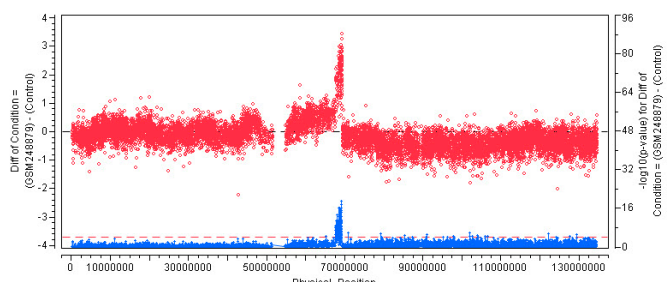
View p-values from statistical tests individually by chromosome, or create custom, multichromosome views.



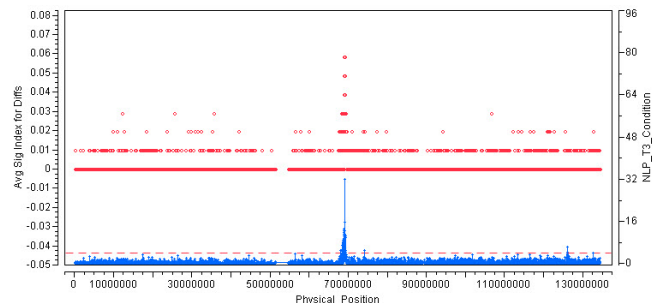
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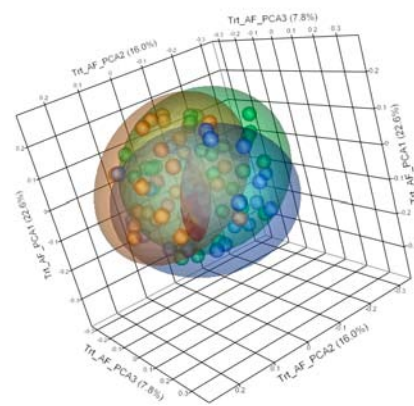
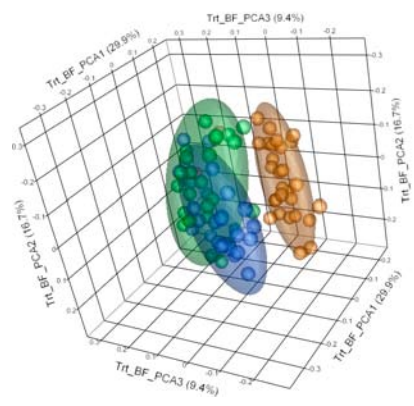


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“The ability to instantly recreate any analysis by opening the saved script means that I can determine exactly what analysis was done, even when the student forgets what options they selected. And JMP Genomics uses SAS data sets, allowing me to move easily between SAS and JMP Genomics tools.”

Lauren McIntyre, PhD
University of Florida Genetics Institute

Examine individual variations in copy number (left) or guide your search for shared regions with summary plots (right). JMP Genomics also allows examination of group-level differences.



See batch effects in your data and remove them prior to statistical analysis. JMP Genomics offers several different options for batch normalization. At left, samples collected in different batches group closely together, outweighing treatment effects. At right, the same samples are shown after batch effect removal.

See and explore your genomics data from every angle. Features in JMP® Genomics include:

Customized SAS Analytics running behind a JMP user interface:

- Offer point-and-click menus and options so users can get started quickly.
- Power robust data import, quality control, analyses, annotation and pattern discovery features using well-documented methods.
- Require no previous SAS programming experience.

The JMP software platform provides:

- Unparalleled flexibility for point-and-click creation of custom graphics: 2-D and 3-D scatterplots, parallel, overlay, contour and bubble plots.
- Dynamic, click-and-drag interface for visual exploration of data patterns with Graph Builder.
- Built-in JMP Scripting Language (JSL) and auto-generated graphics scripts that make it easy to capture and share important findings.
- Options for creating tailored dialogs for custom SAS analysis processes.

Interactive graphics generated automatically during analysis:

- Produce easy-to-understand summaries of large data sets.
- Link graphical features to data for point-and-click selection and easy creation of subset data tables.
- Can be queried dynamically to create tailored views of your data, using the JMP Data Filter or a variety of other selection tools.

Flexible workflows offer options for all users:

- Affymetrix CHP Wizard, which guides new users through analysis of CHP and ARR files through a simple, interactive interface.
- Basic Workflows for expression, exon, genetics and copy number.
- Intermediate Workflows for expression quality control and analysis.
- Workflow Builder, which offers complete control for expert users who wish to create their own custom workflows.

JMP Genomics imports data from a variety of formats, including:

- Illumina Bead Studio files for expression, SNP, genetic marker, copy number and other data types.
- Exon, whole transcript and 3' expression CEL and CHP files from GCOS and Affymetrix Command and Expression Console.
- CEL, CHP, LOHCHP and CNCHP files from Affymetrix Genotyping Console, and CNAT files.
- GenePix, QuantArray, one-color and two-color Agilent files.
- Genomics data contained within single text files or multiple text files.
- Text, Excel and comma-separated files.

Assess genetic markers and individuals to:

- Examine missing data patterns at the individual and genetic marker levels.
- Summarize characteristics of genetic marker data sets: allele and genotype frequencies, HWE, number of missing values, heterozygosity and diversity.
- Filter data sets based on marker characteristics prior to statistical analysis.
- Calculate and visualize linkage disequilibrium measures to zoom into interesting regions with interactive triangular plots.
- Generate distributions of categorical and continuous phenotypes.

Perform candidate-gene or whole-genome SNP analysis to:

- Analyze data sets as large as 1.5 million SNPs for 15,000 samples.
- Explore associations between genetic markers and binary or quantitative traits while adjusting for covariates.
- Visualize and correct for population structure prior to association tests with PCA for Population Stratification or Multidimensional Scaling.

Expand analysis options for marker data to incorporate:

- Detection of, and corrections for, relatedness among samples.
- Reconstruction of haplotypes and discovery of haplotype-trait associations.
- Selection of tagSNPs for haplotypes or linkage disequilibrium blocks.
- Computation of genetic distance matrices for individuals or populations.
- Single-marker, interval and composite-interval QTL mapping.

Analyze large microarray experiments with confidence to:

- Identify data quality issues and remove outlier arrays prior to statistical analysis.
- Normalize large data sets with a variety of options, including RMA and GCRMA for Affymetrix 3' expression arrays.
- Visualize intensity distributions and sample grouping patterns to explore the impact of experimental and technical effects.
- Normalize within and across arrays to remove confounding sources of variation and batch effects.
- Perform gene-by-gene modeling to discover statistically significant differences while correcting for multiple tests.
- Reveal biological insights with pattern discovery tools.

Apply advanced predictive modeling analysis tools to allow:

- Identification of reliable biomarkers from large, wide data sets.
- Assessment of multiple data types from different experiments.
- Customized predictor filtering during model construction.
- Cross-validation with adjustable hold-out and iteration options to enable comparison of relative performance across multiple models.
- Learning Curve analysis assessing the impact of sample size.

Assess copy number data sets to:

- Examine data quality with PCA and data distribution analysis.
- Analyze SNP intensities directly or importing copy number values generated by a variety of algorithms.
- Look for shared genomic areas that display statistically significant differences using ANOVA.
- Compare breakpoints identified by circular binary segmentation within and between samples.

Integrate statistical analysis into next-gen sequencing workflows to:

- Import sequence counts at the SNP, exon or transcript level generated by partner software from Illumina, the National Center for Genome Resources or GenoLogics.
- Utilize existing exon and expression workflows to examine mRNA-seq data.
- Test for association between variant alleles and traits.
- Perform cross-correlation analysis to relate sequence counts to other genomic measures.

Annotate results using public or proprietary sources of information to:

- Merge annotation information with statistical results.
- Download annotation and library files from Affymetrix NetAffx.
- Upload results to Ingenuity Pathways Analysis to seek points of interaction between SNP, gene and protein lists.
- Color KEGG pathways to identify sets of co-regulated genes.
- Create proportional-area Venn diagrams to assess overlap of up to five categories simultaneously.

“ When you’re going from looking at 10 genes to looking at thousands of genes, making biological sense of the results isn’t easy – it’s impossible to do if you don’t have the tools that help you easily visualize and explore the annotation of the results. JMP is great for that.”

Tom Juenger, PhD
University of Texas, Austin



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