Key Pathway Advisor on MetaCore accelerates your gene expression research and helps your organization spark innovation in patient stratification, disease mechanism reconstruction, target and biomarker discovery, and pharmacogenomics.

**COMPREHENSIVE PATHWAY ANALYSIS**

Gene expression analysis workflow is generally focused on identification of genes whose expression differs between two conditions (case/control studies) and pathways affected by defined genes. However, differential gene expression itself in turn could be the result of aberrant cellular signal transduction (due to environmental factors or gene variants with protein function impact).

KPA uses *causal reasoning network analysis* to predict Key Hubs transcriptional factors activity change (and their upstream regulators) that cause differential gene expression change. To perform this analysis, KPA leverages more than 100,000 manually annotated high-quality protein/RNA/compound interactions (including more than 34,000 transcriptional regulation interactions) stored in Thomson Reuters MetaBase™.

KPA also uses *synergy pathway analysis* to automatically define processes and pathway maps enriched with both experimentally defined and predicted Key Hubs gene sets in order to show connected, biologically relevant results.

**INTEGRATED 'OMICs DATA ANALYSIS**

KPA enables faster, more intuitive analysis of gene expression and associated gene variant data. Using MetaCore’s Genomic Analysis Tools (GAT)*, you can annotate your experimentally defined gene variant list and narrow it down to as many as 150 of the most promising protein function impact variants.

The resulting list can be uploaded into KPA with expression data in order to identify pathway impact and hypothesize about gain and loss of function effect (e.g., if a missense mutation occurred on a statistically significant active Key Hub, it may indicate gain of function effect).

**SIMPLE, INTUITIVE INTERFACE**

Simply import your data into the program and apply default analytical parameters. KPA automatically calculates a comprehensive workflow and delivers your results in structured reports:

- Online interactive reports – browse enrichment distributions, pathway maps, and significant molecular components.
- PDF report – shows static lists of enrichment distributions and top ten pathway map pictures overlaid with significant molecular components. Ideal for showing pathway snapshots and distributions to your colleagues.
- XLS report – contains expanded lists of molecular components and cross-references between them. Allows you to sort/filter all molecules on the basis of various molecular characteristics, narrowing down your data and making them appropriate for use in the other system biology tools and workflows.

**WHO CAN BENEFIT**

- Discovery biologists, including those without bioinformatics experience
- Bioinformaticians
- Pre-clinical scientists
- Translational researchers

KPA AND METACORE: INTUITIVE TOOLS FOR 'OMICs RESEARCH

Powered by Thomson Reuters MetaCore, Key Pathway Advisor (KPA) is a Web application that allows you to submit gene expression data and apply a comprehensive workflow that analyzes data in both upstream and downstream directions. Like a biological version of the GPS in your car, KPA doesn’t just visualize all possible pathways, it uses causal reasoning to point you to the most promising “routes,” analyzing the key pathways activated in your data and showing the key genes or proteins driving the distribution of your data.

KPA can be added to any MetaCore subscription.

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* GAT license required.
KPA ANALYSIS APPROACH

Predict Key Hubs and their activity on molecular interaction network which drive differential expression changes

Identify Key Pathways affected by differential expression and deregulated Key Hubs

KPA enables researchers to easily perform upstream and downstream analysis of gene expression – even if they have little to no bioinformatics experience.

With its intuitive interface and powerful analytics, the Key Pathway Advisor can support a range of researchers for disease understanding and discovery.

**DISCOVERY BIOLOGISTS**
**HELPING YOU TO:**
- Easily submit your data and generate an automatic comprehensive analysis even if you have no bioinformatics experience
- Combine gene expression and gene variant data
- Get a better understanding of aberrant signal transduction using detailed pathway maps with differentially expressed genes and Key Hubs highlighted
- Learn the biology of the disease you are working on

**BIOINFORMATICIANS**
**HELPING YOU TO:**
- Provide advanced causal reasoning and synergy pathway analyses to identify possible disease-driving molecules and aberrant signal transduction pathways
- Analyze connections between expression, gene variant, and predicted Key Hubs using a detailed XLS report
- Share analysis results with biologist colleagues using a PDF report with pathway images and descriptions

**TRANSLATIONAL RESEARCHERS**
**HELPING YOU TO:**
- Connect experimental research and clinical studies for insights
- Study detailed candidate disease drivers, biomarkers, and target relations, as well as their effect on signal transduction pathways

**PRE-CLINICAL SCIENTISTS**
**HELPING YOU TO:**
- Find novel biomarker and drug target candidates
- Identify drug response molecular changes and their signal transduction effect

To learn more, visit lsresearch.thomsonreuters.com/pages/solutions/1/metacore