QIAGEN® IPA®

For the analysis and interpretation of ‘omics data

QIAGEN Ingenuity® Pathway Analysis (QIAGEN IPA) is a web-based software application for the analysis, integration and interpretation of data derived from ‘omics experiments, such as RNA-seq, microarrays including miRNA and SNP, metabolomics, proteomics and small-scale experiments that generate gene and chemical lists. Powerful analysis and search tools uncover the significance of data and identify new targets or candidate biomarkers within the context of biological systems.

QIAGEN IPA goes beyond pathway analysis by:
- Identifying key regulators and activity to explain expression patterns
- Predicting downstream effects on biological and disease processes
- Providing targeted data on genes, proteins, chemicals and drugs
- Building interactive models of experimental systems

Insightful data analysis and interpretation

Data analysis and interpretation with QIAGEN IPA builds on the comprehensive, manually curated content of the QIAGEN Knowledge Base. Powerful algorithms identify regulators, relationships, mechanisms, functions and pathways relevant to changes observed in an analyzed dataset. Analytics go beyond pathway analysis to help you understand experimental results within the context of biological systems (Tables 1 and 2) and interactive tools allow detailed exploration of results, including comparisons across multiple analyses (Figure 1), discovery of novel biological connections and generation of testable hypotheses.

Figure 1. Interactive tools to explore and compare datasets. Trends and similarities across analyses can be quickly compared using heatmaps and interactive pathway graphics within the context of canonical pathways, analysis of downstream effects and examination of potential upstream regulators.

Table 1. Applications supported by IPA
- Target identification and validation
- Biomarker discovery
- Drug mechanism of action
- Drug mechanism of toxicity
- Disease mechanisms

Table 2. Experimental approaches supported by IPA
- RNA-seq
- Microarray
- miRNA
- mRNA
- qPCR
- Proteomics
- Metabolomics
Unlock insights and develop novel hypotheses

The Core Analysis in IPA quickly identifies relationships, mechanisms, functions and pathways relevant to a dataset. Upstream Regulator Analysis surfaces molecules, including miRNA and transcription factors, which may be causing observed gene expression changes (Figure 2) while Downstream Effects Analysis predicts downstream biological processes that are increased or decreased based on the analyzed data (Figure 3).

Integrating results about potential regulators and effects, the Regulator Effects tool highlights connections to create hypotheses about upstream triggers responsible for downstream phenotypic or functional outcomes. To further explore potential hypotheses, Molecule Activity Predictor (MAP) enables the user to interrogate subnetworks and canonical pathways by selecting a molecule of interest, indicating up- or downregulation, and simulating directional consequences on downstream molecules and the inferred activity upstream in the examined network or pathway (Figure 4).

Figure 2. Interactive analysis of plausible upstream regulators and networks. Insightful analyses predict upstream molecules, including miRNA and transcription factors, which may be causing observed gene expression changes.

Figure 3. Detailed examination of downstream effects. Detailed heatmaps highlight significant downstream biological processes that are increased or decreased based on gene expression results.
Advanced Analytics goes beyond immediate connections

Building on the Core Analysis, Causal Network Analysis, which is a component of QIAGEN IPA Advanced Analytics, uncovers multi-level causal relationships relevant to experimental data by expanding upstream analysis to include regulators that are not directly connected to targets in the analyzed dataset. Another Advanced Analytics component, BioProfiler, quickly surfaces molecules that are causally relevant to a disease or phenotype of interest, helping to identify potential therapeutic or toxicity targets, as well as associated known drugs and biomarkers.

Automatically compare your analysis with thousands of other analyses

QIAGEN IPA is also available with Analysis Match to automatically discover other QIAGEN IPA Core Analyses with similar (or opposite) biological results as compared to yours. These matches can help confirm your interpretation of the results or provide unexpected insights into underlying shared biological mechanisms among supposedly unrelated datasets. It scans the analyses you have created in your Project Manager, as well as thousands of other human and mouse expression analyses curated from public sources, seeking shared patterns of Canonical Pathways, Upstream Regulators, Causal Networks and Diseases and Functions (Figure 5).

The analyses included in Analysis Match were created in QIAGEN IPA from more than 50,000 highly curated and quality-controlled human and mouse disease and oncology datasets re-processed from SRA, GEO, Array Express, LINCS and TCGA. These datasets were generated by QIAGEN OmicSoft and are the “comparisons” found in DiseaseLand and OncoLand representing various contrasts between disease versus normal, treatment versus non-treatment and much more (Figure 6).
**Figure 5.** Analysis Match automatically displays analyses that are similar or dissimilar to your analysis. Analyses can be filtered by percentage similarity or by metadata keywords.

**Figure 6.** Selected analyses displayed as a heat map. This view enables you to drill down to explore the underlying drivers of the similarity or dissimilarity to your analysis.
Cause and effect for phosphoproteomics and metabolomics

QIAGEN IPA helps you understand the cause and effect of phosphorylation changes and endogenous metabolite concentration changes. You can predict which upstream regulators are responsible and whether those regulators are activated or inhibited, as well as visualize effects on downstream biological processes, diseases and established biological pathways (Figure 7).

Powerful tools for deep analysis of RNA-seq and miRNA data

Every feature of QIAGEN IPA is aimed at maximizing the impact of the information in an analysis so that the interpretation of a dataset is comprehensive. For example, the Isoform View displays expression data associated with each isoform from uploaded RNA-seq data in an intuitive graphical overview. Significantly regulated isoforms are listed in this view along with impacted functional protein domains and links to supporting publications (Figure 8).

Figure 7. Canonical Pathway with phosphorylation overlay. QIAGEN IPA leverages the known effects of phosphorylation to predict activity of phosphorylated proteins in your phosphoprotein dataset. For example, proteins with blue halos are proteins with increased phosphorylation which inhibits their activity.
IsoProfiler helps you determine which isoforms from your RNA-seq datasets have interesting biological properties relevant to your research project. For example, use this function to identify genes where isoforms are both upregulated and downregulated in the same dataset, which may have important functional consequences. Find cases of isoform switching – when the most highly expressed (highest FPKM) isoform for a gene differs between the experiment and the control, or explore all the cases where a gene expresses multiple protein-coding isoforms or is known to impact a disease or function. Explore the tissue-specific expression patterns of the transcripts in your dataset with fully integrated GTEx data (Figures 9–10).

Another powerful capability, the MicroRNA Target Filter, combines interactive filtering and comprehensive content to identify and prioritize miRNA-mRNA target pairings and provide insight into the biological effects of miRNA (Figure 11). Additionally, Upstream Regulator Analysis predicts miRNAs that may be regulating genes in an experimental dataset.
Figure 10. Graph of tissue expression across human tissues for all known splice variants for a gene. Use the fully reprocessed and integrated human GTEx expression data to identify transcripts with known tissue-specific expression.

Figure 11. Intuitive filtering tools to confidently identify mRNA targets. The miRNA Target Filter in QIAGEN IPA provides insights into the biological effects of miRNA, based on experimentally validated interactions from TarBase and miRecords, predicted miRNA-mRNA interactions from TargetScan, and miRNA-related findings from peer-reviewed literature.
Build custom pathways and gene or chemical list libraries

Create custom pathways with My Pathways and gene or chemical list libraries from a range of input data: Gene lists from QIAGEN IPA search results, existing QIAGEN IPA networks or canonical pathways, uploaded lists of targets or biomarkers, or imported pathways using XGMML, BioPax, SBML or GPML. Integrated tools guide the identification of upstream regulators or downstream targets of genes, enable layering of biological information or experimental data, and facilitate interrogation of hundreds of indexed subnetworks and canonical pathways to simulate effects and mechanisms of altered activity of target molecules (Figure 12). Highly interactive, these features afford intuitive exploration of connections between targets in a dataset to generate testable hypotheses and construct event-specific pathways such as:

- miRNA-mRNA target networks
- Transcriptional networks
- Phosphorylation cascades
- Protein-protein or protein-promoter interaction networks
- Chemical/drug effects on proteins

Figure 12. Powerful pathway editing tools to create interactive biological models. Transform your networks and pathways in QIAGEN IPA into publication-quality pathway graphics rich with color, customized text and fonts, biological icons, organelles and custom backgrounds. Expand and explore pathways using the high-quality content stored in QIAGEN IPA.
Explore pathways and interactions of interest

Path Explorer is an interactive tool that uncovers relevant relationships among genes of interest. By exploring these connections, the shortest paths between molecules associated with a disease or toxicity phenotype can be quickly identified, including access to supporting literature. Gene, Chemical & Pathway Search quickly generates and compares targeted lists of genes, druggable proteins, biomarkers and chemicals (Figure 13).

Leverage internal knowledge for a better understanding

QIAGEN IPA can incorporate your own or your institution’s internal data curation efforts for a disease or therapeutic area of interest. With the My Findings module, proprietary molecule-to-molecule relationships and molecule-to-disease or molecule-to-function relationships are uploaded to a secure, customer dedicated repository, making the content accessible throughout QIAGEN IPA. Any hypothetical or empirically demonstrated relationships can be imported or drawn and annotated on a new or existing pathway and then used in subsequent analyses to increase confidence in predicted upstream regulators, interaction or causal networks and downstream effects (Figure 14).

Seamless sharing and communication of results

QIAGEN IPA functions as a central platform for the analysis of biological data, generation of testable hypotheses and construction and visualization of molecular models of experimental systems. The communication and collaboration tools of QIAGEN IPA enable collaborative work on models and creation of interactive reports to share with colleagues. Collaborators with a license for QIAGEN IPA can be invited to share datasets and analyses or a customized Collaboration Workspace can serve as a shared results repository within or across institutions or consortia. QIAGEN IPA creates detailed summaries of analysis results that highlight the broader biological and therapeutic relevance of a particular pathway, gene or molecule list (including uploaded proprietary lists). The detailed tabular data and dynamic features of these reports enable fast decision making and hypothesis generation. Finally, Path Designer transforms networks and pathways into publication-quality colored graphics with species-specific nomenclature, biological icons and organelles, and with customized text, fonts and backgrounds. Path Designer pathways are intuitive, fully interactive graphics supported by the comprehensive content of QIAGEN IPA.
Understanding biological connections in a variety of applications

**Biomarker discovery**
Prioritize molecular biomarker candidates based on key biological properties and elucidate mechanisms linking markers to a disease or phenotype of interest.

**NGS/RNA-seq data analysis**
Streamline data analysis with rapid visualization and biological interpretation of expressed isoforms.

**Metabolomics**
Leverage critical biological context in QIAGEN IPA to overcome the challenges of analyzing metabolomics data and infer impacted cell function from metabolite lists.

**Proteomics and phosphoproteomics**
Uncover mechanistic links in complex proteomics data, identify potentially implicated regulators and predict impacted downstream processes or diseases.

**miRNA research**
Predict miRNAs regulating gene expression patterns and find mRNA targets based on content from miRBase, TargetScan and the QIAGEN Knowledge Base.

**Toxicogenomics**
Generate focused toxicity and safety assessments of candidate compounds and gain insight into pharmacological response and mechanism of action and toxicity.
The QIAGEN Knowledge Base is a data repository like no other. It organizes biological interactions and functional annotations created from millions of individually modeled relationships between proteins, genes, complexes, cells, tissues, drugs and diseases. These modeled relationships, or Findings, are manually reviewed for accuracy and include rich contextual details and links to original publications (Figure 14). The QIAGEN Knowledge Base enables access to relevant and substantiated knowledge from primary literature, as well as public and third-party databases (Tables 3–4), for the comprehensive interpretation of experimental results within the context of larger biological systems.

Table 3. Additional content sources in QIAGEN IPA

<table>
<thead>
<tr>
<th>Source</th>
<th>Description</th>
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<tbody>
<tr>
<td>Entrez Gene</td>
<td>GTEx Isoform-level Tissue Expression</td>
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<tr>
<td>RefSeq</td>
<td>HumanCyc metabolic pathway information</td>
</tr>
<tr>
<td>Conserved Domain Database (CDD)</td>
<td>BIND, DIP, MIPS, BioGRID, IntAct, Cognia protein-protein interactions</td>
</tr>
<tr>
<td>OMIM</td>
<td>clinicaltrials.gov</td>
</tr>
<tr>
<td>GWAS Database</td>
<td>European Medicines Agency</td>
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<tr>
<td>Gene Ontology (GO)</td>
<td>Pharmaceuticals and Medical Devices Agency</td>
</tr>
<tr>
<td>Human Metabolome Database (HMDB)</td>
<td>Mosby’s Drug Consult</td>
</tr>
<tr>
<td>Human Phenotype Ontology (HPO)</td>
<td>Goodman &amp; Gilman’s Pharmacological Basis of Therapeutics</td>
</tr>
<tr>
<td>GNF Tissue Expression Body Atlas</td>
<td>DrugBank</td>
</tr>
<tr>
<td>NCI-60 Cell Line Expression Atlas</td>
<td>Obesity Gene Map Database</td>
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Table 4. Identifiers supported in QIAGEN IPA

<table>
<thead>
<tr>
<th>Source</th>
<th>Description</th>
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<tbody>
<tr>
<td>Affymetrix®</td>
<td>Gene Symbol – human (Hugo/HGNC)</td>
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<tr>
<td>Affymetrix SNP ID</td>
<td>GenPept</td>
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<tr>
<td>Agilent®</td>
<td>GI Number</td>
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<tr>
<td>CAS Registry Number</td>
<td>Human Metabolome Database (HMDB)</td>
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<td>CodeLink</td>
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<td>GenBank</td>
<td>Applied Biosystems®</td>
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<td>miRBase (mature)</td>
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<td>Gene Symbol – rat (Entrez Gene)</td>
<td>miRBase (stemloop)</td>
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</table>

Species-specific Identifiers supported in IPA:
- PubChem CID
- Refseq
- UCSC (hg18)
- UCSC (hg19)
- Unigene
- Uniprot/Swiss-Prot Accession
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