**PATIKA: An informatics infrastructure for cellular networks**

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**THE PATIKA PROJECT** aims to develop methods and software tools for effective analysis of complex biological data at a functional level, consisting of following work-packages:

- Define an ontology for a comprehensive representation of cellular pathways.
- Develop software tools and construct an associated database using this ontology and provide an effective environment for pathway data integration, storage, access, visualization and analysis.
- Design methods for automatic population and annotation of the pathway database.
- Design methods for effective, advanced querying of the pathway database.
- Design methods for inferring pathway activity using temporal data such as gene expression data.
- Develop techniques for effective visualization of pathway and gene expression data.

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**Ontology**

We define an intuitive, comprehensive, uncomplicated representation of cellular networks.

**Basics**

**Bioentities**: actors of the cellular events, genetic (e.g., DNAs, proteins), chemical (e.g., ions), or physical (e.g., heat).

**Bioentity Interactions**: high level, imprecise relations: protein-protein interaction, transcriptional regulation or generic.

**States**: different forms of Bioentities via chemical modification (acetylated protein), localization (cytoplasmic ion), aberration (mutant gene), homomerization (dimers), etc.

**Transitions**: changes that states undergo.

**Interactions**: relations of states with transitions such as substrate, product, activator and inhibitor.

**Molecular Complexes**: Non-covalently bound clusters of molecules behaving as a single state.

**Cellular compartments**: part of the model.

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**Incomplete Information**

Since the data on cellular processes is incomplete, different levels of information may be available for certain events. On the left, it is unknown whether S4 activates either of two transitions.

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**Homologies**

B is transformed into B’ by activation of A. In the actual case, there are two A homologs, three B homologs and three B’ homologs.

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**Software**

A client/server architecture to provide access to PATIKA database through a state-of-the-art visual pathway editor has been implemented in pure Java™.

Advanced, graph theoretic queries may be performed through specialized GUIs, including queries by value or ID, neighborhood and shortest path querying.

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**Multi-User Environment**

Collaborative construction and concurrent modification issues are also addressed. While a user is working on a pathway locally, others might change its topology or properties in the database.

Checks for up-to-date status of graph objects result in each graph object being color-coded with respect to its status:

- *Blue*: Up-to-date
- *Red*: Out-of-date
- *Yellow*: Local
- *Green*: Locally Modified

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**Automated Layout**

Specialized algorithms for layout of cellular pathways produce aesthetically pleasing drawings.

**Gene Expression Analysis**

Please visit POSTER F-67 for details of PATIKA’s Microarray Data Analysis Facilities.

Support for analysis of gene expression data including a pathway activity inference method using gene expression data has been implemented.

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**Previous Contributors**

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