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Data integration, pathway analysis and mining for systems biology
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Doctoral dissertation for the degree of Doctor of Science in Technology to be presented with due permission of the Faculty of Information and Natural Sciences, The Aalto University School of Science and Technology, for public examination and debate in Auditorium F239a at Aalto University (Otakaari 3, Espoo, Finland) on the 14th of May, 2010 at 12 noon.
Abstract

Post-genomic molecular biology embodies high-throughput experimental techniques and hence is a data-rich field. The goal of this thesis is to develop bioinformatics methods to utilise publicly available data in order to produce knowledge and to aid mining of newly generated data. As an example of knowledge or hypothesis generation, consider function prediction of biological molecules. Assignment of protein function is a non-trivial task owing to the fact that the same protein may be involved in different biological processes, depending on the state of the biological system and protein localisation. The function of a gene or a gene product may be provided as a textual description in a gene or protein annotation database. Such textual descriptions lack in providing the contextual meaning of the gene function. Therefore, we need ways to represent the meaning in a formal way. Here we apply data integration approach to provide rich representation that enables context-sensitive mining of biological data in terms of integrated networks and conceptual spaces. Context-sensitive gene function annotation follows naturally from this framework, as a particular application. Next, knowledge that is already publicly available can be used to aid mining of new experimental data. We developed an integrative bioinformatics method that utilises publicly available knowledge of protein-protein interactions, metabolic networks and transcriptional regulatory networks to analyse transcriptomics data and predict altered biological processes. We applied this method to a study of dynamic response of *Saccharomyces cerevisiae* to oxidative stress. The application of our method revealed dynamically altered biological functions in response to oxidative stress, which were validated by comprehensive *in vivo* metabolomics experiments. The results provided in this thesis indicate that integration of heterogeneous biological data facilitates advanced mining of the data. The methods can be applied for gaining insight into functions of genes, gene products and other molecules, as well as for offering functional interpretation to transcriptomics and metabolomics experiments.
Preface

This research work has been carried out at the Quantitative Biology and Bioinformatics (QBIX) Group at VTT Technical Research Center of Finland (VTT). I thank my advisor, Research Professor Matej Orešič, for his excellent guidance throughout this work and the broad range of opportunities he provided me to work. I thank my supervisor, Professor Kimmo Kaski, Head of the Centre of Excellence, Department of Biomedical Engineering and Computational Science (BECS) of Helsinki University of Technology (TKK; called Aalto University School of Science and Technology since January 2010), for accepting me as a doctoral student and for his excellent support during this period. I thank my co-advisor, Dr. Jaakko Hollmén, Chief Research Scientist, Department of Information and Computer Science, TKK, for his cooperation and excellent advise during thesis writing.

The QBIX group has been reorganised into two teams since January 2009: Metabolomics and Biosystems Modelling; I belong to the latter. I thank the leaders of these two teams Dr. Tuulia Hyötyläinen and Dr. Marko Sysi-Aho, respectively, for their continued support to my work. I thank Technology Manager Dr. Richard Fagerström for his support. He has readily offered help and advise on a variety of issues. I am grateful to Dr. Juha Ahvenainen, former Vice President (R&D) (currently Vice President, Customer Management), Dr. Anu Kaukovirta-Norja, Vice President (R&D), and Research Professor Hans Söderlund for the excellent working facilities. I thank Mr. Atul Khanna, chairman of Tooltech Software Ltd, for his support during the early years of this work.

I thank all researchers who co-authored with me. Without their contributions to my publications this work would have been incomplete. I thank Mr. Erno Lindfors for his enthusiasm and excellent work, which have been crucial for the success of my work. Working with him has always been easy and effective. Working with Dr. Vidya Velagapudi has provided insights into biology. Her positive attitude and perseverance made working with her a motivating experience. I also thank Mr. Jussi Mattila, Dr. Jyrki Löjtönen, and other members of the Signal and image processing team at VTT Tampere for their collaboration.

The working environment at VTT has been conducive for the sort of interdisciplinary research presented in the thesis. The interdisciplinary knowledge of the members of QBIX group has created a knowledgeable workspace, and I thank each and every individual of the group. Working at QBIX group has also been full of fun. I thank Mr. Laxman Yetukuri, Dr. Catherine Bounsaythip, Mr. Han Zhao, Ms. Sandra Castillo, and Mr. Pekka Savolalhti for their wonderful company. A lot of discussions with them, which included topics such as culture, life, ethics, or simply fun, at the coffee table have been revitalising. Technical discussions with my present and past team mates as well as many other researchers and faculty members at VTT and TKK, have been helpful at different instances.

I thank Professor Garry Wong and Docent Tero Aittokallio for the pre-examination of the thesis. Their valuable comments were insightful and enhanced the presentation of the thesis. I thank all my colleagues who read my thesis and provided their valuable comments. I thank Dr. Kaija Virolainen, Ms. Anna-Kaarina Hakala, and Ms. Sirpa Nygren for their help with many practicalities.

Friends at work as well as outside, including a number of Indian friends, made the social life
in Finland very smooth and lively. They are, at times, unique sources of some important practical information related to living in Finland. As many of these friends are researchers in biology, bioinformatics or computer science, I also had many useful technical discussions with them.

When I was in my masters at Indian Institute of Technology (IIT) Kharagpur, my friend Dr. Murthy Chavali, then a graduate student, has introduced to me the topic of DNA computing, which subsequently led my interest to the broad variety of applications at the interface of computational sciences and biology. Support provided by Dr. G. P. Raja Sekhar at that time is also gratefully acknowledged.

I thank my wife, Subhadevi Attili, for her unconditional love and trust. I thank my parents and all other family members for their love, affection, and support. What I am today is the net result of my actions in the past and how they were supported by family, friends, teachers, and possibly even people I do not know. Although the names of all of those who played important roles in my life are not mentioned, they are already part of what I am today, and I am indebted to their support.
List of publications

This thesis consists of introductory part and the following publications


Roman numerals are used to refer to these articles.

**Author’s contributions**

The research presented in this thesis consists of two themes: data integration and context-sensitive visualisation (*Articles I–IV*), and dynamic topology of integrated networks (*Article V*). Author’s contributions to each of these papers have been summarised below. Contributions of other authors are mentioned when necessary.

*Article I* presents a heterogeneous data integration system for the integration and visualization of multiple types of biological interaction data and demonstrates its usage with two applications: multiple pathway retrieval and protein neighbourhood search. The author designed the system, performed data modelling, developed the schemas, obtained relevant data, developed parsers, implemented the database back-end and developed the semantics for the integration of heterogeneous
biological entities, and provided guidance to the development of the overall system. Author wrote the first draft of the paper which was then improved with the contributions of other authors. Erno Lindfors (EL) implemented the integration of biological entities, representation of integration results using complex networks, the Sammon’s mapping algorithm, Graphical User Interface client software and wrote these parts of the manuscript.

Article II presents the data integration system with addition of new methods. It presents three applications: system-wide metabolic network and the study of its topological properties, exploration of properties and relationships of a specific set of proteins, and combined visualization and exploration of a Type 1 Diabetes gene expression data in mouse together with related pathways and ontologies. Author developed the ideas concerning integration of gene expression data to networks and performed the analyses and wrote the manuscript. EL developed the ideas for topology study and performed analyses and implemented middle-tier and user interface and wrote some parts of the manuscript. The author and EL equally contributed to this work. Jarkko Miettinen (JM) implemented projection algorithms and improved the client and middle-tier software design and code.

Article III presents the application of the data integration framework for context-dependent visualization of protein function using network representation and nonlinear projection methods based on Curvilinear Distance Analysis. The author performed the analyses and wrote the methods and results parts of the paper which was then improved by all authors under the coordination of Matej Orešič (MO).

Article IV presents the details of the networks and the distance metrics. It demonstrates with experiments how judicious use of various distance functions can allow emergence of context. The author mainly provided biological details and data, and contributed to the manuscript which was mainly written by Catherine Bounsaythip (CB) and EL.

Article V presents an integrative bioinformatics method called Topological Enrichment Analysis of Functional Subnetworks (TEAFS). Author developed the main ideas of the method, implemented some parts of the method, performed the analyses and wrote the manuscript. Vidya R. Velagapudi (VRV) performed in vivo metabolic profiling experiments and data analysis, and wrote the experimental methods and biological details in the manuscript. Author and VRV equally contributed to this work. EL performed biological network construction, provided help with topological analysis, implemented the statistical test and contributed to the writing. Eran Halperin (EH) provided ideas for the statistical test. MO initiated and coordinated the project and contributed to the writing of the paper.

Goals of the thesis

The goals of this thesis are

- to develop integrative bioinformatics tools and methods to integrate heterogeneous biological data
- to produce knowledge, e.g., testable hypotheses, from these tools in an exploratory manner
- to apply these tools and methods in mining newly generated experimental data

Contributions presented in this thesis

Following contributions have been made through this thesis.
PUBLICATIONS

- A technical framework for storage and retrieval of biological data achieved through XML and relational databases.

- The technique of data traversals, achieved by curation of maps database, as the basis for heterogeneous data integration.

- A platform called MegNet for enabling context-sensitive mining of heterogeneous biological data through the usage of conceptual spaces as the framework for knowledge representation.

- A new method called Topological Enrichment Analysis of Functional Subnetworks (TEAFS) for studying the dynamic activity of biological process modules, and its application to studying the dynamic response of Saccharomyces cerevisiae to Oxidative stress.
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<th>Expansion</th>
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<tr>
<td>A</td>
<td>absent</td>
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<tr>
<td>API</td>
<td>Application Programming Interface</td>
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<tr>
<td>BioGrid</td>
<td>General Repository of Biological Interaction Datasets</td>
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<td>BIND</td>
<td>Biomolecular Interaction Database</td>
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<tr>
<td>CCA</td>
<td>Curvilinear Component Analysis</td>
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<td>CDA</td>
<td>Curvilinear Distance Analysis</td>
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<tr>
<td>cDNA</td>
<td>Complementary DNA</td>
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<td>DBMS</td>
<td>Database Management System</td>
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<tr>
<td>DIP</td>
<td>Database of Interacting Proteins</td>
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<td>DNA</td>
<td>Deoxyribonucleic Acid</td>
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<tr>
<td>DOM</td>
<td>Document Object Model</td>
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<tr>
<td>DTD</td>
<td>Document Type Definition</td>
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<td>EC</td>
<td>Enzyme Commission</td>
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<td>EMBL</td>
<td>European Molecular Biology Laboratory, EMBL Nucleotide sequence database</td>
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<tr>
<td>FDR</td>
<td>False Discovery Rate</td>
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<td>GC</td>
<td>Gas Chromatography</td>
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<td>GDS</td>
<td>GEO Data Set</td>
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<td>GEO</td>
<td>Gene Expression Omnibus</td>
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<td>GI</td>
<td>GenInfo sequence identifier</td>
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<tr>
<td>GNEA</td>
<td>Gene Network Enrichment Analysis</td>
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<td>GO</td>
<td>Gene Ontology</td>
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<td>GSEA</td>
<td>Gene Set Enrichment Analysis</td>
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<td>GPL</td>
<td>GEO Platform</td>
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<td>GSM</td>
<td>GEO Sample</td>
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<td>GSE</td>
<td>GEO Series</td>
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<tr>
<td>HPLC</td>
<td>High Performance Liquid Chromatography</td>
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<tr>
<td>JDBC</td>
<td>Java Database Connectivity</td>
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<tr>
<td>KEGG</td>
<td>Kyoto Encyclopedia of Genes and Genomes</td>
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<td>KGML</td>
<td>KEGG Markup Language</td>
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<tr>
<td>M</td>
<td>Marginal</td>
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<td>MINT</td>
<td>Molecular Interaction Database</td>
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<td>MM</td>
<td>Mismatch</td>
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<tr>
<td>mRNA</td>
<td>messenger RNA</td>
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<tr>
<td>MS</td>
<td>Mass Spectrometry, Mass Spectrometer</td>
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### Abbreviations

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Expansion</th>
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<tr>
<td>NOD</td>
<td>Non Obese Diabetic</td>
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<tr>
<td>P</td>
<td>Present</td>
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<td>PDB</td>
<td>Protein Data Bank</td>
</tr>
<tr>
<td>PIR</td>
<td>Protein Information Resource, PIR-international protein sequence database</td>
</tr>
<tr>
<td>PM</td>
<td>Perfect Match</td>
</tr>
<tr>
<td>PSI</td>
<td>Protein Standards Initiative</td>
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<tr>
<td>PSI-MI</td>
<td>PSI-Molecular Interaction format</td>
</tr>
<tr>
<td>RDF</td>
<td>Resource Description Framework</td>
</tr>
<tr>
<td>RNA</td>
<td>Ribonucleic Acid</td>
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<tr>
<td>SAGE</td>
<td>Serial Analysis of Gene Expression</td>
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<tr>
<td>SAX</td>
<td>Simple API for XML</td>
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<tr>
<td>SBML</td>
<td>Systems Biology Markup Language</td>
</tr>
<tr>
<td>SOAP</td>
<td>Simple Object Access Protocol</td>
</tr>
<tr>
<td>SOFT</td>
<td>Simple Omnibus Format in Text</td>
</tr>
<tr>
<td>SQL</td>
<td>Structured Query Language</td>
</tr>
<tr>
<td>TCA cycle</td>
<td>The citric acid cycle or The tricarboxylic acid cycle</td>
</tr>
<tr>
<td>TEAFS</td>
<td>Topological Enrichment Analysis of Functional Subnetworks</td>
</tr>
<tr>
<td>TRANSFAC</td>
<td>Database of Transcription Factors</td>
</tr>
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<td>TRANSPATH</td>
<td>Database of Signal Transduction Pathways</td>
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<tr>
<td>TrEMBL</td>
<td>Translated EMBL</td>
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<tr>
<td>T1D</td>
<td>Type 1 Diabetes</td>
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<tr>
<td>UMLS</td>
<td>Unified Medical Language System</td>
</tr>
<tr>
<td>UniProt</td>
<td>Universal Protein Resource</td>
</tr>
<tr>
<td>UPLC</td>
<td>Ultra Performance Liquid Chromatography</td>
</tr>
<tr>
<td>XML</td>
<td>eXtensible Markup Language</td>
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<tr>
<td>XPath</td>
<td>XML Path Language</td>
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<tr>
<td>XQuery</td>
<td>XML Query Language</td>
</tr>
<tr>
<td>XSD</td>
<td>XML Schema Definition</td>
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<tr>
<td>Y2H</td>
<td>Yeast Two-Hybrid method</td>
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Chapter 1

Introduction

Systems biology aims to study biological systems at system level \cite{Kitano2002}. Integrative bioinformatics provides tools for systems biology. The necessary goal of integrative bioinformatics is to provide platforms and methods for carrying out systems biology analysis. \cite{Ge2003} noted that, “high-throughput data integration is needed in systems biology approaches”. This thesis contributes methods for data integration, visualisation, mining, and study of network dynamics, which are important elements of systems biology Figure 1.1.

Post genomic biology is characterized by rapid accumulation of information and thus biological research can be viewed as information science \cite{Schena1998}. Information is increasing more swiftly than humans resources such as time and cognitive processing power. This forces humans to narrow their focus in processing the information, which in turn causes diminishing awareness. As an example, the literature documenting Raynaud’s disease and fish oils were mutually isolated in the sense that the authors and readers of one literature were not aware of the other. Much later, a cure for Raynaud’s disease with dietary fish oils was found, using a literature-based approach covering both areas \cite{Swanson1986}. This demonstrates the power of data integration. Thus the methods for efficient retrieval and presentation of results can benefit researchers.

Cytoscape \cite{Shannon2003, Killcoyne2009} is a general network visualization, data integration, and analysis software, which has been mainly developed with the modelling requirements of systems biology in mind. The core of Cytoscape mainly features powerful layout algorithms for visualising networks and is quickly becoming a de facto standard for the visualisation of biological networks, while its flexible plug-in architecture brings the real power via community-based development of useful plug-ins. However, Cytoscape does not offer data management capabilities. Biological data management and integration has also attracted significant amount of research \cite{Lacroix2003}. There are three major approaches in this area: data warehousing approach, distributed or federated approach, and mediator approach. The data warehousing approach involves assembling data sources into a centralised system with a global data schema and an indexing system for integration and navigation. In the federation approach, underlying data sources remain autonomous, and the federated system maintains a common data model and makes use of schema mapping to translate heterogeneous source database schema to the target schema for integration. The mediator approach introduces a mediator layer, a collection of software components performing integration, to decouple the underlying heterogeneous distributed data sources and the client layer. Many popular and important biological data integration systems are discussed in \cite{Lacroix2003}. But these systems are limited to providing web based access to multiple reference databases.
CHAPTER 1. INTRODUCTION

Figure 1.1: Systems biology can be viewed as a combination of omics technologies, data integration, analysis, mining, and modelling, often involving use of these techniques iteratively over hypothesis driven systematic experimental design to gain increased understanding of the structure and dynamics of the biological systems. High throughput omics technologies provide the measurements for systems biology. Integrative bioinformatics starts with the integration of multiple data sets from one or more omics and also possibly from multiple organisms, and forms the basis for systems biology analysis. Systems biology analyses include data mining, visualisation, biological networks, and dynamic modelling. The new knowledge generated by these analyses would enable us to build quantitative models. The hypothesis generated by the analysis of these models drive the design of more experiments to gain increased understanding of the biological systems.

The availability of high-throughput data collection techniques of modern biology introduce some new problems. First, there are many false positive findings and reproducibility is poor i.e., a biological sample analysed by using a single experimental technique at different times or laboratories often lead to unidentical results (Ge et al. 2003; Ein-Dor et al. 2005; Irizarry et al. 2005; Tan et al. 2003). Meta-analyses of multiple data sets or evidences from multiple types of biological experiments may improve the statistical power of the analysis. Second, various types of biological activities or interactions within an organism and between an organism and its environment do not happen in isolation. Biological function is a net result of simultaneous activities and interactions of various types (Kanehisa and Bork 2003; Ideker et al. 2001a,b; Ge et al. 2003; Papin and Palsson 2004). Thus the integrated modelling of the biological systems is very important.

This thesis addresses these problems. First, it presents bioinformatics methods to visualise biological interactions of different types in an integrated manner. Next, it presents methods to facilitate advanced context-sensitive mining of the integrated network data. Finally, it presents
mining of dynamic topological changes of functional modules in integrated networks in response to specific interventions. The methods presented in this thesis can be applied for prediction of gene, gene product or metabolite functions, and to associate experimental phenotypes with genotypes i.e., for interpretation of transcriptomics and metabolomics data in terms of molecular level patterns or changes.

**Organisation of the thesis**

The research presented in this thesis consists of integrative bioinformatics methods under two themes. The first theme is that of a bioinformatics software platform which embodies methods for integration of heterogeneous biological data: a variety of interaction, annotation, and molecular measurement data (chapter 2). The methodological details of this software platform are presented in section 3.1. The platform achieves integration across different biological data types using data traversals (section 3.1.4). The resulting information forms the basis for context-sensitive data mining (section 3.1.6), which draws on the concepts from the theory of conceptual spaces. Some results of exploratory data mining using this platform are presented in section 4.1. The second theme is the study of dynamic changes of functional modules in an integrated network (section 3.2). An application of this strategy with a study of dynamic topological response of oxidative stress in *Saccharomyces cerevisiae* is presented in section 4.2. Summary and concluding remarks follow in chapter 5.
Chapter 2

Biological data

Cells consist of different types of biomolecule. Reductionist approach of molecular biology deals with studying the properties and roles of individual molecules and produced large amounts of useful knowledge. A discrete biological function, however, can not be attributed to an individual molecule, but to a complex web of interactions between a set of molecules. Therefore, describing biological systems requires explaining how they arise from interactions among components in the cell (Hartwell et al., 1999; Ge et al., 2003). Availability of complete genome sequences of several organisms (Goffeau et al., 1996; Blattner et al., 1997; Adams et al., 2000; Lander et al., 2001; Venter et al., 2001; Waterston et al., 2002) has opened doors for development of high-throughput omics technologies.

Figure 2.1 provides an overview of some omics experimental techniques for measuring different biological data types, which are elaborated in this chapter. Gene expression microarrays (Schena et al., 1995; Lockhart et al., 1996) and Serial Analysis of Gene Expression (SAGE) (Velculescu et al., 1995), which enable us to measure the abundances of thousands of gene transcripts simultaneously, have empowered the first omics discipline known as transcriptomics. The development of proteomics, large scale study of proteins, followed (Patterson and Aebersold, 2003). Mass spectrometry based protein purification (Rigaut et al., 1999; Aebersold and Mann, 2003) and yeast two hybrid analysis (Ito et al., 2001) have enabled investigating protein-protein interactions in high-throughput manner (Shoemaker and Panchenko, 2007a). The omics technologies have contributed to rapid accumulation of knowledge such as gene and gene product annotations, biomolecular interactions. Simultaneously, the availability of genome sequences also enabled development of computational algorithms for sequence analyses (Altschul et al., 1990; Thompson et al., 1994; Durbin et al., 1998), which also helped to rapidly annotate new sequence data, and predict the structure and interactions (Shoemaker and Panchenko, 2007b; Marcotte et al., 1999). Finally, text mining (Skusa et al., 2005) also helped to retrieve important molecular interaction information, and careful manual literature curation led to higher quality information as compared to high-throughput data (Reguly et al., 2006; Matys et al., 2006). In the spirit of genomic data sharing, many types of the biological data have been made accessible through world wide web.

These web-accessible databases and their associated search and mining tools are primary resources serving thousands of biology researchers worldwide. These tools allow researchers to effectively mine the databases and answer one's biological questions (Kanehisa and Bork, 2003). The databases cover a wide range of information including literature, sequences and annotations of genes and gene products, and a variety of molecular interactions such as biochemical reactions, transcriptional regulatory interactions, signal transduction pathways, to name a few. This chapter introduces some of the commonly used databases for microarray gene expression profiles, protein-protein interactions, and more.
Figure 2.1: Various types of data in molecular biology and the experimental techniques employed for obtaining the data (by no means exhaustive). High-throughput techniques capable of measuring all or at least a large number of components (several hundreds to thousands) simultaneously are known as omics techniques. Generally the word interactome is used to refer to the collection of protein-protein interactions, but in this thesis we also include other types such as metabolic networks and transcriptional regulatory interactions into the interactome category for convenience, as we repeatedly refer to all these types of interactions.
interactions, metabolic interactions, transcriptional regulatory interactions, and signal transduction networks.

GenBank (Benson et al., 2000, 2010) is a sequence database that stores all known DNA sequences gathered by direct submission of sequence data from individual laboratories and from large-scale sequencing projects. There are two other major DNA sequence databases namely EMBL nucleotide sequence database (Stoesser et al., 1999; Kulikova et al., 2007), and DDBJ (Tateno et al., 2002; Kaminuma et al., 2010), and the data among these three databases are synchronised. The Ensembl project (Hubbard et al., 2007; Flicek et al., 2010) offers an integrated source of genome sequences and annotations for a comprehensive set of chordate genomes with a particular focus on human, mouse, rat, zebrafish etc. The University of California Santa Cruz (UCSC) Genome Browser Database (Karolchik et al., 2003; Rhead et al., 2010) is a source for genome sequence and annotation data. UCSC Genome Browser, is a tool associated with the UCSC database that provides rapid visualization and querying of the data. The annotations provided by these genome databases include mRNA and expressed sequence tag (EST) alignments, gene predictions, cross-species homologies, high-level maps, single nucleotide polymorphisms (SNPs) and so on. Besides these general genomic databases, there are organism specific genomic databases for model organisms. The Saccharomyces Genome Database (SGD) (Cherry et al., 1998; Engel et al., 2010) is a database for the molecular biology and genetics of the yeast Saccharomyces cerevisiae that provides functional annotations, mapping and sequence information, protein domains and structure, expression data, mutant phenotypes, physical and genetic interactions and the primary literature from which these data are derived. FlyBase (Gelbart et al., 1997; Drysdale et al., 2008) is a database of genetic and genomic data concerning fruit flies of which Drosophila melanogaster is an extensively studied model organism. FlyBase is populated with information from a variety of sources ranging from large-scale genome projects to the primary research literature. FlyBase provides access to information on gene models, molecular classification of gene product functions, mutant phenotypes, mutant lesions and chromosome aberrations, gene expression patterns, transgene insertions, and anatomical images. WormBase (Stein et al., 2001; Harris et al., 2010) is a central data repository for nematodes of which Caenorhabditis elegans is an extensively studied model organism. WormBase includes genomic sequences, gene predictions and orthology assignments from a range of related nematodes and relies on manual curation of information from the corpus of C. elegans literature.

Transcription factors are proteins that are vital for the transcriptional regulation of gene expression. A transcription factor has a DNA binding domain which can bind to a particular region in the DNA sequence of a gene, called the binding site, and helps in enhancing or inhibiting the expression of the gene (Latchman, 1997). TRANSFAC database (Wingender et al., 2000; Matys et al., 2006) primarily provides information about entities involved in the transcriptional regulation such as transcription factors, binding sites and genes among a variety of other related information.

Living cells interact with their environment by exchanging a variety of signals. Signaling pathways of the receiver cells forward the signals to the nucleus through cascades of interactions and trigger the appropriate adaptation of the genetic program. The TRANSPATH database (Schacherer et al., 2001; Krull et al., 2006) provides information about signal transduction pathways involved in the transcriptional regulation of gene expression via regulating the activity of the transcription factors.

The Universal Protein Resource (UniProt) (Bairoch et al., 2005; Apweiler et al., 2010) provides information about protein sequences and functional information. The central database in Uniprot, termed UniProt Knowledgebase, provides accurate, consistent and rich sequence and functional annotations and consists of two sections: UniProt/Swiss-Prot and UniProt/TrEMBL. UniProt/Swiss-Prot consists of manually curated protein functional information, resulting from literature information extraction and curator-evaluated computational analysis. UniProt/TrEMBL consists of protein
sequences translated from EMBL gene sequences [Stoesser et al., 1999, Kulikova et al., 2007] and annotated with computational annotation tools, pending manual curation.

Database of Interacting Proteins (DIP) [Xenarios et al., 2002] is a database of manually curated protein-protein interactions. A curator enters each interaction entry into the database after manually reading the publication reporting an experimentally verified interaction. This is intended to be a comprehensive and integrated tool for browsing and efficiently extracting information about protein interactions and interaction networks in biological processes. DIP provides access to combined information from multiple observations and experimental techniques, from multiple organisms, as well as to networks of interacting proteins. Each interaction entry in the DIP database contains information about the protein domains and range of amino acids involved in the interaction, and the corresponding experiments. The interactors are identified by Swissprot [Apweiler et al., 2010], PIR [Barker et al., 1998], or GenBank [Benson et al., 2000] accession numbers and each interactor entry contains information about the organism, function, superfamily, cellular location and so on.

The Biomolecular Interaction Database (BIND) [Bader et al., 2003] stores pairwise interactions between biological ‘objects’ which could be protein, RNA, DNA, molecular complex, small molecule, photon (light) or gene. Moreover, it contains higher level functional structures called molecular complexes and pathways which are collections of the pairwise interactions with some additional data. The minimum amount of information required to define an interaction is a description of the interacting objects and a publication reference to PubMed [Wheeler et al., 2007]. Data in BIND is primarily obtained via submissions of individual contributors across the world. However, it also incorporates interaction data imported from other databases such as PDB, and a number of large-scale cell mapping studies using yeast two hybrid, mass spectrometry, genetic interactions and phase display. SeqHound is a data integration system [Michalickova et al., 2002] that provides extensive C, C++, and Perl application programming interfaces (API) for data in BIND. SeqHound system provides also functions to link the biological objects with other biological databases in public domain.

The Molecular Interaction database (MINT) [Zanzoni et al., 2002, Ceol et al., 2010] stores information about experimentally verified molecular interactions extracted from publications from peer-reviewed journals. The main focus is on physical interactions between proteins. Genetic or computationally inferred interactions are not included in MINT. MINT includes an additional database called HomoMINT [Persico et al., 2005], which is a database of interactions between human proteins inferred from interactions between orthologous proteins in model organisms. A large number of MINT data comes from large scale, genome wide experiments, although curating data from low-throughput published experiments is given emphasis. Each interaction entry contains reference to Swiss-Prot/TrEMBL protein accession number [Apweiler et al., 2010] for the interactor and contains the experimental information and pubmed reference [Wheeler et al., 2007] for describing the experimental conditions and other properties of the interaction.

BioGrid [Stark et al., 2006, Breitkreutz et al., 2008] is a database of protein and genetic interactions. It is aimed to be a generic repository providing comprehensive information on molecular interactions in several organisms such as Saccharomyces cerevisiae, Drosophila melanogaster, Caenorhabditis elegans, and Homo sapiens. It currently hosts protein-protein interaction data from high-throughput experiments such as yeast two-hybrid (Y2H) method [Ito et al., 2001] and mass spectrometry analysis of purified protein complexes [Rigaut et al., 1999]. Additionally, the BioGrid team also compiles interaction data by extensive manual curation of literature. Literature curated data for Saccharomyces cerevisiae and Schizosaccharomyces pombe have already been added to BioGrid and curation efforts for other organisms are underway [Reguly et al., 2006, Breitkreutz et al., 2008].

The Kyoto Encyclopedia of Genes and Genomes (KEGG) system [Kanehisa et al., 2004] con-
CHAPTER 2. BIOLOGICAL DATA

consists of three main components: the genomic space (KEGG GENES), the chemical space (KEGG LIGAND), and network space (KEGG PATHWAY). The KEGG GENES is a collection of gene catalogues for completely or partially sequenced genomes, compiled by automatically extracting information from databases such as NCBI GenBank, and RefSeq (Wheeler et al., 2007). The KEGG LIGAND is further divided into several components including COMPOUND, GLYCAN, REACTION, ENZYME, and so on (Goto et al., 2002). The COMPOUND database contains manually entered and computationally verified chemical structures of known metabolic compounds, and some pharmaceutical and environmental compounds. The GLYCAN database consists of carbohydrate structures, a few hundreds of which were manually entered and the rest derived from CarbBank project (Doubet et al., 1989). The REACTION database contains reaction formulæ for enzymatic reactions, the reactants of which are represented in COMPOUND or GLYCAN databases. The ENZYME database contains enzyme nomenclature. Each enzyme is identified by an Enzyme Commission (EC) number (Webb, 1992; Tipton and Boyce, 2000), which can be linked to other public databases such as UniProt (Bairoch et al., 2005). The KEGG PATHWAY database is a collection of manually drawn diagrams, called KEGG reference pathway diagrams (maps), each of which corresponds to a known network of functional significance. Moreover, PATHWAY database also contains organism-specific pathways, which are automatically generated by superimposing genes in given organisms. The KEGG pathways are provided in an XML based markup language called KGML. Each metabolic reaction in a KEGG metabolic pathway is linked with one entry in the REACTION database, and the enzymes in the enzymatic reactions can be linked to the databases in the genome space as well as to other public databases via EC numbers.

iND750 is a manually reconstructed genome-scale metabolic model describing *Saccharomyces cerevisiae* metabolism (Duarte et al., 2004) with 750 genes, their transcripts, proteins and reactions. Manual reconstruction process involves curating reaction lists based on information from genome annotations, biochemical pathway databases, biochemistry textbooks, and publications (Fürster et al., 2003). All reactions in iND750 model are elementally and charge balanced, and compartmentalised to eight cellular locations: extracellular space, cytosol, mitochondrion, peroxisome, nucleus, endoplasmic reticulum, Golgi apparatus, and vacuole. Similar semi-automated manual curation has been employed to construct the first consensus metabolic network for yeast (Herrgard et al., 2008), global human metabolic network (Duarte et al., 2007), and so on.

Gene Expression Omnibus (GEO) (Edgar et al., 2002; Barrett et al., 2005) stores a variety of high-throughput molecular abundance data of which microarray gene expression data is a major data type. The data in GEO is organized into GEO Platforms (GPL), GEO Samples (GSM), GEO Series (GSE) and GEO Data sets (GDS). A Platform describes the set of elements that can be detected and quantified in the experiment. A Sample describes a single hybridization or experimental condition. A Series is a group of related Samples that make up one single study. A Data set is an assembly of biologically meaningful Samples that are statistically comparable. Of these, GPL, GSM and GSE are direct submissions of contributors, while GDS is a curated collection.

Gene Ontology (GO) (Ashburner et al., 2000; Berardini et al., 2010) consists of three independent ontologies: Biological process, Molecular function, and Cellular component to describe the roles of genes and proteins in eucaryotes. GO is built on the premise that a large fraction of the genes specifying core biological functions are shared by all eucaryotes. It is aimed to be a dynamic controlled vocabulary applicable to all eucaryotes even as our knowledge of gene and protein roles in cells continuously evolves.

Although the high-throughput experimental techniques of modern molecular biology empower us to measure multiple components of a biological system simultaneously, they often produce data that is inferior in quality to low-throughput techniques. On the one hand, high-throughput techniques may produce large number of false positives, meaning that not all findings are necessarily correct. On
the other hand, they may also exhibit large number of false negatives or detection biases, meaning that they may miss some true phenomena, leading to the lack of coverage or comprehensiveness in the findings. von Mering et al. (2002) performed a detailed comparison of multiple high-throughput techniques as well as a computational approach commonly employed for the study of protein-protein interactions. They estimated that more than half of all high-throughput protein-protein interaction data are false positives. While the estimated lower-bound to the number of protein-protein interactions is 30,000, the number of interactions supported by more than one method is only approximately 2,400, which demonstrates the sparing coverage of the methods employed (von Mering et al., 2002). For these reasons, manual curation of published literature has, despite the practical difficulties, received significant interest. In order to compile high-quality data, manual literature curation is expected to consider high-confidence data coming from low-throughput techniques, and perhaps also additionally supported by multiple independent studies. However, a recent study that has systematically compared the quality of a literature curated data has speculated that the quality of the literature curated yeast protein interaction data from BioGrid (Reguly et al., 2006) is at the most as good as or even inferior to high-throughput yeast two-hybrid (Y2H) data (Cusick et al., 2009). On the other hand, high-throughput techniques such as Y2H method are also improving, and it is increasingly believed that the little overlap among different high-throughput protein interaction studies is due not to the false positives, but to the false negatives (Lemmens et al., 2010). However, still the coverage of such data is quite little. For instance, while reporting a newly produced high-quality comprehensive binary Y2H protein interaction map for *Saccharomyces cerevisiae*, Yu et al. (2008) indicated that three proteome-level Y2H studies (i.e., Uetz et al. (2000); Ito et al. (2001); Yu et al. (2008)) taken together, only account for approximately 20% of the empirically estimated protein binary interactions in *Saccharomyces cerevisiae*. Y2H interaction maps have also been generated for other model organisms and humans (see references cited by Yu et al. (2008)), and their quality and coverage are similar to those of *S. cerevisiae* interaction maps. Global metabolic network reconstruction approaches described earlier (Duarte et al., 2004; Herrgard et al., 2008; Duarte et al., 2007) presumably produce high-quality metabolic networks but they are not complete and continuously keep growing. Comprehensive system-level data of high-quality is a key ingredient of systems biology (Kitano, 2002). Similarly, efficient computational tools to effectively handle current and future high-throughput data and turn them into knowledge are equally important.

Most of the databases described in this chapter provide easy access to web interfaces and tools for mining the data. However, these tools as well as the underlying data formats are different from each other and thus very diverse. A platform for accessing all these databases in a unified fashion as well as for performing advanced data mining of the resulting data is presented in the next chapter.
Chapter 3

Methods

3.1 \textit{MegNet} platform for biological data integration

Although high-throughput experimental techniques of molecular biology and biochemistry produce increasing amounts of data such as gene, protein, and metabolite expression, as well as interactions between them, characterising a sample’s phenotype in the context of environment or experimental condition remains a challenge. The aim of data integration is to address this challenge by providing links between a variety of these data.

By constructing a system to represent experimental phenotypes and environmental context of samples in GEO \cite{Barrett2005} with annotations from Unified Medical Language System (UMLS) \cite{Bodenreider2004}, and mining the data across multiple datasets representing similar biological contexts, \cite{Butte2006} showed that a large set of phenome-genome and envirome-genome relations could be retrieved within a public repository of transcriptome measurements (GEO). \cite{Oresic2004} explored integrated analysis of gene, protein and metabolite expression profiles, attempted to interpret the results in the biological context using pathways, and emphasised the need for data traversals.

We have developed a bioinformatics platform, a consolidation of multiple heterogeneous molecular biology databases, and a visualisation software called \textit{MegNet} for automatic integrative mining of these data \cite{ArticleI, ArticleII}. This section explains the technical details of the platform and \textit{MegNet}. Henceforth, we use the words \textit{MegNet} and \textit{data integration platform} interchangeably to represent the database system and the visualisation software together.

Fundamentally \textit{MegNet} system has been developed to achieve integration of heterogeneous biological data by enabling traversals across different data sources. It enables traversals across protein-protein interactions, transcriptional regulation reactions, metabolic pathways, metabolic models, signal transduction pathways, biological ontologies, and molecular profile data such as gene expression measurements \cite{Figure3.1}. \textit{MegNet} represents the integrated data as networks. Furthermore, \textit{MegNet} enables context-based visualisation of the integrated networks by building a conceptual space representation \cite{Garfors2000} and making use of the dimensionality reduction techniques \cite{Carreira-Perpinan1997} to visualize the similarity structure in a low-dimensional space, typically a two-dimensional plot.
Figure 3.1: Multiple molecular biology databases provide descriptions of biological systems at different levels of abstraction. Some common biological information, along with names of primary databases providing information in that domain are provided. Four levels of biological information are indicated by boundaries: molecular profiles (black), general information about molecules (red), interactions (blue), and biological pathways and functions (green).

3.1.1 Databases

A database is merely a collection of information that exists over a long period of time. A Database Management System, DBMS, is a software system that supports storage of large databases, provides efficient access to the data through powerful query languages, supports atomic and independent execution of concurrent transactions, and supports durability—the ability to recover from failures or errors (Garcia-Molina et al., 2002).
CHAPTER 3. METHODS

Relational databases

Relational database systems are based on relational data model (Codd, 1970) which organizes data as relations or tables and provides a high level query language called Structured Query Language (SQL) for manipulating and querying the relational data (Garcia-Molina et al., 2002). Due to its rigorous mathematical base in Relational Algebra, relational database systems became de facto standard in DBMS technology. Oracle (http://www.oracle.com) is a famous vendor of a powerful relational database management system.

XML databases

eXtensible Markup Language (XML, http://www.w3.org/xml) is a World Wide Web Consortium standard for document markup. XML quickly gained popularity as data exchange format due to its semantic capabilities and its supporting standards and technologies such as DTD, XMLSchema, DOM, SAX, XPath, XQuery and so on (http://www.w3.org). Native XML as well as XML enabled database systems have responded to increasing XML data management needs (Chaudhri et al., 2003). Tamino XML Server (http://www.softwareag.com) is a powerful native XML data management system. XML is a key technology in biology data management (pp. 291–319 of Chaudhri et al., 2003), and commonly used for data exchange based on many XML data standards (Spellman et al., 2002, Hucka et al., 2003, Hermjakob et al., 2004) and for data integration (Achard et al., 2001, Philippi and Köhler, 2004).

3.1.2 Overview of MegNet

Our data integration and visualisation system is composed of three layers as depicted in Figure 3.2. Data curated from heterogeneous biological sources including the ontologies constitutes the back-end of the system. The logic and algorithms such as those for the database traversals, network...
construction, network projections, and integration of gene expression data constitute the middle-tier. The visualisation client communicates with the middle-tier, which in turn communicates with the back-end. Fundamentally the queries that visualisation software enables can be divided into two types: data integration queries and network projection queries. In response to a data integration query sent by the visualisation front-end, the middle-tier queries the relevant databases and performs the integration using database traversals and optionally using additional correlation calculations and sends the results in the form of an integrated network. A network projection request can be made when an integrated network is already constructed, and the system does not use the database back-end in processing these requests. The result of a network projection request is a low-dimensional representation of the network, typically a 2-d plot of network nodes, where the distances between points reflect the similarity of nodes in the network. The similarity measures will be described later in this chapter.

3.1.3 Infrastructure

A detailed description of the infrastructure of the MegNet system is included in section Architectural design of Article II. However, for completeness, it is depicted in Figure 3.2 and a brief summary is given below.

**Back-end:** The XML data management system Tamino XML server (Software AG) is used for storing XML data. Oracle 10g database server (Oracle Inc.) is used for storing molecular profile data such as gene expression data and metabolomics data. Tamino Java API and Java Database Connectivity (JDBC) enable the communication between the middle-tier and Tamino and Oracle respectively.

**Middle-tier:** JBoss application server (Redhat Inc.) hosts the middle-tier, which is a set of Java Beans invokable by web services, and sends responses as web services. The communication between middle-tier and the visualisation front-end is handled via SOAP messages.

**Front-end:** The front-end consists of the user interface. Initially it was implemented in Java, but more recently it is implemented in C#.NET, and the Java client has been deprecated. The client communicates with middle-tier via SOAP messages.

3.1.4 Databases and data curation

Data from various public data sources were collected into our local database systems (Table 3.1, Article I, Article II). The curation of a public database involves several steps (section 2.3 of Article I). Usually every database might need some specialized steps in the curation process (Table 3.1), but the general steps are as follows:

- creating logical schemas which represent the logical structure and physical properties such as indexing to enable efficient queries.
- development of parsers to convert the non-XML data into XML or relational formats or to convert from one XML to another XML format etc.
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Table 3.1: Databases integrated into our system, along with information on some high level steps employed for integration. Appropriate schemas (XML or relational) were defined for each database. In some cases the logical schemas were readily available e.g., as XML schema definition (XSD) files, in which case they were customised as needed. If the XSD files were not available, they were developed manually. Tamino or Oracle specific physical schema structures including indices for efficient data retrievals were defined based on an analysis of common queries. Tamino mass data loader (or similarly Oracle SQL loader) was used for loading data.

<table>
<thead>
<tr>
<th>Database</th>
<th>Curation tasks</th>
</tr>
</thead>
<tbody>
<tr>
<td>UniProt (Bairoch et al., 2005)</td>
<td>Data for UniProt-SwissProt and UniProt-TrEMBL were available in XML format. Perl parser was written to extract some subset of XML nodes and to format the data for Tamino data loader.</td>
</tr>
<tr>
<td>BIND (Bader et al., 2003)</td>
<td>Data was available in XML format. A Java parser was written to format the data for Tamino data loader. In order to link the protein interactors to UniProt entries, the GenInfo identifiers were converted to UniProt accession numbers using SeqHound API (Michalickova et al., 2002).</td>
</tr>
<tr>
<td>DIP (Xenarios et al., 2002)</td>
<td>Data was available in XML format. A Perl parser was written to format the data for Tamino data loader.</td>
</tr>
<tr>
<td>MINT (Zanzoni et al., 2002)</td>
<td>Data was available in PSI-MI XML format (Hermjakob et al., 2004). A Perl parser was written to format the data for Tamino data loader.</td>
</tr>
<tr>
<td>BioGrid (Stark et al., 2006)</td>
<td>Data was available in PSI-MI XML format. A Perl parser was written to format the data for Tamino data loader.</td>
</tr>
<tr>
<td>KEGG (Kanehisa et al., 2004)</td>
<td>Pathway data was available in KGML, an XML format (Kanehisa et al., 2004). A Perl parser was written for further Tamino specific formatting. Data from KEGG LIGAND (Goto et al., 2002) database were available as formatted text files. For these data, XML schemas were developed and Perl parsers were written for constructing valid XML documents.</td>
</tr>
<tr>
<td>iND750 (Duarte et al., 2004)</td>
<td>Data were available as flat files. Parsers based on libSBML were developed to construct SBML representation (Hucka et al., 2003), and the data pertaining to compound identities were manually annotated with PubChem database identifiers (Wheeler et al., 2007).</td>
</tr>
<tr>
<td>TRANSFAC (Matys et al., 2006)</td>
<td>Data were available as formatted text files. XML schemas were developed and Perl parsers were written for constructing valid XML documents.</td>
</tr>
<tr>
<td>TRANSPATH (Krull et al., 2006)</td>
<td>Data were available in XML format. Perl parsers were written to extract some subset of XML nodes and for Tamino specific formatting.</td>
</tr>
</tbody>
</table>
CHAPTER 3. METHODS

<table>
<thead>
<tr>
<th>Database</th>
<th>Reference</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>GO (Ashburner et al., 2000)</td>
<td>Data were available in an RDF-based format, and correspondly XML schemas were readily available. A Perl script was written for Tamino specific formatting, and for inserting an additional &quot;Ontology&quot; element which describes whether a term is a &quot;Biological Process&quot;, &quot;Cellular component&quot; or a &quot;Molecular function&quot;. Due to the cyclic dependency of the root elements &quot;go:go&quot; and &quot;rdf:RDF&quot;, an empty schema was first defined in Tamino for &quot;rdf:RDF&quot; element. Then the schema for &quot;go:go&quot; was defined, and finally, the full structure of &quot;rdf:RDF&quot; element was redefined.</td>
<td></td>
</tr>
<tr>
<td>GEO (Barrett et al., 2005)</td>
<td>Data was available in simple line oriented text file format called SOFT format. Perl parsers were developed to extract information from the SOFT files. UMLS annotations for meta-data (Butte and Kohane, 2006; Bodenreider, 2004) were incorporated into maps database.</td>
<td></td>
</tr>
<tr>
<td>TIDBase (Hulbert et al., 2007)</td>
<td>The data were available as MYSQL sqldumps. A MYSQL database was created from the sqldumps, and then the MYSQL data has been exported to ORACLE using Oracle Database Migration Workbench.</td>
<td></td>
</tr>
</tbody>
</table>

Database traversals using schema maps

Biological system description involves various levels of abstraction (Figure 3.1) which include biological molecules (i.e., DNA, RNA, Proteins, Metabolites, and so on), biomolecular interactions (i.e., Protein-protein, Protein-DNA interactions, Transcriptional regulation steps such as a transcription factor binding to the binding site of a gene in order to control its expression, Metabolic reactions), pathways (i.e., cascades or sets of interactions working in concordance in order to perform biological functions), and biological processes (i.e., biological mechanisms involving, for instance, cell-cell, cell-tissue, organ level interactions, and so on). Some of such information can be accessed from the biological databases. Therefore, the integration of such databases allows us to automatically mine that information. In order to build a platform, which allows mining of a variety of such biological interactions, one should fundamentally achieve traversals across the databases providing biological information (Orešič et al., 2004).

Resolving even simple biological relationships that contain a few biological components often requires traversing across multiple databases. The traversals can be achieved by identifying names or identifiers in different databases, which represent the same biological entity. In order to enable traversals in our data integration system, we developed a database called "maps" database, which maps names used for the same entities across multiple databases. This database is populated by parsing information from several databases (Fig. 3B of Article I).

Currently this database contains mapping of proteins and gene expression experiments. The protein maps contain information of proteins indexed by Uniprot Swiss-Prot and TrEMBL identifiers (Bairoch et al., 2005). The information was primarily parsed from Uniprot Swiss-Prot and TrEMBL identifiers.
and further populated by supplementary information parsed from BIND database (Bader et al., 2003), KEGG Pathways (Kanehisa et al., 2006), and yeast metabolism models iND750 (Duarte et al., 2004) and YMN1 (Herrgård et al., 2008). Maps for gene expression experiments contain experimental meta descriptions and annotations from Unified Medical Language System (UMLS) where available (Butte and Kohane, 2006; Bodenreider, 2004).

As an example to illustrate how the data in maps database is compiled, we explain how protein entries in this database are linked to GenInfo identifiers provided in BIND molecular interaction database (Bader et al., 2003). In protein-protein interaction databases such as MINT and DIP, protein nodes are indexed by Uniprot identifiers (Zanzoni et al., 2002; Chatr-aryamontri et al., 2007; Xenarios et al., 2000). But in BIND, interactions are indexed by GenInfo sequence identifiers (GI’s) (Bader et al., 2003). We obtained mapping between the GI’s and Uniprot identifiers in each BIND interaction using DBXREF table that provides all external references made in BIND and SeqHound Perl API (Michalickova et al., 2002) as follows. We first extracted the list of unique UniProt accession numbers from the DBXREF table. Then, for each Uniprot accession number in this list, we did the following. We first found the corresponding primary GI using SHoundFindAcc function of SeqHound Perl API. Next, we retrieved the list of GI’s that have the exact sequence as that of the primary GI using SHoundRedundantGroup function of the API. We finally stored mapping between all the GI’s whose taxonomy id (as found by SHoundTaxIDFromGi function of the API) matched that of the primary GI. Finally, all GI’s associated with each UniProt accession were added as external database links for the corresponding protein entry, indexed by the UniProt accession number, in maps database.

The data traversals use protein nodes as central nodes for constructing integrated networks. More specifically, in order to construct an integrated network consisting of various types of edges in it, those edge types are joined based on common protein nodes they share. As an example, consider traversing from a metabolic reaction to a protein-protein interaction. Such traversal could bring information about the protein-protein interaction partners, when exist, of the enzyme which catalyzes the metabolic reaction. It can be typically achieved by translating the Enzyme Classification (EC) number of the enzyme to the corresponding protein identifier (Uniprot Id), and then searching protein-protein interaction databases for any possible interactions partners (Fig 3 of Article I).

Similarly, for linking correlation networks based on gene expression measurements with interaction networks we find out identifiers of the proteins which are expressed by the genes (indexed by EMBL sequence identifiers) under consideration.

In order to achieve data traversals, the maps database is designed to contain identifiers and names of protein entities from multiple databases. Conversions from other types of identifiers, e.g., EC enzyme identifiers to Uniprot protein identifiers, and linking of e.g., EMBL gene identifiers to Uniprot protein identifiers are obtained from maps database.

3.1.5 Integration of Gene expression data

We achieved integration of gene expression data with interaction networks in other ways (sections 3.2, 4.2). At this point, however, we limit the discussion to network visualisation based integration approach. The visualisation approach is based on correlation networks. For selected biological samples, possibly from one or more gene expression studies, the integration approach will be explained below.

Gene co-expression across multiple experimental conditions and multiple organisms indicates strong functional relationship, and hence is a powerful tool for elucidation of gene function (Stuart et al., 2003). Gene Expression Omnibus (GEO) (Edgar et al., 2002; Barrett et al., 2005) stores high-throughput gene expression data from many organisms and a huge variety of biological conditions.
In *MegNet* we allow the users to assess co-expression of genes from any choice of combinations of data sets and integrate the information with pathway and interaction data.

**Gene expression data preprocessing**

Many variations introduced at different stages of microarray experiments blur the real biological variation and microarray data preprocessing tries to remove the non-biological variations from the data (Speed, 2003; Quackenbush, 2002; Bolstad et al., 2004). Within each GEO data set, the statistical preprocessing of the data such as background correction and normalisation are performed in a consistent fashion (Barrett et al., 2005), thereby removing the experiment-specific non-biological variations. But in order to enable simultaneous mining of gene expression data from multiple studies together, we need further normalisation so that different data sets can be combined. For example, in building ONCOMINE system to mine a large number of diverse cancer microarray data sets, Rhodes et al. (2004) applied median centering and standard deviation scaling per microarray. In an across-laboratory reproducibility study of microarrays, Irizarry et al. (2005) indicated that studying relative expression values instead of absolute expression values is a simple solution to remove the probe-specific effects in particular experiments.

In the data collected from GEO, there are two major types of gene expression microarray datasets: single channel (i.e., intensity based) microarrays (Lockhart et al., 1996) such as Affymetrix oligonucleotide microarray data, and dual channel (i.e., two colour) microarrays (Schena et al., 1995) such as cDNA microarray data. Therefore, the cross-platform comparability problem reduces to applying normalisation so that both these dataset types can be analysed in an identical fashion. Since GEO data provides log2 ratio between individual channel intensities (i.e., between case and control) for the dual channel arrays, we similarly normalise the single channel data of case samples with control samples from the same study, where the selection of the case and control samples is upto the user.

To be more precise, for each case sample, we compute the log2 ratio of the gene expression intensity measurement versus the average intensity of control samples from the same GDS dataset. After this transformation, single channel data and dual channel data can be analyzed in identical fashion as well as simultaneously. In order to allow data sets coming from different studies to be combined, we scale each microarray to unit standard deviation. Finally, when multiple data sets are queried, only the genes common to all microarray platforms are used for the analysis.

**Statistical hypothesis testing**

Statistical hypothesis testing (Box et al., 1969; Montgomery, 1983) is a framework that allows us to answer particular questions related to one or more populations on the basis of samples randomly drawn from those populations. A statistical hypothesis is an assumption about the probability distribution of a population. Hypothesis testing generally involves the following steps

**Formulation of null hypothesis ($H_0$):** The null hypothesis is the hypothesis that the results observed in a study (e.g., difference between treatment group and control group) are purely by chance. The null hypothesis would be rejected if data does not provide enough evidence to its truth; otherwise, we fail to reject it. An alternative hypothesis ($H_a$) is complementary to the null hypothesis, and is effectively favoured when the null hypothesis is rejected.

**Calculating a test statistic:** The test statistic is a measure of the size of the “effect” relevant to our test.

**Calculating the P-value:** The significance probability or $P$-value is the probability of getting the data at least as extreme as observed if the null hypothesis were true.
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Rejecting or failing to reject the null hypothesis: The level of significance ($\alpha$) is the maximum probability with which we are willing to falsely reject the null hypothesis. The null hypothesis would be rejected if the significance probability ($p$-value) is less than the level of significance ($\alpha$); the evidence to reject the null hypothesis is considered inadequate otherwise.

Parametric tests make particular assumptions on the properties of the populations. When the parametric assumptions are not satisfied, non-parametric tests may be used instead. The advent of powerful computers allows a new approach called a permutation test (Moore and McCabe 2005) for performing a non-parametric test. A permutation test employs resampling to estimate the significance of the test statistic. Permutation tests are often more robust than formula based non-parametric or parametric tests. They are applicable even though the parametric assumptions are not satisfied, as long as the resampling is done in a way that is consistent with the null hypothesis (Moore and McCabe 2005).

Multiple hypothesis testing

In the context of hypothesis testing, two types of errors are possible: Type I error or Type II error. Rejecting a true null hypothesis is called the Type I error. Failing to reject a false null hypothesis is called the Type II error. When multiple hypotheses are tested at a specified Type I error probability ($\alpha$) for each test, the chance of committing at least one Type I error increases sharply with the number of hypotheses, and such phenomenon is referred to as multiple comparison (or testing) problem. In order to address the multiple comparison problem, one needs to define an appropriate Type I error rate and devise powerful multiple testing procedures that control this error rate (Shaffer, 1995; Dudoit et al., 2003). Controlling the false discovery rate (FDR) is one such approach, in which, the FDR—the expected proportion of falsely rejected null hypotheses (Benjamini and Hochberg, 1995)—is controlled. Procedure for controlling FDR proposed by Benjamini and Hochberg (1995) is as follows. Sort the $p$-values of the test in ascending order, and denote the sorted order as $p_1 \leq p_2 \leq p_3 \leq \cdots p_m$.

Represent the corresponding $m$ hypothesis with the same indices $H_1, H_2, H_3, \ldots, H_m$.

For controlling the FDR at level $\alpha$, define

\[ j_0 = \max \{ j : p_j \leq \frac{j}{m} \cdot \alpha \} \]

and reject hypothesis

\[ H_i, \forall i \in 1, 2, 3, \ldots, j_0. \]

Other classical Type I error rates controlled in multiple comparison approaches include Per Family Error Rate (PFER)—expected number of type I errors, and Family Wise Error Rate (FWER)—the probability of at least one type I error (Shaffer, 1995), both of which tend to impose more strict control over the Type I errors, but at the expense of larger number of Type II errors. Therefore, controlling the FDR is common choice in exploratory analysis (Dudoit et al., 2003).

Correlation

Correlation between two variables $X$ and $Y$ is the extent to which their values vary together systematically (Box et al., 1969). A correlation coefficient is a measure to the extent of correlation between
two variables. Product-moment correlation coefficient or pearson correlation coefficient, which is calculated from given samples of $X$ and $Y$ as the ratio between their covariance and the product of their standard deviations, indicates the extent to which relationship between the variables $X$ and $Y$ is linear. Pearson correlation coefficient obtained from samples ($r$) is an estimate of the true correlation ($\rho$) of the bivariate normal population ($X, Y$). The null hypothesis that $\rho = 0$ can be tested with the statistic

$$t = \frac{r \cdot \sqrt{n - 2}}{\sqrt{1 - r^2}}$$

which follows Student’s distribution with $(n - 2)$ degrees of freedom (Box et al., 1969). The $p$-value of this test is the probability that pearson correlation coefficient of the sample ($r$) would be at least as extreme as observed if the null hypothesis is true (i.e., if $\rho = 0$).

Co-expression network

MegNet allows studying co-expression of genes from multiple GEO data sets emanating from different labs, different studies, and possibly using diverse microarray platforms. Genes that are common to all microarray platforms in the query are identified and used for the combined analysis using correlation via Pearson correlation coefficient. For visualizing gene expression data, we compute correlation and $p$-value between every pair of genes in the selected datasets. We then apply FDR method of Benjamini and Hochberg (1995) to account for the multiple hypothesis testing. A correlation network is then defined as the collection of edges representing statistically significant correlation between the pairs of gene nodes they connect. Edges sharing common genes are joined at the common gene nodes.

Finally data traversals can be also combined with co-expression network construction in order to link the correlation network to molecular interaction networks, pathways and ontologies. For example, MegNet system may be used to look for interaction neighbourhood of entities on the correlation network. But more generally, it is possible to combine interaction network construction parameters (e.g., certain proteins, pathways or ontologies) and co-expression network construction parameters (e.g., some particular selection of gene expression data sets) in the same query, and then let the data traversals automatically discover the cross-talk between these different levels wherever possible (see e.g., section 4.1.4).

3.1.6 Context-sensitive data mining

Conceptual spaces

Gärdenfors (2000) advocates conceptual spaces as a representation paradigm to modelling human cognition and as complementary to the Symbolic and Associationist levels of representation used in cognitive science. He argues that these three levels are representations at different levels of resolution. In conceptual spaces, concepts are represented as high-dimensional geometric spaces over a variety of quality dimensions with geometric or topological structure for one or more domains. A domain can be modelled as a set of integral dimensions i.e., dimensions which need not necessarily be completely independent. For example, an apple might be represented as a multi-dimensional space where the dimensions could be “colour”, “taste”, “shape” and so on. Thus, while an “apple” is simply a token (or a text string) at the Symbolic level, it has a rich underlying geometric description at the conceptual level.

There are two approaches to explaining meanings: Realist and Cognitive semantics. According to Realist semantics, meanings of expressions are independent of the individuals involved in communication; meanings are in the world. According to Cognitive semantics, meanings are mental entities;
different people have different conceptualisations of the world. The realist semantics fail to deal with the fact that different people have different approaches to learn and that meanings of concepts often change over time and between contexts. Context plays an important role in human learning or understanding of new concepts. In terms of conceptual spaces, different conceptualisations correspond to considering different subsets or alternative choices of quality dimensions. The dynamic nature of the meaning, i.e., changes in meaning of concepts with respect to time and context, can be represented by giving different saliencies (i.e., weights) to dimensions and domains.

**Context-sensitivity in conceptual spaces**

Context-sensitivity can be achieved in conceptual spaces by specifying weights to the quality dimensions and recomputing the similarities (Gärdenfors, 2000). However, how to appropriately assign weights to the dimensions according to the context is not necessarily trivial.

In building an information inference application, Song and Bruza (2003) employed a heuristic based on query term frequency in text corpus to automatically find out the dominant concept in a concept combination. The concept combination heuristic was then applied to emphasise quality properties shared by the concepts in the concept combination and to increase the weights of properties of the dominant concept by rescaling (Song and Bruza, 2003). Raubal (2004) in their way-finding application used different sets of empirical weights to represent the users preferences on facades according the day and night.

However, the true power of conceptual spaces representation is in allowing each user to have one's own conceptualisation of the world. Therefore, in an interactive system, the user must be given full control of weight assignment. This is especially useful for facilitating exploratory data mining. Meanwhile, the system may offer some predefined or default contexts defined by some particular choices of weights in order to facilitate queries from users with little experience.

Using some specific experimental context of interest, if available, maybe a useful alternative. For example, a gene expression experiment data maybe used to assign weights based on some measure such as correlation coefficient (see e.g., section [1.5]).

**Conceptual representation of MegNet networks**

An integrated network constructed using database traversals as explained earlier corresponds to symbolic representation of biological relationships obtained using semantic web technologies. Therefore representing the information in the networks as a conceptual space makes it possible to mine the data in a context-sensitive manner, or in other words, to facilitate Cognitive semantics.

In this direction, we represent every MegNet integrated network as a high-dimensional space (Figure 3.3) whose dimension is defined by the number of nodes in the network (Article IV). Every node in the network is a vector in this space. A vector \( v \) representing a node \( p \) contains weighted shortest path distances from node \( p \) to all other nodes in the network. These weights are the subject of discussion in the next paragraph. For now, it is enough to note that, by default, the weights for all interactions are set to 1. Therefore, the default representation contains shortest path length for each coordinate.

As discussed earlier, context-sensitivity in a conceptual space can be achieved by specifying one’s own conceptualisation in terms of weights on quality dimensions. But translating one’s conceptualisation to weight assignment may not always be trivial. In fact, finding the best weights may need exploration. Therefore, to facilitate interactive exploration, as a pragmatic alternative, we give the full control of the weights to the user, and only define the default context (or system’s view of the world), in which we assign a unit weight (i.e., weight of 1) to each interaction type.
Figure 3.3: Conceptual space representation and context visualisation of MegNet networks. A context is defined by a specific choice of weight assignment to the edges. Every network node is a vector in a high-dimensional space where each coordinate represents the weighted shortest distance of this node from a particular node in the network. Dimensionality reduction methods enable us to visualise this space in a low-dimensional (typically 2-d) plot.
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As the integrated network consists of heterogeneous edge types, translating the user’s context specification (i.e., preferences on weighting the quality dimensions) to the assignment of weights to individual edges (or weight normalisation) is not obvious, because the underlying biological meaning of the edges has to be preserved. For example, a metabolic reaction consists of two edge units: the first edge connecting substrate to enzyme, and the second connecting enzyme to the product. Therefore, assigning a weight of 1 unit to a metabolic reaction is, for instance, achieved by assigning a weight of 0.5 to each of these two edges. For each datatype incorporated in our database system, we define the standardisation for the distances in MegNet, so that the user is not required to know the underlying representation of the data while specifying weights to interactions.

Visualisation of a context

Typically, MegNet integrated networks are very complex in the sense that they contain thousands of nodes and edges. Hence, understanding the conceptual space encoded in such a network under a particular context is a formidable task. Visualising the semantic similarities encoded into the conceptual representation allows us to visualise a particular context specified by the user. Therefore, as a solution to the visualisation problem, we apply dimensionality reduction techniques (Carreira-Perpiñan, 1997) and visualise the distances in a lower-dimensional space (typically 2-d) (Figure 3.3). Dimensionality reduction is most often possible, because the original high-dimensional representation of the data often contains redundancies and dependencies between the variables. Under the assumption that the data lie on a manifold whose dimension is smaller than the dimension of the embedding space, the dimension reduction can be achieved by the construction of a continuous mapping between the embedding space and the unknown manifold space. This mapping has to be reversible in order to project and reconstruct the original data with minimal error (Lee et al., 2004). The true dimension of the unknown manifold is known as the intrinsic dimension of the data.

The intrinsic dimension needs to be provided as an input to most dimension reduction techniques to build a correct mapping i.e., to avoid over or under fitting. Thus, finding the correct value for the intrinsic dimension might be an exploratory undertaking. However, 2-dimensional plots are the easiest and the most common choice for visualisation. A variety of dimension reduction techniques exist, and are usually utilised for a variety of purposes such as visualisation, data compression, and variable selection (Carreira-Perpiñan, 1997). As we are particularly interested in the visualisation of the semantic distances, our focus is on dimension reduction techniques that preserve the pairwise distances of the data space in the projection space. Therefore, we implemented three non-linear dimensionality reduction methods in MegNet, namely, Sammon mapping (Sammon, 1969), Curvilinear component analysis (CCA) (Demartines and Hérault, 1997), and Curvilinear distance analysis (CDA) (Lee et al., 2000, 2002, 2004). The aim of these projection methods is to represent the high-dimensional vectors in a low-dimensional space in such a way that the distances are preserved by the mapping. Each method achieves this objective by minimising a cost function. Technical details such as the analytical formulae of the cost functions, the computational complexities of these algorithms, and our implementation of these ideas are discussed in detail in Article II and Article IV. Some important properties of these methods are given here for completeness.

Sammon’s mapping (Sammon, 1969) estimates the configuration of entities in the projected space using steepest gradient descent algorithm on a cost function that is based on the interpoint distances between the entities in the original space and the discrepancies introduced by the dimensionality reduction mapping (see section 3.3.1 of Article II). In this way, the visual configuration approximates the original relationships in the complex networks. Sammon’s mapping puts higher emphasis on preserving smaller distances. Curvilinear component analysis (CCA) (Demartines and Hérault, 1997) attempts to preserve the topology by first favouring shorter distances, and then the longer
distances. CCA uses stochastic gradient descent. Curvilinear Distance analysis (CDA) (Lee et al., 2000, 2002, 2004) maps entities in higher-dimensional space into lower-dimensional space in such a way that the curvilinear distances, as opposed to Euclidean distances, between points in high-dimensional space are preserved. Curvilinear distance is the shortest path distance in the centroid graph in the high-dimensional space. CDA uses stochastic gradient descent algorithm to minimise the projection error.

3.2 Dynamic topology of integrated networks

High-throughput functional genomics techniques such as gene expression microarrays (Schena et al., 1998; Lockhart and Winzeler, 2000), mass spectrometry based proteomics (Aebersold and Mann, 2003) and metabolomics (Goodacre et al., 2004; Orešić et al., 2006) allow us to measure the state of a biological system in terms of molecular abundances or concentrations. Databases describing signalling (Krull et al., 2006), metabolic pathways (Kanehisa et al., 2004; Goto et al., 2002) and biological functions (Ashburner et al., 2000) provide reference information on cellular processes. One important goal of developing our data integration platform is to enable interpretation of new experimental data in the context of the established pathway knowledge.

3.2.1 Transcriptomics

Transcriptome is the collection of messenger RNA (mRNA) molecules or “transcripts” in a cell or a tissue. Transcriptomics is the discipline which deals with the large scale study of the transcriptome under selected conditions. DNA microarray technology (Schena et al., 1995; Lockhart et al., 1996), capable of measuring activity of thousands of transcripts simultaneously, has served as the platform of choice for transcriptomics.

3.2.2 Metabolomics

The word metabolome represents the whole collection of metabolites in an organism. Metabolomics is defined as the comprehensive study of the whole metabolome under particular conditions (Fiehn, 2001; Goodacre et al., 2004). It embodies global study of all metabolites, their dynamics, composition, interactions, responses to interventions or environmental changes (Orešić et al., 2006). Owing to the fact that small changes in the activities of individual enzymes can lead to large changes in metabolite concentrations, metabolomics is a more sensitive tool than transcriptomics and proteomics for studying complex diseases (Orešić et al., 2006) and the regulatory roles of nutrition in human health (Gibney et al., 2005). But due to the huge diversity among metabolites, different technologies are required for studying different classes of metabolites, yet it is impossible to study the whole metabolome (Orešić et al., 2006). An interesting subfield of metabolomics is lipidomics, a discipline which deals with global study of lipids. Lipids are highly diverse molecules which play crucial roles in cellular energy storage, structure and signalling (Orešić et al., 2008; Seppänen-Laakso and Orešić, 2009).

3.2.3 Pathway analysis

A challenge in the analysis of data arising from functional genomics experiments is to understand the results in the context of established information such as biological processes, metabolic pathways and so on. Understanding what biological processes are significantly modulated under a particular
intervention is often more informative than finding a set of individual molecules that are differentially regulated (Curtis et al., 2005).

Recently a set of techniques broadly known as Pathway analysis methods have been developed to achieve this goal (Curtis et al., 2005). A most prominent technique that enables pathway analysis of genome-wide gene expression data is Gene Set Enrichment Analysis (GSEA) (Mootha et al., 2003; Subramanian et al., 2005). By determining which gene sets (i.e., groups of genes that share common biological function, chromosomal location or regulation) are significantly enriched among the significantly modulated genes in a gene expression study, GSEA not only offers an easy interpretation of the data, but also accounts for the fact that single-gene analysis (i.e., analysis of differential expression using a statistical test) may miss important “group or pathway effects”. By definition, a gene set represents the set of genes that act in concert to “perform” a biological function. A moderate but concordant change in a gene set therefore may be more important than a very high change in a single gene in the set. This is called the “group effect or pathway effect”. Moreover, Subramanian and colleagues (Subramanian et al., 2005) also showed, by applying GSEA on three different lung cancer data sets, that the pathway level changes are more reproducible than the individual gene markers. Furthermore, GSEA is not only limited to transcriptomics data analysis. Subramanian et al. (2005) said, “GSEA can clearly be applied to other data sets such as serum proteomics data, genotyping information, or metabolite profiles”. Recently GSEA algorithm has also been adapted for the data analysis in genome-wide association studies (Wang et al., 2007; Holden et al., 2008).

Numerous conceptually similar but competing approaches to GSEA exist, which employ alternative enrichment statistic and permutation testing scheme, seeking to improve the statistical power of the pathway analysis (Tian et al., 2005; Efron and Tibshirani, 2006). Moreover, high quality pathway databases would enhance the usefulness of pathway analysis. Multiple proprietary pathway databases and pathway analyses software also exist (http://www.ingenuity.com, http://www.ariadnegenomics.com, http://www.genego.com).

Finally, several studies reported that gene level biomarkers are not reproducible (Tan et al., 2003; Michiels et al., 2005; Ein-Dor et al., 2005). Despite this poor reproducibility, however, pathway level changes are more consistent (Subramanian et al., 2005; Zhang et al., 2008). Thus, pathway level analysis is a more promising tool for identifying the disease mechanisms, and adaptive physiological compensatory responses (Curtis et al., 2005).

### 3.2.4 Topology of biological networks

Topological properties of a variety of biological networks such as protein-protein interaction and metabolic networks have been extensively studied by many researchers (Barabási and Oltvai, 2004). Due to intriguing similarities such as scale-free topology between the topological properties of biological networks and other networks such as social networks, scientific collaboration networks and so on, Network biology has been an active area of research. Besides analysing the structures of static networks, studying the network dynamics is of interest as it may enable elucidation of dynamic design principles. Kharchenko et al. (2005) studied the expression dynamics of a metabolic network and discovered relationships between pairwise distances on the metabolic network and the co-expression of genes. Lascombe et al. (2004) studied regulatory network dynamics by integrating transcriptional regulatory information and gene-expression data, and showed that, in response to diverse stimuli, transcription factors alter their interactions to varying degrees causing large topological changes in the regulatory network.

In order to study the topological dynamics of integrated networks at the level of functional modules and to facilitate analysis and interpretation of molecular profile data such as gene expression data via integrated network connectivity, we have established a method called Topological Enrichment
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Analysis of Functional Subnetworks (TEAFS). This method benefits from genome wide integrated networks constructed using *MegNet* and genome wide gene expression profiles. Article V reports an application of this method for studying dynamic response of oxidative stress, the results of which are presented in section 4.2.

3.2.5 Integration of transcriptomics and interactomics data

In a recent study, subnetwork markers identified by integration of protein-protein interaction networks and gene expression data (Chuang et al., 2007) were found to be more reproducible than individual gene markers found by differential gene expression analysis and they achieved higher classification accuracy in discriminating metastatic versus non-metastatic tumours. Genes with known breast cancer mutations are typically not significantly differentially expressed, but were found to play central roles in protein networks by interconnecting many differentially expressed genes. This demonstrates the value of studying the gene regulation at the level of interaction network connectivity.

Although gene set enrichment analysis (GSEA) (Subramanian et al., 2005) and the related methods account for the subtle but coordinated patterns of gene expression changes, they do not take the connectivity of the system into account. For example, loss of a central node from the interaction network representing the biological process may lead to compensatory rise in the expression of multiple genes involved in the biological process simultaneously, but the loss of connectivity would render the network dysfunctional. However, due to the simultaneous rise in the expression of multiple genes, GSEA would report this dysfunctional pathway to be significantly up-regulated.

Recently a method called Gene Network Enrichment Analysis (GNEA) has been developed (Liu et al., 2007). This method takes the connectivity of proteins in protein-protein interaction networks into account in order to find pathways that are consistently affected across multiple interventions or models related, for instance, to a particular human disease, using gene expression data. Thus, the philosophy of this method is not to identify biological processes perturbed under a particular intervention but rather to discover which biological processes are transcriptionally altered across diverse tissue types in the context of a disease.

Outline of TEAFS

Topological Enrichment Analysis of Functional Subnetworks (TEAFS) facilitates understanding of how a specific biological intervention modulates biological functions. Below we provide a general outline of the TEAFS method. More specific details of TEAFS for studying dynamically most changing modules can be found in Article V.

Construction of an integrated network: TEAFS uses connectivity of protein nodes in an interaction network in terms of various topological measures (Albert and Barabási, 2002) to compute the topological changes of subnetworks representing functional modules. The changes in the topology of the subnetworks indicate the changes in activity of functional modules. Thus an interaction network is an input to this method and forms the starting point for this analysis. Any genome wide interaction network of the organism of interest, such as a protein-protein interaction network can be used for this purpose, but we take the unique advantage of the *MegNet* integrated networks.

Gene detection: The second input to TEAFS is the gene expression data that is intended to be analysed. This data is used to compute whether a protein is “present” (P) or “absent” (A) based on whether or not the transcript encoding the protein shows sufficiently high expression level. For
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probeset detection in Affymetrix Oligonucleotide arrays, MAS5 software (http://www.affymetrix.com) uses Wilcoxon signed rank test for perfect match (PM) and mismatch (MM) probes, and “calls” the probeset “present”, “absent” or “marginal” (M) based on the p-value (m Liu et al., 2002). More recently, an alternative gene detection method called PANP was introduced for computing P/A calls for oligonucleotide arrays (Warren et al., 2007), which uses only the PM probes for Affymetrix chips. For spotted DNA arrays, if the hybridization signal intensity of a spot is above the background intensity distribution, the corresponding gene may be considered present. Luscombe et al. (2004) applied one such criterion for computing the presence or absence of transcription factors in order to reconstruct transcriptional regulatory networks. For each sample in the selected gene expression study, we compute the P/A corresponding to all transcripts analysed on the chip. For studying the effects of yeast oxidative stress, we employed the P/A criteria similar to Luscombe et al. (2004) (Article V).

Reconstruction of networks: We identify the proteins encoded by the transcripts, for instance by translating the accession numbers, and thus infer which proteins in the network are present and absent in each condition. For each sample in the gene expression study, we construct the condition specific network by removing the absent protein nodes and the edges incident on them from the original network.

Identification of Functional Subnetworks: Functional subnetworks in the integrated network can be identified by using the established knowledge of the functional association of proteins. For example, by identifying which proteins in the network are associated with a particular Gene Ontology term (Ashburner et al., 2000), we can identify the functional subnetwork consisting of all these proteins and their neighbouring interactions. Other functional categories such as metabolic pathways (Kanehisa et al., 2006) may also be used similarly.

Computation of Topological Measures: In a directed network, the number of edges coming into a particular node is called the “in degree” of the node, and the number of edges going out of the node its “out degree”. Two nodes connected to each other by an edge are called neighbours. Clustering coefficient of a node is the ratio between the number of edges between its neighbours and the maximum number of theoretically possible edges between those neighbours. A topological measure (i.e. in degree, out degree or clustering coefficient) of functional subnetwork in an integrated network is the average of the topological measure of all nodes that are members of the subnetwork (Albert and Barabási, 2002).

Enrichment score: Enrichment score or enrichment statistic assigned to a functional subnetwork is the quantity of interest that we compute based on what questions we want to answer. In order to find out the most changing subnetworks during a time course following a particular intervention, an appropriate enrichment score would represent the degree of topological change the subnetwork is subjected to over time by the underlying biological intervention. In order to find out the most changing functional subnetworks during the time course following oxidative stress in yeast we employed standard deviation of topological measure as the enrichment score (Article V).

Computation of Statistical Significance: To compute the statistical significance of the enrichment score, a null distribution for the enrichment score is calculated, based on the random P/A model of proteins in the yeast oxidative stress study (Article V). A permutation test (Moore and McCabe, 2005) was devised for this.
Chapter 4

Results

4.1 MegNet based data mining

According to Hand et al. (2001), “Data mining is the analysis of (often large) observational data sets to find unsuspected relationships and to summarise the data in novel ways that are both understandable and useful to the data owner”, and “the relationships and summaries derived through a data mining exercise are referred to as models or patterns”. MegNet data integration system facilitates mining of biological data and hence exploration of some useful patterns, novel relationships between different biological entities from the data, and may provide novel insights into protein functions and context-specific biological functions.

Here, we summarise some MegNet data mining examples that were published in Article I, Article II, and Article III. The examples provided in sections 4.1.1 and 4.1.2 describe mining based solely on data traversals. Section 4.1.3 presents a case of a conceptual space representation with unit weights. Section 4.1.4 presents the case of integration between co-expression network with interactions and ontologies. Finally, section 4.1.5 presents context-sensitive mining of interaction and ontology information using the context of the gene expression experiment.

4.1.1 Integrated pathway retrieval

When attempting to model real biological phenomena one needs to understand the cross-talk across different levels of biological organisation, for instance, between metabolic pathways and cell signaling (Papin and Palsson, 2004). MegNet models the cross-talk across different levels through database traversals.

As an example, we queried the following Saccharomyces cerevisiae metabolic pathways: Glycolysis / Gluconeogenesis, Pentose phosphate pathway, and TCA cycle. More specifically, the query processing would include retrieval of primary components (i.e., enzymes and compounds) of the metabolic pathways from KEGG (Kanehisa and Goto, 2000), and then database traversals to search protein-protein interaction databases BIND (Bader et al., 2003) and MINT (Zanzoni et al., 2002) for interactions of the enzymes with the nearest neighbor proteins.

The resulting networks showed surprisingly high level of connectivity across different stages of linear metabolic pathways via protein-protein interactions (Fig. 5 of Article I). Specifically, two enzymes from the glycolysis pathway: phosphoglycerate kinase (PGK; EC: 2.7.2.3) and acetate-CoA ligase (ACS; EC: 6.2.1.1) appeared to aggregate with SRB2 (Uniprot accession: P34162) via protein-protein interactions.
ACS catalyzes formation of acetyl-CoA from acetate, which is a starting point in the TCA cycle, while PGK catalyzes acetylation of 3-phospho-d-glycerate, which is a part of the second phase of glycolysis. SRB2 is involved in transcriptional initiation. This could mean that PGK and ACS, enzymes at two different stages of glycolysis, are co-regulated. Thus, our results point towards a testable hypothesis (Article I).

4.1.2 Protein neighbourhood search

Assignment of protein function is a non-trivial task owing to the fact that the same protein may be involved in different biological processes, depending on the state of the biological system and protein localisation (Camon et al., 2004). *MegNet* allows visualisation of interaction neighbourhood of a protein, i.e., the entities in the network close to the protein, which may potentially provide insights into the function and mode of action of a protein. These entities include molecules, genes or more complex concepts.

As an example, we searched the neighbourhood of mannose-6-phosphate isomerase for *Saccharomyces cerevisiae* (PMI40; UniProt Id: P29952), which catalyzes the conversion between fructose-6-phosphate and mannose-6-phosphate and thus connects glycolysis with the cell wall synthesis in *S. cerevisiae*. The search involved concurrent retrieval of relationships for the following databases: UniProt (Bairoch et al., 2005), KEGG (Kanehisa and Goto, 2000), BIND (Bader et al., 2003), MINT (Zanzoni et al., 2002) and GO (Ashburner et al., 2000). Figure 6 of Article I shows the neighbourhood of PMI40. The zoomed-in window shows one region of potential interest, which includes protein-protein interactions between the PMI40 and NUP100 (UniProt Id: Q02629), a subunit of the nuclear pore complex, as well as between NUP100 and alpha-1,6-mannosyltransferase (MNN10; UniProt Id: P50108). Also both PMI40 and MNN10 are involved in cell wall mannoprotein synthesis (GO:0000032).

PMI40 is a gate between cell wall synthesis and glycolysis, i.e., cell decision point between growth or energy production, and MNN10 is a part of the protein complex in mannoprotein synthesis toward the end of the cell wall biosynthesis pathways. Examination of interaction entries (BIND id’s: 137955 and 137823) suggests that NUP100 protein, which is a part of nuclear pore complex, binds to the PMI40 and MNN10 open reading frames (Casolari et al., 2004). This and other evidence by Casolari et al. (2004) provide support for the gene gating hypothesis, which suggests that the interaction of the nuclear pore complex with different genes might serve as a level of gene regulation (Blobel, 1985). Thus, again, *MegNet* based mining leads to a testable hypothesis that PMI40 and MNN10 are co-regulated in relation to cell decision-making between energy production versus growth.

4.1.3 Context-dependent protein function visualization

In order to suggest protein function annotations to a set of human proteins related to maintenance of energy homeostasis and specific G-protein coupled receptors (GPCRs) that are not yet well characterized—PPAR-γ, PPAR-α, PGC1-α, SREBP2, GPR40, GPR41, and GPR43—we searched their neighbourhood using protein-protein interaction databases BIND (Bader et al., 2003), MINT (Zanzoni et al., 2002), DIP (Xenarios et al., 2000), metabolic pathway database KEGG (Kanehisa and Goto, 2000), transcriptional regulation database Transfac (Matys et al., 2006) and Gene Onology databases. The resulting network contained three isolated subnetworks, with all three GPCRs jointly in one subnetwork (Fig. 1 of Article III). While some of the well known relationships were revealed in the largest subnetwork, the results of the query have not facilitated characterization of poorly annotated proteins such as GPR40, GPR41, and GPR43.
We then formulated the conceptual space representation of the data, by assigning unit weights to all the edges in the network and visualised the resulting similarity structure using curvilinear distance analysis (CDA) (Lee et al., 2004) (Fig. 2 of Article III). According to this figure, in the underlying conceptual space, PPAR-γ and GPR41 are closely associated with response to nutrients (GO:0007584). PPAR-γ (UniProt id: P37231) is annotated in UniProt as “Receptor that binds peroxisome proliferators such as hypolipidemic drugs and fatty acids. Once activated by a ligand, the receptor binds to a promoter element in the gene for acyl-CoA oxidase and activates its transcription. It therefore controls the peroxisomal beta-oxidation pathway of fatty acids. Key regulator of adipocyte differentiation and glucose homeostasis”. Thus, within the specific context of relationship to a specific GPCR, the context-based mining approach improves this annotation by indicating that PPAR-γ and GPR41 are closely associated with response to nutrients, as also supported by recent research (Xiong et al., 2004).

4.1.4 Integrated co-expression and interaction networks

As an illustration of combining gene expression data with the existing interactions, pathways and ontologies, we have utilised gene expression data from mouse congenic strains in a study related to Type 1 Diabetes (Eaves et al., 2002). Type 1 Diabetes (T1D) is an autoimmune disease caused by destruction of pancreatic beta cells. Non Obese Diabetic (NOD) mouse (Makino et al., 1980) is a model of autoimmune diseases; it develops spontaneous autoimmune diabetes, which shares many similarities with T1D in humans. Hence, NOD mouse serves as an excellent animal model for T1D (Anderson and Bluestone, 2005).

“A congenic strain is a strain identical or almost identical to a standard inbred strain except for the presence of a chromosome segment introduced by appropriate crosses from a second strain. A strain is usually not regarded as congenic unless there have been at least eight crosses to the inbred strain” (Snell, 1978). Eaves et al. (2002) utilised microarray profiling of a NOD mouse, four NOD-derived diabetes-resistant congenic strains, and two non-diabetic control strains, to explore functional links between genotype and phenotype for T1D, using a novel method for differential expression analysis.

Here, the objective is to examine the correlation network of the gene expression profiles, link that information with available interactions and ontologies, and study the emergent patterns. The analysis is performed as explained in Article II. The resulting network is shown in Figure 9 of Article II in which, some relevant entities in network are indicated with their names.

The largest upregulated cluster is related to lipid and glucose metabolism. Interestingly, the upregulated BRCA1 and BRCA2 genes are also placed within this cluster. BRCA genes are associated with breast cancer, but are known to be highly expressed in spleen and associated with immune response. How these genes specifically relate to Type 1 Diabetes is unclear and requires further study. Another upregulated small cluster of genes is found to be associated with beta-cell proliferation, which is a known response to increased rate of beta-cell apoptosis in Type 1 Diabetes.

4.1.5 Interaction neighbourhood in experimental context

In the context-based mining example given in a previous section (4.1.3), unit weights were used for all interactions to define the context. In general, as described in section 3.1.6, choosing proper weights to represent a context is a difficult task and most often an exploratory undertaking. Experimental data, such as gene expression or metabolomics experiments, can also be utilized to define a specific context. In such cases the distance measure relating biological entities in the molecular profile space
may correspond to the measure of co-expression (such as correlation coefficient) between different entities.

To demonstrate the use of MegNet based mining using gene co-expression to define the context, we constructed an integrated network of interaction neighbourhood of mouse proteins PPAR-γ, PPAR-α, PGC1-α, GPR40, GPR41, and GPR43, together with co-expression network based on the gene expression data from spleens of various NOD related strains of mice (Eaves et al., 2002). We then used a combination of the correlation coefficient values and unit weights for the edges to define the conceptual space, and visualised the similarities using CDA mapping (Fig. 3 of Article III).

Interestingly, several tumor suppressor genes such as BRCA1 associated with PPAR-γ, are found in this mapping. This finding deserves further attention. Only recently a link between a specific tumor suppressor (LKB1) and diabetes has been established, linking cancer and physiological control of metabolism (Shaw et al., 2005).

### 4.2 Dynamic topology of integrated networks

In this section, we present a study of dynamic topology changes. In order to study the dynamic changes in the topology of functional modules in an integrated network, we developed a method called Topological Enrichment Analysis of Functional Subnetworks (TEAFS; section 3.2). TEAFS performs topological analysis of MegNet networks using transcriptomics data. We applied TEAFS to study dynamic responses of Oxidative stress in yeast, and validated the results with comprehensive in vivo metabolomics analysis (Article V).

#### 4.2.1 Oxidative stress

Oxidants are normally produced by aerobic metabolism, but are produced at elevated rates under pathophysiological conditions. Oxidative stress is defined as an imbalance between oxidants and antioxidants in favour of the oxidants, potentially leading to damage (Sies, 1997). Oxidative stress has been implicated in many human diseases such as atherosclerosis, diabetes, cancers, cardiovascular diseases, Parkinson’s diseases, Alzheimer’s disease, and aging (Sohal and Weindruch, 1996; Christen, 2000; Aruoma, 1998; Maritim et al., 2002; Miwa et al., 2008). Therefore, studying the phenomenon of oxidative stress is very important.

Saccharomyces cerevisiae or Baker’s yeast shares remarkable similarities with higher eucaryotes and has served as a valuable model organism (Botstein et al., 1997) in facilitating understanding of numerous human diseases. It has been used as a model organism to study the effects of oxidative stress on aging (Gonidakis and Longo, 2008).

#### 4.2.2 Transcriptomic response to Oxidative stress

Gasch et al. (2000) studied the responses, in terms of changes in transcript abundances over time, of Saccharomyces cerevisiae to a set of diverse environmental stresses. This set includes oxidative stress which was induced by growing cells to early-log phase and adding Hydrogen peroxide ($H_2O_2$) to a concentration of 0.30mM. Samples from this culture were collected at 10, 20, 30, 40, 50, 60, 80, 100, and 120 minutes and analysed with two-colour DNA microarrays (Schena et al., 1995). The resultant pattern of mRNA level responses to $H_2O_2$ treatment were characterised by the strong induction of genes that are involved in the detoxification of $H_2O_2$ superoxides such as superoxide dismutases, glutathione peroxidases, and thiol-specific antioxidants, as well as genes involved in oxidative and...
reductive reactions within the cell including thioredoxin, thioredoxin reductases, glutaredoxin, and glutathione reductase (Gasch et al., 2000).

4.2.3 Topological response to Oxidative stress

The goal of the study presented in Article V is to gain an increased understanding of the dynamic response of the oxidative stress at the level of biological function. We integrated the transcriptomic information with interactome topology, formulating a method called Topological Enrichment Analysis of Functional Subnetworks (TEAFS), and showed that TEAFS analysis of topological changes derived from transcriptomics changes outperformed the traditional Gene Set Enrichment Analysis (GSEA) which is based on transcriptomic changes alone (Subramanian et al., 2005).

Metabolomics study

In order to assess the validity of the results of TEAFS analysis, we performed comprehensive study of metabolite concentrations during the course of oxidative stress on yeast. We conducted the metabolomics experiments with protocols for cell cultivation and $H_2O_2$ treatment identical to Gasch et al. (2000). Lipid profiling, primary metabolite and fatty acid profiling were done with Ultra Performance Liquid Chromatography coupled with Mass Spectrometry (UPLC/MS), High Performance Liquid Chromatography and Mass Spectrometry (HPLC/MS) and Gas Chromatography (GC) respectively (Goodacre et al., 2004; Orešić et al., 2006).

TEAFS analysis

An outline of TEAFS analysis is depicted in Figure 1 of Article V and a brief summary is provided here. First, we constructed a yeast genome-wide interaction network by integrating information pertaining to protein-protein interactions (DIP) (Xenarios et al., 2000), metabolic reactions (KEGG) (Kanehisa et al., 2006), genes encoding the proteins (EMBL, UniProt) (Stoesser et al., 1999; Bairoch et al., 2005), and transcriptional regulation (TRANSFAC) (Wingender et al., 2000; Matys et al., 2006) using MegNet. Next, this network was reconstructed corresponding to each time point in the oxidative stress experiment, by employing the gene detection criteria (i.e., presence/absence criteria) similar to Luscombe et al. (2004), (Article V). Next, functional modules were identified by making use of Gene Ontology Biological Process term annotations (Ashburner et al., 2000) available in the UniProt protein database (Bairoch et al., 2005). Finally, the test statistic associated with TEAFS (Article V) was computed for each module, significance of the statistic was computed using a permutation test (Moore and McCabe, 2005), and False Discovery Rate $q$-values were computed to account for multiple hypothesis testing (Benjamini and Hochberg, 1995; Shaffer, 1995). The results of TEAFS were compared to those of GSEA.

Results

TEAFS found changes in modules involved in environmental stress responses including oxidative stress response. These modules include regulation of cell cycle and check points, response to DNA damage stimulus (i.e., repair mechanisms), cell wall organization, pentose phosphate shunt, biosynthesis of stress protectors (i.e., glycogen and trehalose), signal transduction pathways, post-translational modifications, regulation of transcription and vacuolar acidification. In comparison, GSEA failed to identify many relevant changes.

Analysis of primary metabolites revealed consistent increase in the levels of trehalose-6-phosphate and decrease in that of pyruvate and mannose-6-phosphate during oxidative stress with respect to $S$. 43
cerevisiae under normal conditions. Analysis of fatty acids revealed consistent increase in the levels of palmitic acid (C16:0) during oxidative stress with respect to S. cerevisiae under normal conditions and increase in relative palmitate concentration over time. Analysis of lipids revealed increase in average ceramide concentrations over time. Lipid level changes, especially the ceramide and phospholipid levels, and the changes in functional modules—particularly lipid metabolism, phospholipid biosynthesis and ceramide biosynthesis—detected by TEAFS are in mutual accordance (Article V).

Fatty acid analysis results showed a significant increase in the levels of palmitic acid (16:0). Palmitate is a precursor of de novo ceramide biosynthesis which involves fatty acid elongation. The fatty acid elongase 3-ketoreductase (IFA38), which is encoded by the gene YBR159w, was identified as a hub protein in our integrated reference network which was absent at all time points under oxidative stress. The YBR159w mutant shows characteristic accumulation of ceramides and related reactive sphingolipids similar to other mutants with defects in fatty acid elongation (Han et al., 2002). This may indicate that accumulation of palmitate, a substrate to elongase system involving IFA38, and subsequent accumulation of ceramides are in part consequences of IFA38 response to oxidative stress.
Chapter 5

Summary and Conclusions

Post-genomic molecular biology is a data-rich field of research, and a variety of such data is publicly available. The goals of this thesis are: developing methods to make use of the available data in order to produce knowledge and to aid mining of newly generated data. Seeking to provide data mining for systems biology, the research has been built up on integration of heterogeneous types of data. This thesis has made contributions in the areas of data integration, visualisation, mining, and study of network dynamics, which are important elements of systems biology (Figure 1.1).

This thesis has presented an integrated database which is a consolidation of a number of heterogeneous biological databases, and a software system called MegNet that enables retrieval and visualization of biological relationships across the data sources. Data traversals, an approach to linking interactions of heterogeneous types based on the identity of proteins, form the basis for the construction of integrated networks of interactions retrieved from multiple databases, and the MegNet software system allows visualisation of the networks (Article I). Context-sensitive mining of the data is facilitated by representing the data as conceptual spaces (Article IV), and visualising the similarities using dimension reduction (projection) techniques. As demonstrated by the results presented in section 4.1, Article I, Article II, and Article III, MegNet based data mining approach may facilitate discovery of novel or unexpected relationships, formulation of new hypotheses, data annotation, interpretation of new experimental data, and construction and validation of new network-based models of biological systems. High-throughput experimental techniques of postgenomic era are poor at quality and reproducibility. Therefore, integration of multiple types of data is desirable for bringing more confidence into analysis. Gene function prediction using MegNet has the potential to offer higher confidence as well as context-sensitivity.

The study of dynamic topological response of Saccharomyces cerevisiae to oxidative stress (Article V) was based on the integration of transcriptomics and interactomics to predict altered biological processes. Comprehensive metabolomics was used to validate the integrative analysis. In this study, by performing an integrative analysis of transcriptomics and interactomics data, we have showed that the connectivity of the Saccharomyces cerevisiae cellular network is being dynamically modulated in response to oxidative stress, leading to progressive accumulation of (lipo)toxic lipids such as ceramides (Article V, section 4.2). Our approach takes advantage of connectivity of functional modules in heterogeneous interactome network constructed by MegNet and shows that connectivity based approach is superior to traditional pathway analysis. The findings from this study establish the applicability of our network analysis strategy, and support the hypothesis that modelling of local network topology dynamics can be used as an effective tool to study the activity of biological modules.
Omics data is ever-expanding and this poses challenges to updation and curation of data in datawarehousing approaches for data integration such as ours. It is not possible to completely avoid these problems, but by taking standards-based approach to data integration, the overheads in e.g., keeping the schemas up-to-date can be reduced to some extent. In this thesis, we have relied largely on XML-based biological data exchange standards such as PSI-MI (Hermjakob et al., 2004), SBML (Hucka et al., 2003), KGML (Kanehisa et al., 2004) and so on. However, the diversity of the data, and the fact that not all data sources adapt the standards forces us to create our own schemas and write ad-hoc parsers in many cases. As explained in Articles I–II, we adapt the source schemas directly if available or we try to keep the problems arising from frequent schema changes to the minimum by extracting only relevant parts of the data to our databases. We adapted a combination of multiple approaches in data integration. Although we imported all the databases to the local warehouse, the individual schemas were kept intact. We only created an additional semantic mapping called maps database to facilitate resolution of entities across databases, which often doesn’t need to change even when a new data source is added. The integration of data across databases and sophisticated queries are handled by java programs in the middle tier.

The results of the data integration techniques such as data traversals presented in this thesis are all from yeast, human or mouse. But the technique of data traversals is applicable more broadly to any organism for which we have large scale protein annotations and interactomics data such as gene regulatory reactions, metabolic reactions and protein-protein interactions. As protein identifiers are the central to the data traversals, the data traversals are possible if the different interaction databases use consistent identifiers or if the identifiers used in the interaction databases can be mapped to one common type of identifiers. We used Uniprot accession numbers are the standard identifiers for proteins, mainly because Uniprot is the primary database for protein annotations.

It is well known that publicly available molecular biology data include many false positives and thus quality filtering of these data is essential. There are exhaustive manual curation efforts (Keshava Prasad et al., 2009) as well as sophisticated filtering approaches to address these problems (Wu et al., 2009). In this thesis, however, we have not considered these aspects. Using high quality source data would enhance the value of our approaches.

**Future work**

Some topics of future developments include: extending Topological Enrichment Analysis of Functional Subnetworks (TEAFS; sections 3.2 1.2) to be applicable to many commonly used experimental designs, extending gene detection criteria for many commonly used experimental platforms, and more fundamentally, defining a variety of interesting topological enrichment hypotheses and defining new schemes for computing enrichment score, to address these new hypotheses.

In this work we considered a variety of heterogeneous data for integrative modelling. However, integrating larger variety of data will enhance our abilities in more accurately modelling the biological systems (Kim et al., 2010). For example, Bauer-Mehren et al. (2009) demonstrated that integration of Single Nucleotide Polymorphism (SNP) data enables us to study the impact of the functional effect of SNPs in the structure and dynamics of biological networks. Thus, integration of SNPs would clearly be a natural extension to our framework.
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Title

Data integration, pathway analysis and mining for systems biology

Abstract

Post-genomic molecular biology embodies high-throughput experimental techniques and hence is a data-rich field. The goal of this thesis is to develop bioinformatics methods to utilise publicly available data in order to produce knowledge and to aid mining of newly generated data. As an example of knowledge or hypothesis generation, consider function prediction of biological molecules. Assignment of protein function is a non-trivial task owing to the fact that the same protein may be involved in different biological processes, depending on the state of the biological system and protein localisation. The function of a gene or a gene product may be provided as a textual description in a gene or protein annotation database. Such textual descriptions lack in providing the contextual meaning of the gene function. Therefore, we need ways to represent the meaning in a formal way. Here we apply data integration approach to provide rich representation that enables context-sensitive mining of biological data in terms of integrated networks and conceptual spaces. Context-sensitive gene function annotation follows naturally from this framework, as a particular application. Next, knowledge that is already publicly available can be used to aid mining of new experimental data. We developed an integrative bioinformatics method that utilises publicly available knowledge of protein-protein interactions, metabolic networks and transcriptional regulatory networks to analyse transcriptomics data and predict altered biological processes. We applied this method to a study of dynamic response of Saccharomyces cerevisiae to oxidative stress. The application of our method revealed dynamically altered biological functions in response to oxidative stress, which were validated by comprehensive in vivo metabolomics experiments. The results provided in this thesis indicate that integration of heterogeneous biological data facilitates advanced mining of the data. The methods can be applied for gaining insight into functions of genes, gene products and other molecules, as well as for offering functional interpretation to transcriptomics and metabolomics experiments.
Systems biology aims to facilitate understanding of cellular behaviour in terms of interactions among cellular components and bio molecules, and the dynamics resulting from the interactions. Systems biology studies typically involve perturbation of a biological system, application of high-throughput omics technologies to measure biological components and interactions among them, and finally integration of the data into a biological network to understand the system’s behaviour. Thus, data integration is a necessary step in systems biology. Data integration forms a basis for data mining, visualisation, as well as study of dynamics. This thesis presents methods for integration and mining of heterogeneous biological data, and for study of dynamic changes in response to system perturbations using these data.