Systems biology

MetaboNetworks, an interactive Matlab-based toolbox for creating, customizing and exploring sub-networks from KEGG

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ABSTRACT

Summary: MetaboNetworks is a tool to create custom sub-networks in Matlab using main reaction pairs as defined by the Kyoto Encyclopedia of Genes and Genomes and can be used to explore transgenomic interactions, for example mammalian and bacterial associations. It calculates the shortest path between a set of metabolites (e.g. biomarkers from a metabolomic study) and plots the connectivity between metabolites as links in a network graph. The resulting graph can be edited and explored interactively. Furthermore, nodes and edges in the graph are linked to the Kyoto Encyclopedia of Genes and Genomes compound and reaction pair web pages.

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1 INTRODUCTION

Investigating the dynamic metabolic responses in living systems because of external perturbations gives complementary information to genomic and/or proteomic approaches. This field is known as metabolomics (Nicholson et al., 1999) and provides a ‘top-down’ view of the living system to show global changes instead of cell-specific interactions between genes, proteins and/or metabolites. Metabolic pathway information can aid the interpretation of biological changes indicated by significant differences in metabolite concentration identified in metabonomic studies. The effective transformation of metabolic spectroscopy data to biological knowledge presents a significant bioinformatics challenge. In particular, the increased interest in clinical applications of metabolic phenotyping makes practical data visualization in a biological/medical framework of great importance.

The Kyoto Encyclopedia of Genes and Genomes (KEGG) (Kanehisa and Goto, 2000) is an online resource where the interaction information of genes, proteins and metabolites is integrated and it can be used to investigate molecular networks in specific organisms or for all (Kanehisa et al., 2012). KEGG provides many static pathways and metabolic reaction networks as well as a global metabolic map. The standalone application NeAT (Faust et al., 2010) can be used to draw custom metabolic reaction sub-networks for a specific organism. However, it is useful to have the ability to edit graphs, as can be done for pathways e.g. using KEGGParser (Arakelyan and Nersisyan, 2013) in Matlab. MetaboNetworks aims to fill this gap for metabolic reaction networks and combine the ability to draw custom maps and being able to edit them.

Rarely do metabolic networks and programs consider that the typical mammal is not a single organism, but a system comprising of a combination of mammalian, bacterial and potential parasitic organisms. Specifically, there is symbiosis between a host organism and its gut microbiota, and that gut microbes have functions and enzymes that are not found in mammalian organisms (Gill et al., 2006; Nicholson et al., 2012). Therefore, it is important to be able to combine metabolic reaction networks for different organisms. MetaboNetworks has the option to include data from multiple organisms and investigates if reactions can occur in any of the selected organisms. In the present study, we have illustrated the new approach using an NMR-based toxicological biomarker input dataset. This is an area of particular importance as metabolic phenotyping has shown to be particularly useful for studying toxicological processes as models for human disease (Nicholson and Wilson, 2003; Nicholson et al., 2002).

MetaboNetworks only requires the basic version of Matlab (The Mathworks, Natick, MA, USA) and does not require any additional toolbox. Although the MatlabBGL toolbox (Gleich, 2006) has a number of graph layout algorithms in mex/c++ code that can be used to speed up the calculation of large graphs in MetaboNetworks, all MetaboNetworks’ functionalities are implemented using a simple graphical user interface and Matlab code.

2 METHODS AND FEATURES

First, the appropriate data from KEGG has to be imported to Matlab. In MetaboNetworks this is done using a function that uses the KEGG REST-API to calculate a metabolite adjacency matrix that can later be used to draw the graphs. The user can select one or multiple organisms for which complete genomes are available in KEGG; for these organisms a list of enzymes (with E.C. numbers) that are associated with a gene from any of the organisms is determined. Using this enzyme list, all reactions are queried and enzymes involved in the reactions are matched against the enzyme list. Only reactions that require an enzyme from the list or that are listed as ‘non-enzymatic’ or ‘spontaneous’ are used to find their main
reaction pairs. The compounds from these reaction pairs are considered adjacent. Each row/column in the adjacency matrix indicates a specific compound (with a KEGG compound ID) and a list of all names for these compounds are found from the KEGG compound database. A reaction database has previously been collected using a similar approach (Ma and Zeng, 2003) to MetaboNetworks, however, that database includes reactions from all species, whereas MetaboNetworks focusses on organisms of interest as not all reactions can occur in all organisms.

Second, when the data collection is complete, MetaboNetworks can be used to create and explore custom networks. A list of metabolites, e.g. biomarkers arising from a metabonomic experiment, can be passed to MetaboNetworks and it searches for the shortest path between each of these metabolites using the breadh-first search algorithm. All compounds that are a part of a shortest path between any of the metabolites are included in the network. By default, MetaboNetworks plots the network as a circular graph. Other graph layouts include a spring-embedded layout, high-dimensional embedding and two types of uniform edge-length layouts, the last aim to place nodes with as little overlap as possible. If the Matlab statistical toolbox is installed, multidimensional scaling can also be used.

Last, when the initial network layout is satisfactory the graph layout can be manually adjusted. Supported adjustments include node position, node/edge removal, highlighting nodes (see green edges of nodes in Fig. 1), and shortest paths (orange edges in Fig. 1), node text and node/edge/ text properties (font, width, size, etc.). If additional data is supplied, the association of the metabolites with a response variable can be shown as node colour (see Fig. 1). Furthermore, the network can be exported as a tif, png, pdf, eps or other image formats, the network can always be reset to the original graph (all changes are lost). Another option is to click on a node to open a web browser showing the compound entry in KEGG or show reactions pairs in KEGG of selected nodes. The Supplementary Information includes a full walkthrough of the software and all the capabilities.

3 DISCUSSION
MetaboNetworks can be used to investigate and modify complex metabolic reaction networks. It improves the visual interpretation of metabonomic experiments with coverage across disparate metabolic pathways. The network should ideally be viewed as a connected network of probabilistic events (reactions) (Wilson and Nicholson, 2003) that lie on the shortest path between two metabolites. The database can be tailored to the study by including specific organisms, as we have done for the hydrazine toxicity network by including all bacteroidetes and firmicutes species, as it has been shown that germ-free rats react differently to hydrazine (Swann et al. 2009). MetaboNetworks can possibly also be used for identifying unknown metabolites by generating a network of metabolites correlated to the unknown metabolite. The unknown can be part of the shortest path between the known metabolites as the correlation can be due to a similarity in structure.

Note that use of the KEGG-API is free for academic use; however, non-academic users should ask for permission (see http://www.pathway.jp/).
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**REFERENCES**


