**Data and text mining**

**KDDN: An open-source Cytoscape app for constructing differential dependency networks with significant rewiring**

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**ABSTRACT**

Summary: We have developed an integrated molecular network learning method, within a well-grounded mathematical framework, to construct differential dependency networks with significant rewiring. This knowledge-fused differential dependency networks (KDDN) method, implemented as a Java Cytoscape app, can be used to optimally integrate prior biological knowledge with measured data to simultaneously construct both common and differential networks, to quantitatively assign model parameters and significant rewiring p-values, and to provide user-friendly graphical results. The KDDN algorithm is computationally efficient and provides users with parallel computing capability utilizing ubiquitous multi-core machines. We demonstrate the performance of KDDN on various simulations and real gene expression datasets, and further compare the results with those obtained by the most relevant peer methods. The acquired biologically plausible results provide new insights into network rewiring as a mechanistic principle and illustrate KDDN’s ability to detect them efficiently and correctly. While the principal application here involves microarray gene expressions, our methodology can be readily applied to other types of quantitative molecular profiling data.

**Availability:** Source code and compiled package are freely available for download at http://apps.cytoscape.org/apps/kddn

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**Supplementary Information:** Supplementary data are available at Bioinformatics online.

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**1 INTRODUCTION**

Modeling biological networks is an important tool in systems biology to study the orchestrated activities of gene products in cells (Mitra, et al., 2013). Significant rewiring of these networks provides a unique perspective on the changing dynamics that occur in biological systems. Thus, instead of asking “which genes are differentially expressed”, a more interesting question is “which genes are differentially connected?” (Hudson, et al., 2012; Mitra, et al., 2013). To systematically characterize selectively activated or deactivated regulatory components and mechanisms, the modeling tools must effectively distinguish significant rewiring from random background fluctuations. While specific biological networks cannot be constructed by existing knowledge alone, novel incorporation of prior knowledge into data-driven approaches can improve the robustness and biological relevance of network inference.

**Figure 1. Architecture and flowchart of KDDN software.**

Differential Dependency Network (Zhang, et al., 2009; Zhang and Wang, 2010) and its Knowledge-fused extension (Tian, et al., 2014; Tian, et al., 2011) KDDN have been developed to infer biological networks with significant rewiring by integrating experimental data and biological knowledge. The unique and attractive features of the KDDN software tool are: (1) it is easy to use with a user-friendly Graphic User Interface (GUI) and cutting-edge performance; (2) both conserved and differential biological networks can be inferred via efficient closed-form numerical solutions; (3) model parameters are determined statistically aligned with the expected performance; (4) prior knowledge (condition specific or nonspecific) is incorporated for inferring dependency networks under different conditions; and (5) statistical significance on the differential connections and the type I error rate are rigorously assessed.

The KDDN Cytoscape 3.x app adapts and extends recent KDDN algorithms in the literature (Tian, et al., 2014; Tian, et al., 2011; Zhang, et al., 2009; Zhang and Wang, 2010) (Fig. 1). Via a user-
friendly GUI, users can easily install KDDN software and perform network analysis by just a few clicks. KDDN software has been tested thoroughly on both synthetic and real gene expression data, and has been successfully applied to a wide range of research projects including yeast oxidative stress response, breast cancer recurrence, muscular dystrophy, and estrogen exposures. Helpful tips for users to use the software in an efficient manner yet with fully informed limitations are provided in Supplementary Information.

2 DESCRIPTION

2.1 Methods and software

KDDN algorithm jointly learns the conserved biological network and statistically significant rewiring across different conditions. Condition-specific data and prior knowledge are quantitatively fused via an extended Lasso model with \( \ell_1 \)-regularized convex optimization formulation (Tian, et al., 2014). Based on the unique nature of the problem, we derive an efficient closed-form solution for the embedded sub-problem solved by the block-wise coordinate descent algorithm. We conduct the computational complexity analysis on the KDDN algorithm (Supplementary Information). Since existing knowledge is often nonspecific or imperfect, KDDN uses a “minimax” strategy to maximize the benefit of prior knowledge while confining its negative impact under the worst-case scenario. Furthermore, KDDN matches the values of model parameters to the expected false positive rates on network edges at a specified significance level, and assesses edge-specific p-values on each of the differential connections (Tian, et al., 2014).

KDDN is implemented as an open-source cross-platform Cytoscape 3.x app with parallel computing capability in Java. Control panel will guide users to navigate through the experiment settings that allow flexible configuration of the analytic tasks. KDDN software takes the quantitative expression values of relevant genes/proteins as input, and incorporates the prior knowledge applicable to either or both conditions (e.g., KGML files downloaded from KEGG pathway website can be directly imported as prior knowledge). In addition to inferring common and/or significant rewiring with edge-specific p-values, conventional differential analysis is simultaneously performed allowing users to compare expression fold-changes and network rewiring side-by-side. The constructed networks are visualized seamlessly in Cytoscape and detailed numerical results together with model parameters are presented in dedicated panels (Fig. 2). All experimental settings and results can be conveniently exported for further analysis.

2.2 Case study

We applied KDDN to analyze the network rewiring in budding yeast *Saccharomyces cerevisiae* in response to oxidative stress, focused on cell cycle related genes. Integrating the prior biological knowledge in the KEGG yeast pathway and gene expression data (GSE7645), the significant differential networks constructed by KDDN are given in Fig. 2, where red edges indicate the connections existing under control and green edges indicate the connections created under stress, exclusively.

Oxidative stress is a harmful condition in a cell, tissue, or organ, caused by an imbalance between reactive oxygen species and other oxidants and the capacity of antioxidant defense systems to remove them (Lee, et al., 2011). The result shows that *Yap1*, *Rho1* and *Msn4* are among differential hubs and at the center of the network response to oxidative stress; they are activated under oxidative stress and many connections surrounding them are created (green edges). *Yap1* is a major transcription factor that responds to oxidative stress; *Msn4* is considered as a key responder to environmental stresses including oxidative stress; *Rho1* is known to resist oxidative damage and facilitate cell survival; *Ctt1* acts as an antioxidant and it coordinates with *Yap1* to protect cells from oxidative stress. The stress-induced interaction between *Hog1* and *Fus1*, detected by KDDN, is also observed in an earlier study. The biologically plausible results suggest not only the network rewiring as a mechanistic principle in determining phenotypes but also KDDN’s ability to detect them efficiently and correctly.

3 DISCUSSION

The KDDN Cytoscape app presents an integrated software tool to construct biological networks and detect significant rewiring across different conditions. Supported by a well-grounded mathematical framework, KDDN integrates the abundant biological knowledge and condition-specific experimental data to depict the overall dependency networks and their dynamics. Tested on both simulations and real gene expression data, KDDN outperforms peer methods (Supplementary Information) and demonstrates its effectiveness in revealing significant topological changes. We expect KDDN to be a very useful tool for performing differential network analysis in many biological contexts (Mitra, et al., 2013).

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REFERENCES


