**INfORM: Inference of NetwOrk Response Modules**

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**Abstract**

Summary: Detecting and interpreting responsive modules from gene expression data by using network-based approaches is a common but laborious task. It often requires the application of several computational methods implemented in different software packages, forcing biologists to compile complex analytical pipelines. Here we introduce INfORM (Inference of NetwOrk Response Modules), an R shiny application that enables non-expert users to detect, evaluate and select gene modules with high statistical and biological significance. INfORM is a comprehensive tool for the identification of biologically meaningful response modules from consensus gene networks inferred by using multiple algorithms. It is accessible through an intuitive graphical user interface allowing for a level of abstraction from the computational steps.

**Availability and implementation:** INfORM is freely available for academic use at https://github.com/Greco-Lab/INfORM.

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

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**1 Introduction**

Co-expression gene network analysis has become popular for analyzing DNA microarray and RNA sequencing data, for instance, to detect gene modules. A number of software solutions are available offering some important functionalities, but no tool to date implements a robust ensemble strategy for network inference (Supplementary Table S1). Here, we introduce INfORM (Inference of NetwOrk Response Modules), an R-Shiny (Chang et al., 2017) application to assist performing all the computational steps needed to infer relevant responsive modules from transcriptomic data. Its graphical interface provides the user with a layer of abstraction and enables to analyze data without obligatory knowledge of gritty technical and statistical aspects. First, robust gene co-expression network is derived by combining multiple network inferences from an ensemble of methods (Marbach et al., 2012). Second, gene modules are identified in the network by using community detection algorithms and evaluated based on multiple metrics of node importance. Third, the similarity between the identified modules is computed based on the overrepresented gene ontology (GO) terms in each module. Based on this information, the user can select a set of gene communities and merge them into a response module. The workflow of INfORM is depicted in Figure 1.

**2 Methods and features**

INfORM requires two mandatory input files: i) a gene expression data table; ii) a list of significantly differentially expressed genes. Gene symbols (Yates et al., 2017) must be used as row names for
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Both the inputs. Optionally, the user can also upload directly one or more adjacency matrices inferred from any network inference algorithm. Then, the user is guided through the following steps: i) network inference; ii) module detection, iii) module evaluation and selection based on multiple gene-rank scores and overrepresented GO terms. The R functions for each step of the analysis are described in details in Supplementary Table S2. The user guide is also provided in Supplementary File S1.

2.1 Gene network inference

In INIFORM, an ensemble of mutual information-based methods is used to infer gene co-expression networks. A compendium of networks is generated by using all possible distinct combinations of inference algorithms [ARACNE (Margolin et al., 2006), MRNET (Meyer et al., 2007), MRNETb (Meyer et al., 2010) and CLR (Faith et al., 2007)], mutual information and entropy estimators, and discretization methods available in the R package MINET (Meyer et al., 2008), and integrated into the final network. These inferences can be integrated or even replaced by pre-computed adjacency matrices provided by the user. First, inferences by the same algorithm are reduced by taking the median, the mean or the maximum of the estimated edge weights. An evaluation of the effect of different summarization strategies is reported in Supplementary File S2.

Second, the ranked connections from the summarized networks are aggregated by using the Borda method implemented in TopKLists R package (Schimek et al., 2015) and the top ranked edges are progressively included in the final network until all the nodes are incorporated, i.e. all the nodes have degree \( \geq 1 \) (Supplementary File S2). This procedure ensures the robustness of the network inference. Alternatively, the user can decide the top \( n\% \) of the ranked edges to include in the final network.

2.2 Gene module detection and responsive module definition

INIFORM provides widely used community clustering algorithms for identifying the relevant modules from the inferred gene network, the ‘Walktrap’ (Pons and Latapy, 2005), ‘Spinglass’ (Newman and Girvan, 2004; Reichardt and Bornholdt, 2006), ‘Louvain’ (Blondel et al., 2008) and the ‘Greedy’ (Clauset et al., 2004) algorithms. Benchmark analysis showed that all the algorithms generate highly similar modules (Supplementary File S2). The relevance of the identified modules is evaluated by scoring them based on characteristics of the member nodes and edges: i) centrality, ii) differential log2(fold-change), iii) differential P-value, iv) median rank of edge weights and v) number of nodes. These are graphically represented as a radar chart, making it easy to evaluate the modules. Moreover, enrichment analysis is performed to find the GO terms overrepresented in each module and compute the similarity between sets of GO terms from different modules. A heatmap representation of the GO-based module similarity is provided to aid the selection of functionally related modules. INIFORM allows the user to merge statistically significant and biologically relevant modules into an optimized response module. The biological functions associated with the response module are visualized by the means of a tile plot (Supek et al., 2011), in which the semantically similar GO terms are grouped together (Yu et al., 2010).

3 Conclusion

Here we present INIFORM, a novel tool for robust inference of gene co-expression networks from transcriptomics data. The graphical user interface helps to perform the analysis without any obligatory technical expertise and in-depth knowledge of the implementation. Results from a case study analysis of a publicly available data are provided in Supplementary File S3.

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References


Fig. 1. (A) Matrices from same inference algorithm are summarized by median, mean or maximum of the correlation scores. Ranked edges from each summarized matrix are combined by the Borda method. Finally, the top ranked edges are selected until all the nodes are covered by at least one edge. (B) Modules are identified and annotated with node ranks, edge ranks. Modules can be selected and merged into a response module. (C) GO annotations of the response module are summarized and visualized by the means of tile plots.