

CoExp Web, a web tool for the exploitation of co-expression networks

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Abstract

Summary: This paper presents the CoExp Web tool. CoExp is a software tool and a Web resource that fosters research on co-expression network models by providing a highly usable means to exploit and share multiple network models.

Availability and Implementation: the Web resource is accessible at <https://snca.atca.um.es/coexp/>. The software at the front and back-ends is accessible at https://github.com/SoniaRuiz/CoExp_Web.

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Introduction

Gene co-expression network analysis has been widely used to identify biologically important patterns in gene expression in a hypothesis-free and genome-wide manner¹⁻⁴. The driving principle behind co-expression network analysis is that genes whose expression levels are highly correlated, are also likely to share functional and biological relationships^{5,6}. Thus, co-expression networks (CEN) are models of how genes cluster together into modules of highly co-expressed genes, generally by using graph-based approaches to reveal their similarities⁷. A CEN is useful as a way to describe gene-gene relationships in a given gene expression profile which represent a given condition (e.g. a tissue, a specific cell type or a group of cases and controls). But CENs can also be useful when used for annotation of external gene sets. For example, (1) by looking at whether the genes cluster together in the network, (2) by identifying new genes co-expressed with the input genes, or (3) specific cell type markers clustering together with those genes.

Normally, managing CEN models is work intensive, and becomes highly complex when multiple CENs are being explored. In these cases, a Web based application integrating all those CENs, becomes the optimum solution. We present here CoExp, accessible at <https://snca.atca.um.es/coexp/>. CoExp is a Web application to increase the usability and accessibility of CENs. CoExp offers 109

different CENs created with the CoExpNets R package⁷, based on the widely used WGCNA R package⁸ (see a, Figure 1).

The network catalogue is organised into four different network groups: (1) the Religious Orders Study and Memory and Aging Project (ROSMAP)⁹⁻¹¹ composed of four CENs derived from RNAseq data on post-mortem human frontal cortex originating from control individuals, as well as those with cognitive impairment and Alzheimer's disease; (2) The Genotype-Tissue Expression project (GTEx, V6 and V7¹²) composed of two suites of CENs derived from RNAseq data on 47 post-mortem control human tissue samples; (3) the UK Brain Expression Consortium project (UKBEC)^{2,13} composed of ten microarray-based CENs derived from post-mortem control human brain tissue; (4) the North America Brain Expression Consortium (NABEC)¹⁴, composed of one CEN derived from RNAseq data on post-mortem control human frontal cortex (see b, Figure 1).

Network Catalogue Browser

The user can become familiar with the CEN models available by browsing, and downloading, the catalogue through the 'Network Catalogue' tab. The user selects a network category, and then a specific network. Finally, the user decides between two different views of the network: the 'Ontology Classification' or the 'Cell Type Classification'. The 'Ontology Classification' view displays an active data table in which each module (i.e. gene cluster) from the selected network occupies one row. The columns provide summarized information about annotation terms enriched for the genes in the modules. Those terms come from an enrichment analysis performed with gProfileR¹⁵, which incorporates data from well recognised ontologies, including Gene Ontology¹⁶, REACTOME¹⁷ and KEGG¹⁸. The 'Cell Type Classification' view, displays an active data table in which the rows correspond now to sets of gene markers of brain cell types tested for enrichment (Fisher's Exact test) and each module occupies a column. Each cell within the table contains the Bonferroni corrected p-value for the enrichment of the corresponding cell markers within the module. In both view types, the raw data behind the tables is easily downloadable.

Gene Set Annotation

CENs are often used to annotate a gene set of interest, in the context of a specific condition (e.g. a tissue of interest). For those users, CoExp offers the 'Gene Set Annotation' tab. Thanks to this function the user can investigate whether the gene set is enriched within a single or multiple CEN modules, across all the co-expression networks from amongst those available in the catalogue. In this way, those genes can be annotated based on how they are distributed across the network modules and their biological context explored within just a couple of clicks. The user is supplied a results table in which each row relates to a gene of interest which has been successfully found in any of the modules belonging to the network or networks selected. The columns provide information on the module in which the gene has been found, including the statistical significance of the overlap between the input genes and the genes in the module and a brief description of the module's function based on the its top five GO terms. This analysis outcome can easily be downloaded as an excel file.

Plot Network

Once the user has decided which network is of interest, and which module within the network requires detailed visualization, the corresponding gene graph can be plotted. The third tab, 'Plot Network', enables the graph-based visualisation of the genes within a network module of interest, whether identified by browsing through the catalogue or because the user's gene set of interest significantly clusters within a module. The 'Plot Network' tab generates an interactive directed graph formed by the hub (i.e. most important) genes within a module. The user can select how many of the most relevant genes will appear within the plot. The resulting plot is interactive in the sense that it can be zoomed, rotated and the direct neighbours of any gene highlighted by just clicking on the gene of

interest. Both the raw data displayed in the graph and a high quality PNG image are available for download.

Conclusions

CoExp Webpage is a web platform for the exploitation of co-expression networks. CoExp currently offers 109 CENs focused on brain transcriptomics with plans to expand its scope. It is a powerful, easy to use and innovative tool for gene set annotation across a variety of brain-specific transcriptomic datasets. CoExp makes CENs easily shareable with the scientific community. Everything is downloadable in CoExp, including the CENs themselves and the front- and back-end software so any research laboratory can construct their own CoExp web site. This makes it a powerful tool for the wider research community interested in producing, using or sharing co-expression models to support their research.

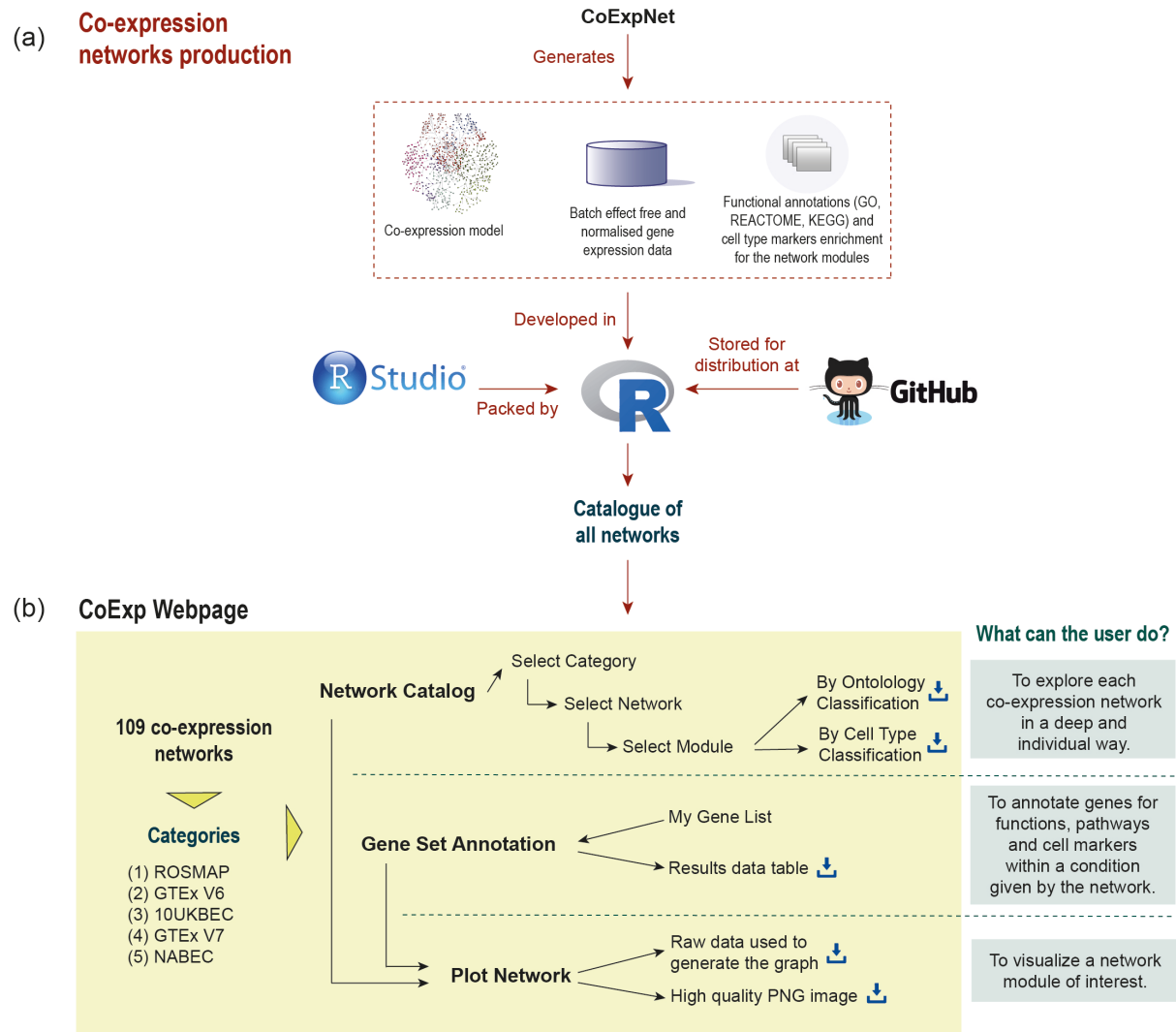


Figure 1: (a) All 109 networks in CoExp were generated with CoExpNets R package for co-expression network generation and annotation. Networks and annotations are organised into R packages (i.e. one package for each network category) and stored in GitHub. (b) There are five network categories currently available at CoExp. All of them can be browsed, used for gene set annotation and plotted by gene modules.

Funding

This research was supported in part by the Intramural Research Program of the NIH, National Institute on Aging. R.H.R. was supported through the award of a Leonard Wolfson Doctoral Training Fellowship in Neurodegeneration. J.H. and M.R. were supported by the UK Medical Research Council (MRC), with J.H. supported by a grant (MR/N026004/) and M.R. through the award of a Tenure Track Clinician Scientist Fellowship (MR/N008324/1). J.H. was also supported by the UK Dementia Research Institute, The Wellcome Trust (202903/Z/16/Z), the Dolby Family Fund, and the NIHR. A.C. is supported by Fundación Séneca - Science and Technology Agency of the Region of Murcia, (grant reference: 20762/FPI/18). A.L.G.M. is funded by Fundación Séneca (grant reference: 21230/PD/19). S.G.R. was supported by the UK Medical Research Council (MRC) through the award of Tenure-track Clinician Scientist Fellowship to MR (MR/N008324/1). R.H.R. was supported through the award of a Leonard Wolfson Doctoral Training Fellowship in Neurodegeneration.

Data availability

All co-expression models available through CoExp and their respective annotations can be downloaded from the CoExp application, accessible from <https://snca.atca.um.es/coexp/>. Alternatively, gene expression profiles and networks created from those expression profiles, for all networks available at CoExp can be obtained at GitHub in the form of R packages, one package per co-expression network family. All the packages and networks were created using CoExpNets software, based on WGCNA, accessible at <https://github.com/juanbot/CoExpNets>. GTEx networks and expression profiles can be downloaded from <https://github.com/juanbot/CoExpGTEx>. ROSMAP networks and expression profiles can be downloaded from <https://github.com/juanbot/CoExpROSMAP>. UKBEC networks and expression profiles can be downloaded from <https://github.com/juanbot/CoExp10UKBEC>. Finally, NABEC network and expression profiles can be obtained through <https://github.com/juanbot/CoExpNABEC>. The back and front ends code of the CoExp Web application is fully available for download on GitHub at https://github.com/SoniaRuiz/CoExp_Web.

Conflict of Interest: none declared.

Acknowledgements

Study data to create the ROSMAP networks were provided by the Rush Alzheimer's Disease Center, Rush University Medical Center, Chicago. Data collection was supported through funding by NIA grants P30AG10161, R01AG15819, R01AG17917, R01AG30146, R01AG36836, U01AG32984, U01AG46152, the Illinois Department of Public Health and the Translational Genomics Research Institute.

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