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Applications Note



Gene expression

myTAI: evolutionary transcriptomics with R

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Abstract

Motivation: Next Generation Sequencing (NGS) technologies generate a large amount of high quality transcriptome datasets enabling the investigation of molecular processes on a genomic and metagenomic scale. These transcriptomics studies aim to quantify and compare the molecular phenotypes of the biological processes at hand. Despite the vast increase of available transcriptome datasets, little is known about the evolutionary conservation of those characterized transcriptomes.

Results: The myTAI package implements exploratory analysis functions to infer transcriptome conservation patterns in any transcriptome dataset. Comprehensive documentation of myTAI functions and tutorial vignettes provide step-by-step instructions on how to use the package in an exploratory and computationally reproducible manner.

Availability and implementation: The open source myTAI package is available at https://github. com/HajkD/myTAI and https://cran.r-project.org/web/packages/myTAI/index.html.

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Supplementary information: Supplementary data are available at Bioinformatics online.

1 Introduction

To investigate phenotypic changes, diseases, environmental stresses, or developmental processes, transcriptome studies are often the approach of choice. Although transcriptomics is based on a solid methodology, little is known about the evolutionary conservation and dynamics of transcriptomes across species (Drost et al., 2017). Understanding the evolutionary processes that change transcriptomes over time, however, might lead to new insights on how diseases emerge or how phenotypic changes are caused by changes in transcriptomes. For this purpose, evolutionary transcriptomics studies aim to capture and quantify the evolutionary conservation of transcriptomes during specific stages of the biological process of interest (Domazet-Lošo and Tautz, 2010). Here, we present the exploratory analysis package myTAI, which can combine evolutionary information of genes with their transcript levels to infer transcriptome conservation patterns. Evolutionary information is given as input to the package and can range from classical phylogenetic or orthology relationships between genes to more recent approaches such as phylogenetic comparative methods (PCMs) (Dunn et al., 2013), phylogenetic reconciliation methods (Doyon et al., 2011), or phylostratigraphy (Domazet-Lošo et al., 2007). In summary, starting with a pre-computed table of gene age information and a transcriptome dataset, the R package myTAI can be used to screen for stages of high or low transcriptome conservation within a biological process of interest. If highly conserved or variable transcriptomes were found in particular stages or treatments, more specialized experimental studies could subsequently be designed to investigate the functions and mechanistic implications of these conserved or variable stages.

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2 Implementation

The R package *myTAI* is released under the GNU General Public License within the CRAN project (R Core Team). The package can be downloaded from https://cran.r-project.org/web/packages/myTAI/index.html. The source code is publicly available at https://github.com/HajkD/myTAI. Internal *myTAI* functions are implemented in C++ and integrated via the *Rcpp* (Eddelbuettel, 2013) Application Programming Interface (API) and unit tested using test-that. The *myTAI* package furthermore depends on the R packages *nortest*, *fitdistrplus* (Delignette-Muller and Dutang, 2015), *dplyr*, *RColorBrewer*, *taxize* (Chamberlain and Szöcs, 2013), *reshape2*, *ggplot2* (Wickham, 2009), *biomartr* (Drost and Paszkowski, 2017), *readr*, *tibble*, *scales* and *gridExtra*.

3 Functions and Examples

More than fifty functions are provided by the *myTAI* package. We recently used *myTAI* to investigate the developmental hourglass model of embryo development (Raff, 1996) on the transcriptomic level (Quint *et al.*, 2012; Drost *et al.*, 2017). Others used *myTAI* to investigate transcriptome conservation in plant organ development (Lei *et al.*, 2017). To illustrate an example workflow with *myTAI*, we here use the developmental transcriptome of *Arabidopsis thaliana* embryo development (Quint *et al.*, 2012) (see Supplementary Material for more details about data formats):

```
# Import the myTAI package and load example dataset library(myTAI); data(PhyloExpressionSetExample)
```

One metric to quantify transcriptome conservation on a global scale is the Transcriptome Age Index (TAI) (Domazet-Lošo and Tautz, 2010), which denotes the average transcriptome age throughout the biological process of interest.

```
# Plot the Transcriptome Age Index of A. thaliana embryo development
```

PlotSignature (PhyloExpressionSetExample)

To quantify the transcript level of each gene age category to the overall transcriptome for each developmental stage, the gene expression level distributions for each gene age category can be visualized by:

```
# Plot gene expression level distributions PlotCategoryExpr\ (PhyloExpressionSetExample,\\ legendName = "PS",\\ log.expr = TRUE)
```

A linear transformation of the mean expression levels into the interval [0, 1] enables the comparison of mean expression level patterns between gene age categories independent of their actual mean expression magnitude. A relative expression level of 0 denotes the minimum mean expression level compared to all other stages, and a relative expression level of 1 denotes the maximum mean expression level compared to all other stages:

```
# Plot relative expression levels

PlotRE (PhyloExpressionSetExample,

Groups = list (c(1: 3), c(4: 12)),

legendName = "PS",

adjust.range = TRUE)
```

Finally, we compare relative expression levels between groups of age categories and quantify their difference:

Compare relative expression levels between groups of age categories

PlotBarRE (PhyloExpressionSetExample,

```
Groups = list(group\_1 = 1: 3, group\_2 = 4: 12),

xlab = "Ontogeny",

ylab = "Mean Relative Expression",

cex = 1.5)
```

In addition to these exploratory functions, *myTAI* provides functionality for taxonomic information retrieval, *gene age* enrichment analyses, differential gene expression analyses of age categories, and additional metrics for quantifying trancriptome conservation. A detailed description and interpretation of *myTAI* functions is available at https://github.com/HajkD/myTAI#tutorials and also in the Supplementary Material.

4 Conclusions

Evolutionary transcriptomics studies can serve as a first approach to screen *in silico* for the potential existence of evolutionary constraints within a biological process of interest. This is achieved by quantifying transcriptome conservation patterns and their underlying gene sets in biological processes. The exploratory analysis functions implemented in *myTAI* provide users with a standardized, automated and optimized framework to investigate evolutionary signatures in any transcriptome dataset of interest.

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