Visualizing RNA-seq data: Pertinence, software, and applications

Exit Seminar Lindsay Rutter

Program of Study Committee:

Dianne Cook (Major Professor) Amy Toth (Major Professor) Heike Hofmann Daniel Nettleton James Reecy

May 1, 2018

My Background

Education

- B.S. in Bioengineering Pennsylvania State University (2003-2007)
- Major in Bioinformatics and Computational Biology Iowa State University (2012-Present)
- Minor in Statistics *Iowa State University* (2012-Present)

Internships

- Nagaoka University of Technology (Summer 2012)
- Okinawa Institute of Science and Technology (Summer 2014)
- MathWorks (Summer 2016)
- After Inc. (Fall 2016)
- Google Summer of Code (Summer 2017)
- GeneLab (Pending Summer 2018)

Chapter Outline

- Chapter 1 Visualization methods for genealogical datasets
- Chapter 2
 The case for visualization methods in RNA-seq data analysis
- Chapter 3
 Software for visualization methods in RNA-seq data analysis
- Chapter 4
 Gene expression responses to diet quality and viral infection in Apis mellifera

Chapter 1 : Visualization methods for genealogical datasets



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ggenealogy: An R Package for Visualizing Genealogical Data

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Abstract

This paper introduces genealogy (Rutter et al. 2015), a developing R software package that provides tools for searching through genealogical data, generating basis statistics on their graphical structures using parent and child connections, and displaying the results. It is possible to draw the genealogy in relation to variables related to the nodes, and to determine and display the shortest path distances between the nodes. Production of pairwise distance matrices and genealogical diagrams constrained on generation are also available in the visualization toolikt. The tools are being tested on a dataset with milestone cultivars of soybean varieties (Hymowitz et al. 1977) as well as on a web-based database of the academic genealogy of mathematicians (North Dakota State University and American Mathematical Society). The latest stable package version is available in source and binary form on the Comprehensive R Archive Network.

Keywords: genealogy, data visualization, statistical graphics, exploratory data analysis, interactive, R.

ggenealogy: Visualization Tools for Genealogical Data

Methods for searching through genealogical data and displaying the results. Plotting algorithms assist with data exploration and publication-quality image generation. Includes interactive genealogy visualization tools. Provides parsing and calculation methods for variables in descendant branches of interest. Uses the Grammar of Graphics.

Version: 0.3.0

Depends: $R (\geq 3.3.0)$

Imports: $ggplot2 (\geq 2.2.0)$, $igraph (\geq 0.7.1)$, $plyr (\geq 1.8.1)$, $reshape2 (\geq 1.4)$, $plotly (\geq 4.5.6)$, $tibble (\geq 1.2)$

Suggests: $\underline{\text{stringr}} \ (\ge 0.6.2), \underline{\text{knitr}} \ (\ge 1.13), \underline{\text{roxygen2}} \ (\ge 3.0.0), \underline{\text{dplyr}} \ (\ge 0.5.0)$

Published: 2016-12-12

Author: Lindsay Rutter, Susan Vanderplas, Di Cook

Maintainer: Lindsay Rutter https://docs.py.ncb/rutter-at-iastate.edu

License: GPL-2 | GPL-3 | [expanded from: GPL]

NeedsCompilation: no

Materials: README NEWS
CRAN checks: ggenealogy results

Downloads:

Reference manual: ggenealogy.pdf

Vignettes: ggenealogy: Visualization tools for genealogical data

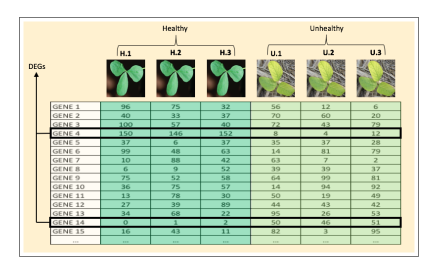
Package source: ggenealogy 0.3.0.tar.gz

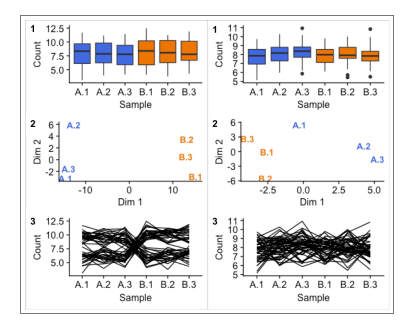
Windows binaries: r-devel: ggenealogy 0.3.0.zip, r-release: ggenealogy 0.3.0.zip, r-oldrel: ggenealogy 0.3.0.zip

OS X binaries: r-release: ggenealogy 0.3.0.tgz, r-oldrel: ggenealogy 0.3.0.tgz

Old sources: ggenealogy archive

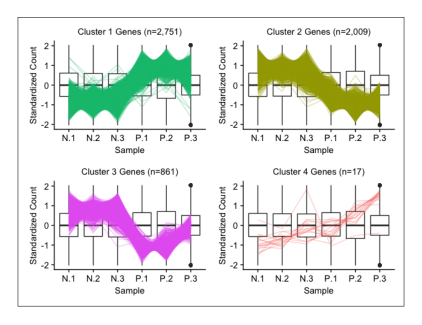
Chapter 2: The case for visualization methods in RNA-seq analysis

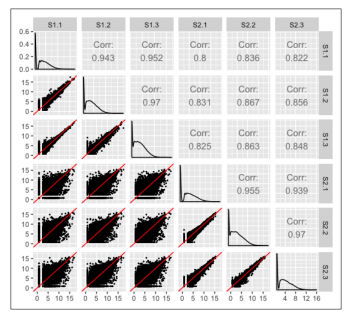




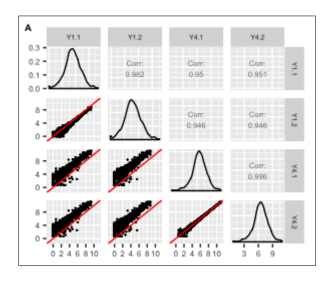
Three new plotting types

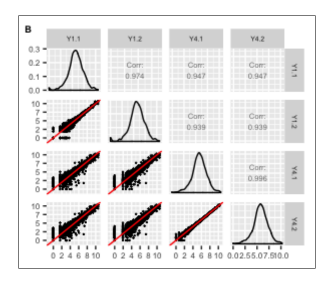
- Parallel coordinate plots
- Scatterplot matrices
- repLIcate TREatment ("litre") plots

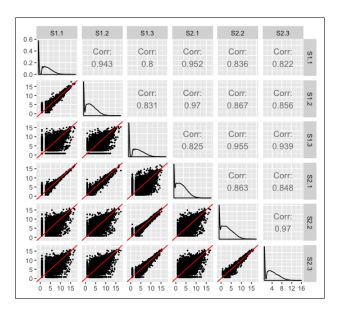


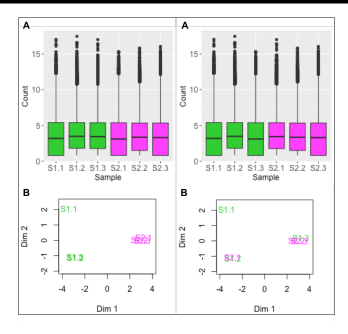


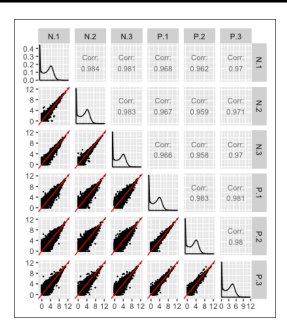
https://rnaseqvisualization.shinyapps.io/scatmat

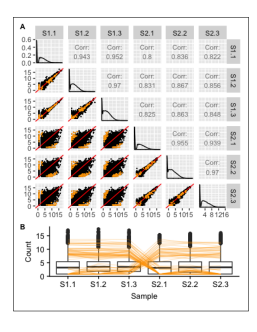


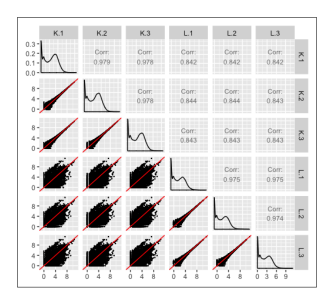


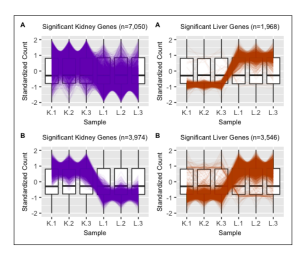


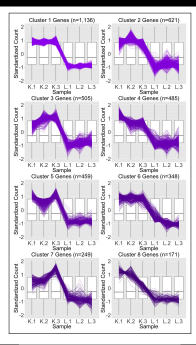












$Completed\ work$

Product	Description	Date
R package	First release of ggenealogy, which pro-	March 2015
	vides visualization tools for genealogical	
	datasets	
Presentation	Presented ggenealogy at JSM	August 2015
Award	Student paper award at ASA Statistical	August 2015
	Computing and Graphics Section	

Scheduled deliverables

Product	Description	Date
R package	Second release of ggenealogy package,	May 2016
	which provides visualization tools for ge-	
	nealogical datasets	
Paper	Submit ggenealogy paper to JSS	May 2016
R package	First release of package that provides vi-	TBD
	sualization tools for RNA-sequencing da-	
	tasets	
Paper	Submit paper about visualization tools for	TBD
	clustering analysis of RNA-sequencing	
Paper	Submit paper about visualization tools for	TBD
	significance testing of RNA-sequencing	

Other work

R package First release of ePort package that gen rates electronic reports for instructors evaluate student performance	v

References

- ggplot2 (Wickham 2009)
- GGally (Schloerke et al. 2016)
- nullabor (Wickham et al. 2014)
- ggbio (Yin et al. 2012)
- GGobi (Swayne et al. 2003)
- tourr (Wickham et al. 2011)
- plotly (Sievert et al. 2016)
- Parallel coordinate plots (Inselberg 1985, Wegman 1990)
- Visual statistical inference (Chowdhury et al. 2015)

References

- pedigree (Coster 2013)
- kinship2 (Therneau et al. 2015)
- GraphViz (Gansner and North 2000)
- Cytoscape (Shannon et al. 2003)
- explorase (Lawrence et al. 2008)
- edgeR (Robinson et al. 2010)
- DESeq2 (Love at al. 2014)
- RUVseq (Risso et al. 2014)
- Gene expression visual inference (Yin et al. 2013)
- Biological clustering : (Newell et al. 2013)

Leaves at 120 minutes - 3 Clusters