# Modular analysis of gene expression data with GNU R or Matlab

Gábor Csárdi, 1,2, Zoltán Kutalik 1,2 and Sven Bergmann 1,2

<sup>1</sup>Department of Medical Genetics, and <sup>2</sup>Swiss Institute of Bioinformatics, University of Lausanne, Rue de Bugnon 27 - DGM 328, CH-1005 Lausanne, Switzerland.

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#### **ABSTRACT**

Summary: Availability:

Contact: Sven.Bergmann@unil.ch

# 1 INTRODUCTION

What is biclustering. What is the ISA.

# 2 METHODS

Normalization.

Random and smart seeding.

The ISA iteration.

Merging the modules.

The robustness measure and filtering the modules.

Module trees.

# **3 R IMPLEMENTATION**

Two packages: isa2 does the computation, eisa does everything related to biology.

# 3.1 Workflow

Two workflows, the simple workflow does everything with default parameters. In the detailed workflow the user has access to every step of the analysis.

# 3.2 Visualization

Expression plots, GO tree plots, module tree plots, HTML summary generation.

# 3.3 Connection to other software

The biclust package.

# 4 MATLAB IMPLEMENTATION ACKNOWLEDGEMENT

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