

A. Moghaddas Gholami (1), C. Schmitt (2), M. Visvanathan (1), N.C. Hauser (2), S. Rupp (2), J. Hoheisel (1), K. Fellenberg (1)

Deutsches Krebsforschungszentrum (DKFZ), Dept. Functional Genome Analysis, Heidelberg, Germany (1)
Fraunhofer Institut für Grenzflächen- und Bioverfahrenstechnik (IGB), Dept. Molecular Biotechnology, Stuttgart, Germany (2)

Motivation

The Multi-Conditional Hybridization Intensity Processing System (M-CHIPS), provides a common platform for integrated analysis of microarray data along with supplementary information - like GO terms and clinical data. For the biologist, it integrates all major platforms such as Affymetrix-, two-color microarrays, or 2D gels, providing a guided tour through the data's variance by mouse click. On the developer's side, it integrates Perl, Matlab, and R, providing the convenient programming environments for various tasks.

Background

- M-CHIPS (www.mchips.org) is in use since 1999
- Fields of research: Cancer, Yeast (*S.cerevisiae*, *Candida albicans*, *N.crassa*), *Arabidopsis*, *Antirrhinum*, *Drosophila*, *Trypanosoma*, *Aspergillus*, *P.putida*, *H.echinata*, Mouse etc.
- Application: statistical analysis of a microarray database's entire content including the experiment annotations [1, 2]
- Advantages:
 1. Flexibility allows the use of one and the same system for different fields of microarray research where different experimental factors are relevant.
 2. Allows to add experiment annotations and/or allowed values within seconds on the fly i.e. without altering the database structure

Results

New functionalities concerning biologists:

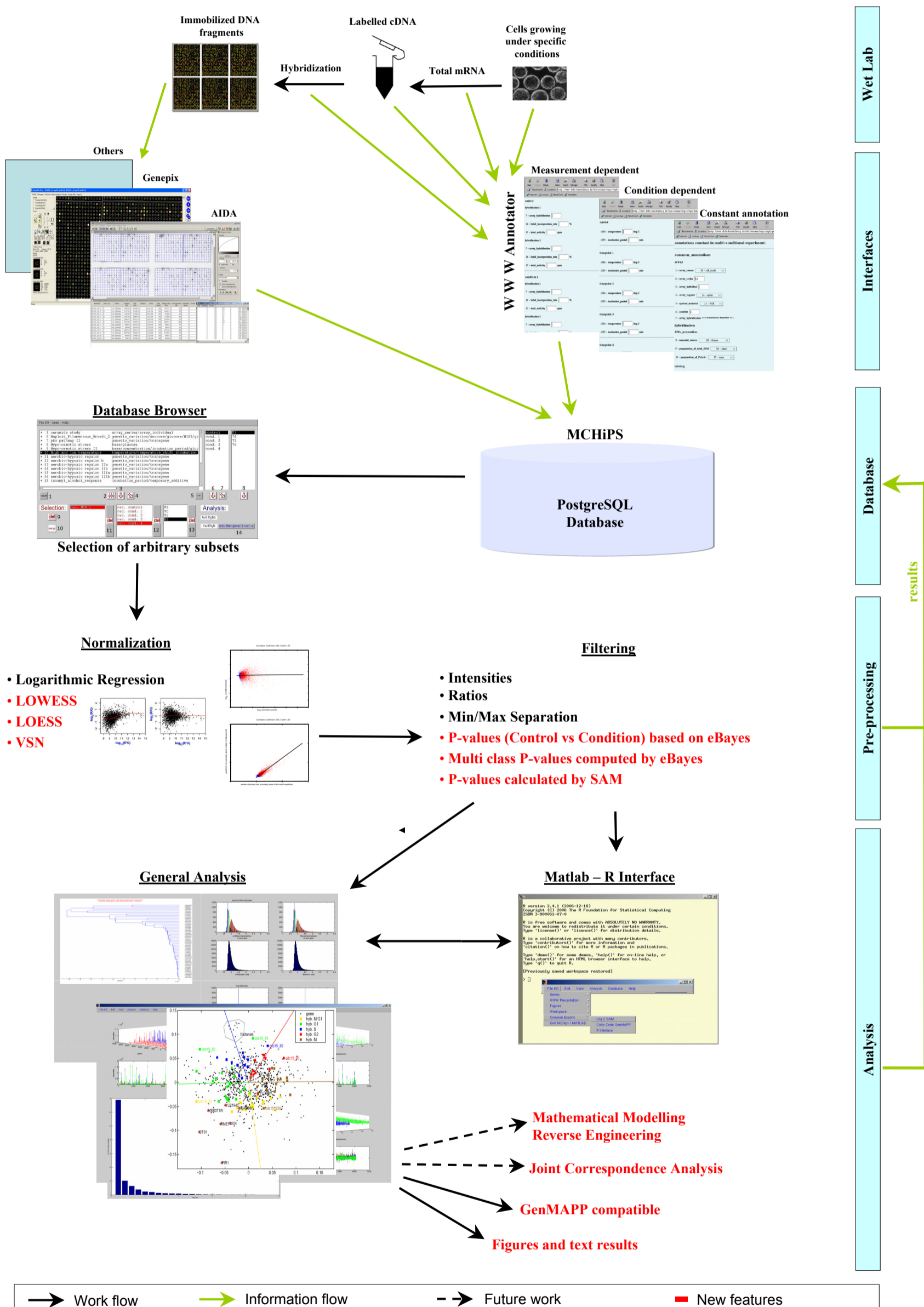
- Menu/GUI re-organization for intuitive use
- Figures and text results
- Export function to GenMAPP
- Integration of AIDA-Analysis Output
- normalization methods:
 - The non-linear LOWESS (locally weighted scatterplot smoothing) and LOESS as well as VSN (Variance Stabilization Normalization)
 - P-values can be computed by SAM (Significance Analysis of Microarray data) using Empirical Bayes methods (eBayes from the limma package)

New functionalities concerning bioinformaticians and statisticians:

- Matlab-R Interface
- Detailed meta data recording everything that has been done to the data during M-CHIPS workflow

Conclusion

Being applied by an increasing number of users since 1999, M-CHIPS has been grown a communication platform, bridging between biologists and statisticians / bioinformaticians. While providing well-established and seamless data analysis by mouse click to the former it allows to apply and/or write any R package or Matlab function on top of the existing functionality to the latter.



References

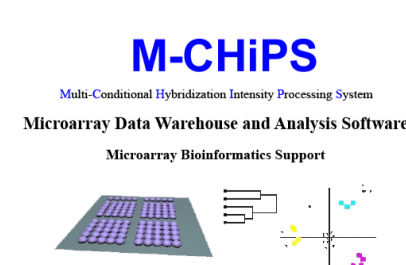
1. Fellenberg K, Busold CH, Witt O, Bauer A, Beckmann B, Hauser NC, Frohme M, Winter S, Dippon J, Hoheisel JD.: Systematic interpretation of microarray data using experiment annotations. *BMC Genomics* (2006) 7:319
2. Fellenberg K, Hauser, NC, Brors B, Hoheisel JD, Vingron M.: Microarray data warehouse allowing for inclusion of experiment annotations in statistical analysis. *Bioinformatics* (2002) 18: 423-433
3. Fellenberg K, Hauser, NC, Brors B, Neutzner A, Hoheisel JD, Vingron M.: Correspondence analysis applied to microarray data. *Proc Natl Acad Sci USA* (2001) 98: 10781-10786

Acknowledgements

Part of this work was funded by DFG - Priority Programme (Schwerpunktprogramm) 1160 ("Colonisation and infection by human-pathogenic fungi"). Special thanks to S. Winter and J. Dippon from the Institute of stochastics and applications, Universität Stuttgart for the mathematical inputs and discussions.

Contact:

Amin Moghaddas Gholami
Deutsches Krebsforschungszentrum (DKFZ)
Dept. Functional Genome Analysis
Im Neuenheimer Feld 580, 69120 Heidelberg, Germany
Phone: +49 (0)6221 422718 Email: a.moghaddasi@dkfz.de



Dipl. Inf. (Bioinf.) Corinna Schmitt
Fraunhofer Institut für Grenzflächen- und Bioverfahrenstechnik (IGB)
Dept. Molecular Biotechnology
Nobelstraße 12, 70569 Stuttgart, Germany
Phone: +49 (0)711 970 4044 Email: corinna.schmitt@igb.fraunhofer.de