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Integrated pathway-level analysis of transcriptomics and metabolomics data with IMPaLA

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ABSTRACT

Summary: Pathway-level analysis is a powerful approach enabling interpretation of post-genomic data at a higher level than that of individual biomolecules. Yet, it is currently hard to integrate more than one type of omics data in such an approach. Here, we present a web tool 'IMPaLA' for the joint pathway analysis of transcriptomics or proteomics and metabolomics data. It performs over-representation or enrichment analysis with user-specified lists of metabolites and genes using over 3000 pre-annotated pathways from 11 databases. As a result, pathways can be identified that may be dysregulated on the transcriptional level, the metabolic level or both. Evidence of pathway dysregulation is combined, allowing for the identification of additional pathways with changed activity that would not be highlighted when analysis is applied to any of the functional levels alone. The tool has been implemented both as an interactive website and as a web service to allow a programming interface.

Availability: The web interface of IMPaLA is available at <http://impala.molgen.mpg.de>. A web services programming interface is provided at <http://impala.molgen.mpg.de/wsdoc>.

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Supplementary Information: Supplementary data are available at *Bioinformatics* online.

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1 INTRODUCTION

Systems biology aims at the concerted analysis of biological systems at different levels, for example the combination of transcriptomics, proteomics and metabolomics. Biochemical pathways are the primary focus of systems biology. Pathways are extensively used to interpret omics data, for example to gain mechanistic insight into gene dysregulation, which is causative or indicative of complex diseases. In particular, pathway over-representation (ORA) and enrichment analyses have become important tools for the interpretation of data from transcriptomics (Riedel *et al.*, 2008) and metabolomics (Sabatine *et al.*, 2005) experiments.

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Several web-based tools exist for such pathway analyses on transcriptomic or metabolomic data separately (Chagoyen and Pazos, 2011; Huang *et al.*, 2008; Kamburov *et al.*, 2011; Xia and Wishart, 2010), but to our knowledge no tools exist yet for integrated pathway analysis with both types of data simultaneously. In a recent study (Cavill *et al.*, 2011), a method for the integrated analysis of transcriptomic and metabolomic data was proposed that exploits the fact that genes and metabolites are linked through biochemical reactions and thus are contained in many pathways. Here, we report the implementation of this method, providing a web server called Integrated Molecular Pathway-Level Analysis (IMPaLA) for the combined analysis of gene/protein and metabolite datasets using a comprehensive basis of biochemical pathways currently taken from 11 publicly available resources. We illustrate the use of the website and exemplify the benefit of combined analysis of transcriptomics and metabolomics data. IMPaLA is accessible through an interactive website or through web services.

2 DESCRIPTION

Web interface: the web interface at <http://impala.molgen.mpg.de> gives the user the possibility to upload genes and/or metabolites in several identifier namespaces and to perform over-representation or Wilcoxon enrichment analysis (WEA) (Adjaye *et al.*, 2005) with the available pathways (Fig. 1A). These pathways (currently 3073) originate from 11 public databases such as Reactome (<http://www.reactome.org>), KEGG (<http://www.genome.jp/kegg/>) or Wikipathways (<http://www.wikipathways.org>). For ORA, the user uploads lists of identifiers that typically represent genes/proteins/metabolites significantly associated with the effect of interest. In addition, background lists of identifiers representing all measured genes/proteins and/or metabolites can be uploaded. This is especially useful when the list of measured entities is small compared with the number of genes/proteins or metabolites in the organism, to avoid potential bias. If no background lists are specified by the user, all entities present in pathways and annotated in the user-specified identifier namespaces are used as default background lists. Based on the uploaded lists, the hypergeometric distribution is used to assess the significance of each pathway in terms of its overlap with those lists.

For WEA, the user uploads lists of all measured genes/proteins and/or metabolites (rather than just the significant ones as in ORA) with either one or a pair of numerical values per entity. These values

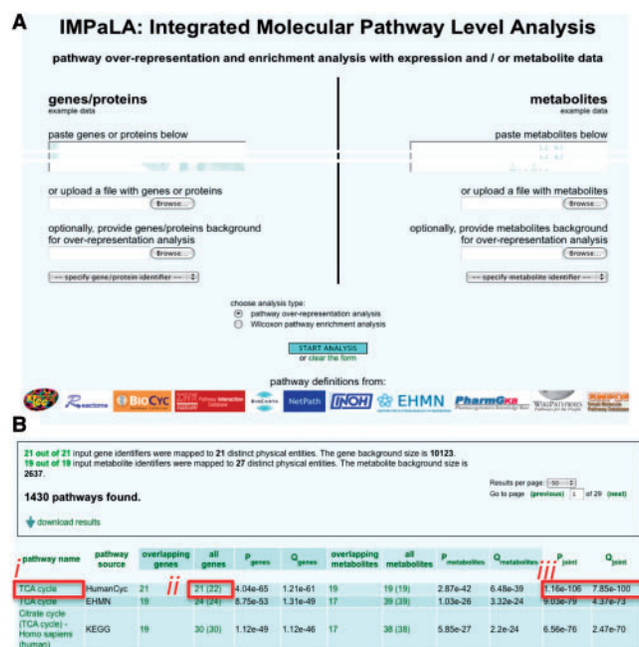


Fig. 1. (A) IMPaLA (version 1) input screen including the logos of the 11 source databases. (B) Output screen with a ranked list of pathways showing; (i) a link to each pathway in the source database; (ii) the size of each pathway in terms of entities also present in the background list, followed by the number of all pathway entities as in the source database; (iii) the P - and Q -values from the joint analysis with genes and metabolites, calculated as in Cavill *et al.* (2011).

may represent fold changes or average expression/concentration values for two different experimental conditions. The values are used to assess the joint expression/concentration difference of all entities contained in each pathway through Wilcoxon's signed-rank test. Even if a pathway contains no individual entities with significant differential expression or concentration, the joint expression/concentration of the group of pathway members may be significantly changed, indicating potential pathway dysregulation on a low but nonetheless consistent level. Results from both ORA and WEA analyses are presented by IMPaLA as tables listing pathways that contain at least one gene and/or metabolite from the input lists (Fig. 1B). Information about pathway name, source, size and overlap with the input entities is provided along with P -values calculated with appropriate statistical test for each pathway. Notably, if both metabolites and genes/proteins are uploaded by the user, a joint P -value is given, calculated as per Cavill *et al.* (2011). To control for multiple testing, Q -values are calculated with the false discovery rate method (Benjamini and Hochberg, 1995). Results can be sorted on any column by clicking on the appropriate column header, and can be downloaded as a tab-delimited file. By clicking on a pathway name, the user is guided to a summary web page at the original source database, which in most cases also shows a detailed pathway diagram.

Example: using publicly available data from the NCI60 (Scherf *et al.*, 2000), we selected the genes and metabolites that were significantly correlated with the GI50 values for the common cancer therapeutic 5-fluorouracil (5-FU) across the 58 cell lines as in previous work (Cavill *et al.*, 2011) (see Files S1–S4 in Supplementary Material that includes background). Full results of the IMPaLA ORA analysis on this data can be found in File 5 in Supplementary Material. The top seven pathways are based entirely on the over-representation of genes, and mainly relate to the ribosome or to eukaryotic translation. For the ABC Transporter pathway, neither genes nor metabolites alone gave $P < 0.05$, yet $P_{\text{joint}} = 0.049$. This example demonstrates that metabolic information gives added value to the genome-wide analysis enhanced the pathway recovery. For further examples and help, please see the tutorial document provided in File S7 in Supplementary Material.

Web service: in addition to the standard interactive web interface, the functionality provided can also be accessed through a SOAP web service. Here, functions are available that carry out ORA or WEA with lists of genes, of metabolites or both. The web service definition (WSDL) file and the appropriate documentation are available at <http://impala.molgen.mpg.de/wsdoc>.

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REFERENCES

- Adjaye, J. *et al.* (2005) Primary differentiation in the human blastocyst: Comparative molecular portraits of inner cell mass and trophectoderm cells. *Stem Cells*, **23**, 1514–1525.
- Benjamini, Y. and Hochberg, Y. (1995) Controlling the false discovery rate: a practical and powerful approach to multiple testing. *J. R. Stat. Soc. Ser. B*, **57**, 289–300.
- Cavill, R. *et al.* (2011) Consensus-phenotype integration of transcriptomic and metabolomic data implies a role for metabolism in the chemosensitivity of tumour cells. *PLoS Comput. Biol.*, **7**, e1001113.
- Chagoyen, M. and Pazos, F. (2011) MBRole: enrichment analysis of metabolomic data. *Bioinformatics*, **27**, 730–731.
- Huang, D.W. *et al.* (2008) Systematic and integrative analysis of large gene lists using DAVID bioinformatics resources. *Nat. Protocols*, **4**, 44–57.
- Kamburov, A. *et al.* (2011) ConsensusPathDB: toward a more complete picture of cell biology. *Nucleic Acids Res.*, **39**, D712–D717.
- Riedel, R.F. *et al.* (2008) A genomic approach to identify molecular pathways associated with chemotherapy resistance. *Mol. Cancer Therap.*, **7**, 3141–3149.
- Sabatine, M.S. *et al.* (2005) Metabolomic identification of novel biomarkers of myocardial ischemia. *Circulation*, **112**, 3868–3875.
- Scherf, U. *et al.* (2000) A gene expression database for the molecular pharmacology of cancer. *Nat. Genet.*, **24**, 236–244.
- Xia, J. and Wishart, D.S. (2010) MSEA: a web-based tool to identify biologically meaningful patterns in quantitative metabolomic data. *Nucleic Acids Res.*, **38**, W71–W77.