Gene expression

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# qCMAP: user-friendly connectivity mapping with R

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#### **ABSTRACT**

Connections between disease phenotypes and drug effects can be made by identifying commonalities in the associated patterns of differential gene expression. Searchable databases that record the impacts of chemical or genetic perturbations on the transcriptome-here referred to as 'connectivity maps'-permit discovery of such commonalities. We describe two R packages, gCMAP and gCMAPWeb, which provide a complete framework to construct and query connectivity maps assembled from user-defined collections of differential gene expression data. Microarray or RNAseq data are processed in a standardized way, and results can be interrogated using various wellestablished gene set enrichment methods. The packages also feature an easy-to-deploy web application that facilitates reproducible research through automatic generation of graphical and tabular reports. Availability and implementation: The gCMAP and gCMAPWeb R packages are freely available for UNIX, Windows and Mac OS X operating systems at Bioconductor (http://www.bioconductor.org).

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## 1 INTRODUCTION

Microarray and RNAseq technologies enable the study of transcriptional changes triggered by experimental perturbations. Gene expression changes observed after one perturbation often resemble those produced by others, suggesting a shared biological mechanism (Lamb et al., 2006). This strategy for establishing links between perturbations can then be used, for example, to identify bioactive compounds (Kunkel et al., 2011; Stumpel et al., 2012), elucidate their mode of action (Coombs et al., 2012) or reposition approved pharmaceuticals for use in new indications (Dudley et al., 2011). Standalone tools (Subramanian et al., 2005), modular software libraries (Pacini et al., 2013) or hosted web applications are available to query the original Connectivity Map (Lamb et al., 2006), but tools that allow users to easily compile and search their own reference datasets are lacking.

The gCMAP package provides utilities and memory-efficient structures to create, store and query large experimental datasets. gCMAP also provides a unified R command-line interface to multiple gene set enrichment (GSE) approaches, permitting their use for querying user-created connectivity maps as well as more traditional gene set collections—e.g. Gene Ontology, Reactome or WikiPathways (Jupe et al., 2012; Kelder et al.,

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2012). In addition, the companion gCMAPWeb package provides a customizable browser-based graphical user interface. and can leverage R's built-in web server or be integrated into a production-scale server.

#### 2 FEATURES

To compile a new connectivity map, users supply expression data in the form of standard Bioconductor ExpressionSet objects. gCMAP provides convenience functions to split large studies into individual perturbation instances and process them in batch, taking advantage of the widely used limma (Smyth, 2004) and DESeq (Anders and Huber, 2010) Bioconductor packages. OC metrics permit removal of unsuitable experiments, e.g. those suffering from normalization artifacts (Supplementary Material). To allow memory-efficient access to large datasets, results are stored on disk and data subsets are loaded on demand.

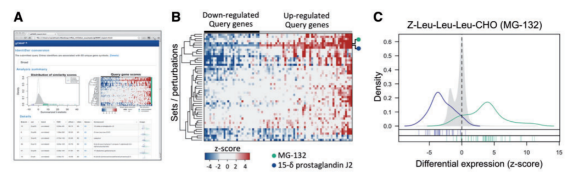
gCMAP represents gene set membership and (optionally) direction of expression change via sparse numerical matrices, enabling efficient GSE analysis using matrix operations. Analysis of complete differential expression profiles, which retains more information than gene sets, is also supported, permitting permutation-based assessment of statistical significance.

gCMAP implements several well-established GSE methods, including Fisher's exact test, the original GSEA statistic (Lamb et al., 2006), the JG score (Jiang and Gentleman, 2007), mgsa (Bauer et al., 2011) and Roast (Wu et al., 2010). All gCMAP implementations use common input and output formats, enabling direct comparisons among the supported statistics.

The gCMAPWeb package complements gCMAP by providing a graphical user interface through a distributable web application. gCMAPWeb leverages R's built-in web server, permitting singleuser access on any computer running R, and can also be integrated into an Apache web server in a multiuser environment. To submit queries, users can paste gene identifiers directly into a web form or upload text files (Supplementary File S1). gCMAPWeb returns results in graphical and tabular form, with detail at the gene set and single-gene levels (Fig. 1). Every report, including graphs, tables and a binary R object, is available for download. The gCMAPWeb user interface is automatically configured to match reference datasets but can be easily customized.

#### 3 EXAMPLES

We used gCMAP to reconstruct the original Connectivity Map Lamb et al. (2006), containing 453 individual perturbations in five human cell lines, from raw microarray data (ArrayExpress E-GEOD-5258, Supplementary File S1). We then used gCMAPWeb to query this dataset with genes found to be



**Fig. 1.** Output returned by gCMAPWeb for a directional query with genes up- or downregulated in MCF7 cells after treatment with 15-δ prostaglandin J2 for 6 h. **(A)** gCMAPWeb's main result page. **(B)** Heatmap displaying differential expression scores for genes (columns) from the top 50 most similar experiments in the Broad Connectivity Map (rows). Treatment with MG-132 (green dot) received the highest JG summary score and was clustered next to the experiment corresponding to the query itself (blue dot). **(C)** Distribution of Z-scores indicating differential expression following MG-132 treatment for all assayed genes (gray) and up- (green) or downregulated (blue) query genes. Query gene scores are shifted toward positive and negative scores

up- or downregulated in MCF7 cells treated with the NF $\kappa$ B inhibitor 15- $\delta$  prostaglandin J2. Top hits included MG-132/Z-Leu-Leu-CHO and celastrol, two known inhibitors of the same pathway (Fig. 1).

To assess differential expression from count data, *gCMAP* leverages the widely used *DESeq* package (Anders and Huber, 2010). To re-examine the transcriptional response of HepG2 cells to the carcinogen Benzo[a]pyrene [van Delft *et al.* (2012), ENA accession SRP011233, Supplementary File S2], we constructed a local gene set collection from WikiPathways (Kelder *et al.*, 2012) and queried it with genes significantly up- or downregulated in response to Benzo[a]pyrene. In concordance with the original study, *gCMAPWeb* reported significant overlap between the query and the Benzo[a]pyrene metabolism and Nrf2/Keaf1 pathways [van Delft *et al.* (2012), Supplementary File S1].

### 4 CONCLUSION

The *gCMAP* Bioconductor packages combine powerful command-line functionality with a convenient portable web application. The efficient handling of large datasets empowers users to assemble large connectivity maps from private or public data, query them programmatically or through an interactive user interface and store queries and results in a reproducible report.

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Conflict of Interest: none declared.

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