# KEGGandMetacoreDzPathwaysGEO : Disease Datasets from GEO

### Gaurav Bhatti

 $17 \ {\rm April} \ 2014$ 

## 1 Overview of KEGGandMetacoreDzPathwaysGEO data package

*KEGGandMetacoreDzPathwaysGEO* is a collection of 18 GEO datasets for which the phenotype is a disease with a corresponding pathway in either of the two popular gene to pathway annotation databases, KEGG and Metacore. These datasets were used as gold standard in comparing gene set analysis methods [1].Details about the individual datasets including sample tissue, target disease pathway, etc may be obtained by typing:

#### > ?KEGGandMetacoreDzPathwaysGEO

at the R prompt. In order to access all the datasets available in the package, type the following:

# > mysets=data(package="KEGGandMetacoreDzPathwaysGEO")\$results[,"Item"] > mysets

The microarray data from the GEO database along with the associated metadata is stored as ExpressionSet class. "The ExpressionSet class is designed to combine several different sources of information into a single convenient structure. An ExpressionSet can be manipulated (e.g., subsetted, copied) conveniently, and is the input or output from many Bioconductor functions." [2].An example dataset is shown below:

```
> library(KEGGandMetacoreDzPathwaysGEO)
> data(GSE1145)
> show(GSE1145)
ExpressionSet (storageMode: lockedEnvironment)
assayData: 54675 features, 26 samples
    element names: exprs
protocolData: none
```

```
phenoData
  sampleNames: GSM18442 GSM18443 ... GSM18436 (26 total)
  varLabels: Sample Group
  varMetadata: labelDescription
featureData: none
experimentData: use 'experimentData(object)'
Annotation: hgu133plus2
```

A similar data package, KEGGDzPathwaysGEO, is already available for installation in Bioconductor.It contains additional 24 GEO datasets for which the phenotype is a disease with a corresponding pathway in the KEGG database.These datasets were used to test the performance of an in-house pathway analysis method which has also been implemented as a Bioconductor package, PADOG [3].

These datasets may be used to compare existing gene set pathway analysis methods or to test the performance of novel methods.

## References

- Tarca, A. L. et al. (2013) A Comparison of Gene Set Analysis Methods in Terms of Sensitivity, Prioritization and Specificity. PLoS ONE 8(11): e79217.doi:10.1371/journal.pone.0079217
- [2] Falcon, S., Morgan, M., and Gentleman, R. (2007), An Introduction to Bioconductor's ExpressionSet Class.
- [3] Tarca, A. L. et al. (2012). Down-weighting overlapping genes improves gene set analysis. BMC Bioinformatics, 13, 136-2105-13-136.