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1 Introduction

Discrimination between different classes of samples such as different cell types or tissues using gene expression profiles is an important problem in genetic and cell research. It has several implications and can contribute to our understanding of cell phenotype differences and will allow precise identification of various cell types and tissues. <code>sampleClassifier</code> provides functions for the classification of samples using their gene expression profiles. The package supports the classification of microarray and RNA-seq gene expression profiles. The tool requires a reference and a test data set, and uses a simple algorithm called Shared Marker Genes (SMG). As the name suggests, the number of shared marker genes between a reference and a query sample is used as a similarity measure. Marker genes are detected using the tools <code>MGFM</code> [1, 2] or <code>MGFR</code> [3] that we have developed previously. <code>sampleClassifier</code> can be applied: i) to evaluate the similarity of experimentally derived cells with their desired target cell type (e.g. to compare primary hepatocytes with induced hepatocytes); ii) to compare in vitro derived organoids to their in vivo counterparts; iii) to classify different types of diseases.

This vignette provides an introduction to gene expression profile based sample classification using the *sampleClassifier* R package and the accompanying data package, *sampleClassifier-Data*. It contains guidelines on how to select all inputs to the tool (such as reference and test matrices), and instructions on running *sampleClassifier* using microarray and RNA-seq data from the *sampleClassifierData*.

2 Contents of the package

The *sampleClassifier* package contains the following functions:

2.1 classifyProfile()

classifyProfile() is a function to classify microarray gene expression profiles.

2.1.1 Parameter Settings

- 1. ref_matrix : Normalized microarray data matrix to be used as reference, with probe sets corresponding to rows and samples corresponding to columns.
- 2. *query_mat*: Normalized microarray query matrix with query samples to be classified, with probe sets corresponding to rows and samples corresponding to columns.
- 3. chip1: Chip name of the reference matrix (e.g. 'hgu133plus2').
- 4. *chip2*: Chip name of the query matrix. This parameter can be ignored if the reference and query matrix are from the same chip.

- 5. *fun1*: mean or median. This will specify the number of marker genes that will be used for classification. Default is median.
- 6. fun2: mean or median. This will be used to summarize the expression values of probe sets that belong to the same gene. This parameter can be ignored if the reference and query matrix are from the same chip. Default is mean.
- 7. write2File: If TRUE, the classification results for each query profile will be written to a file.
- 8. *out.dir*: Path to a directory to write the classification results, default is the current working directory.

2.1.2 Output

The function classifyProfile() returns a list with hits for each of the query samples in the query matrix. The hits are sorted according to their similarity to the query.

2.2 classifyProfile.rnaseq()

classifyProfile.rnaseq() is a function to classify RNA-seq gene expression profiles.

2.2.1 Parameter Settings

- 1. ref_matrix : RNA-seq data matrix to be used as reference, with genes corresponding to rows and samples corresponding to columns.
- 2. *query_mat*: RNA-seq query matrix with query samples to be classified, with genes corresponding to rows and samples corresponding to columns.
- 3. *gene.ids.type*: Type of the used gene identifiers, the following gene identifiers are supported: ensembl, refseq and ucsc gene ids. Default is ensembl.
- 4. fun1: mean or median. This will specify the number of marker genes that will be used for classification. Default is median.
- 5. write2File: If TRUE, the classification results for each query profile will be written to a file.
- 6. *out.dir*: Path to a directory to write the classification results, default is the current working directory.

2.2.2 Output

The function classifyProfile.rnaseq() returns a list with top hits for each query profile, sorted according to a similarity score.

2.3 classifyProfile.svm()

classifyProfile.svm() is a function to classify microarray gene expression profiles using Support Vector Machines (SVM).

2.3.1 Parameter Settings

- 1. ref_matrix : Normalized microarray data matrix to be used as reference, with probe sets corresponding to rows and samples corresponding to columns.
- 2. *query_mat*: Normalized microarray query matrix to be classified, with probe sets corresponding to rows and samples corresponding to columns.
- 3. chip1 : Chip name of the reference matrix (e.g. 'hgu133plus2').
- 4. *chip2*: Chip name of the query matrix. This parameter can be ignored if the reference and query matrix are from the same chip.
- 5. fun1: mean or median. This will specify the number of marker genes that will be used for classification. Default is median.
- 6. fun2: mean or median. This will be used to summarize the expression values of probe sets that belong to the same gene. This parameter can be ignored if the reference and query matrix are from the same chip.

2.3.2 Output

The function classifyProfile.svm() returns a data frame with the predicted classes for each query profile.

2.4 classifyProfile.rnaseq.svm()

classifyProfile.rnaseq.svm() is a function to classify RNA-seq gene expression profiles using Support Vector Machines (SVM).

2.4.1 Parameter Settings

- 1. ref_matrix : RNA-seq data matrix to be used as reference, with genes corresponding to rows and samples corresponding to columns.
- 2. *query_mat*: RNA-seq query matrix with query samples to be classified, with genes corresponding to rows and samples corresponding to columns.
- 3. *gene.ids.type*: Type of the used gene identifiers, the following gene identifiers are supported: ensembl, refseq and ucsc gene ids. Default is ensembl.
- 4. fun1: mean or median. This will specify the number of marker genes that will be used for classification. Default is median.

2.4.2 Output

The function classifyProfile.rnaseq.svm() returns a data frame with the predicted classes for each query profile.

Please note that all replicates of a sample type should have the same label in the reference matrix.

2.5 get.heatmap()

get.heatmap() is a function to display the classification predictions as a heatmap.

2.5.1 Parameter Settings

res.list: the result list returned by the function classifyProfile() or classifyProfile.rnaseq()

2.5.2 Output

The function get.heatmap() is used only for the side effect of creating a heatmap.

3 Getting Started

The *sampleClassifier* package can be downloaded and installed by running the following code from within R:

```
> source("http://bioconductor.org/biocLite.R")
> biocLite("sampleClassifier")
```

We also recommend installing the accompanying data package, *sampleClassifierData*, which contains pre-processed microarray and RNA-seq data, which are derived from normal human tissues, and are available from public repositories. More details about the data can be found in the Vignette of the data package.

After installing and loading sampleClassifierData, the individual sampleClassifierData datasets can be loaded using the function data(). For instance, the RNA-seq dataset named se_rnaseq_refmat is stored as a SummarizedExperiment object. The numeric matrix can be extracted using the function assay() from the SummarizedExperiment R package:

```
> library(sampleClassifierData)
> data("se_rnaseq_refmat")
> rnaseq_refmat <- assay(se_rnaseq_refmat)</pre>
```

4 Classification using sampleClassifier

We will use the data provided in the data package *sampleClassifierData* to demonstrate how to classify samples using *sampleClassifier*.

4.1 Classification of microarray data

To classify microarray gene expression profiles, we use the function classifyProfile(). It expects a reference and a test matrix. We recommend using 3 replicates for each sample type in the reference matrix. Please note that replicates of the same sample type should have the same name in the reference matrix. We also recommend processing the reference and the test matrix in the same way.

As a reference matrix we use here a microarray dataset derived from the study GSE3526 [4], which is available from GEO [5]. This dataset is named se_micro_refmat and can be loaded with the following code:

```
> library(sampleClassifierData)
> data("se_micro_refmat")
> micro_refmat <- assay(se_micro_refmat)
> dim(micro_refmat)
[1] 54675 78
```

As a test matrix we use a microarray dataset derived from the study GSE2361 [6], which is available from GEO [5]. This dataset is named <u>se_micro_testmat</u> and can be loaded with the following code:

Now, we can call the function classifyProfile():

```
> res1.list <- classifyProfile(ref_matrix=micro_refmat, query_mat=micro_testmat,
+ chip1="hgu133plus2",chip2="hgu133a", write2File=FALSE)

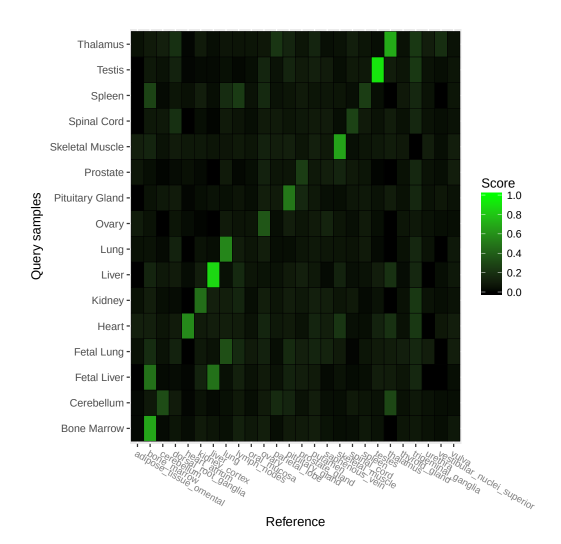
The reference matrix and the query are from different platforms...
Collapse rows ...
detecting marker genes...
16 profiles to be classified...
done!</pre>
```

For simplicity, we show only the two top hits for each query sample:

```
1 ovary 0.364 20 / 55
2 saphenous_vein 0.137 10 / 73
$`GSM44675 : Kidney`
               Hits Score Ratio
      kidney_cortex 0.462 36 / 78
2 trigeminal_ganglia 0.231 3 / 13
$`GSM44676 : Skeletal Muscle`
            Hits Score Ratio
1 skeletal_muscle 0.667 52 / 78
bone_marrow 0.141 11 / 78
$`GSM44678 : Prostate`
             Hits Score Ratio
     prostate_gland 0.256 20 / 78
2 trigeminal_ganglia 0.154 2 / 13
$`GSM44689 : Cerebellum`
       Hits Score Ratio
1 cerebellum 0.321 25 / 78
2 thalamus 0.3 3 / 10
$`GSM44693 : Bone Marrow`
        Hits Score Ratio
1 bone_marrow 0.679 53 / 78
2 lymph_nodes 0.122 6 / 49
$`GSM44698 : Thalamus`
               Hits Score Ratio
          thalamus 0.7 7 / 10
2 trigeminal_ganglia 0.231 3 / 13
$`GSM44699 : Pituitary Gland`
               Hits Score Ratio
    pituitary_gland 0.5 39 / 78
2 trigeminal_ganglia 0.154 2 / 13
$`GSM44700 : Spinal Cord`
                Hits Score Ratio
         spinal_cord 0.278 20 / 72
2 dorsal_root_ganglia 0.194 6 / 31
$`GSM44701 : Testis`
              Hits Score Ratio
            testes 0.91 71 / 78
2 trigeminal_ganglia 0.231 3 / 13
$`GSM44702 : Liver`
    Hits Score Ratio
1 liver 0.859 67 / 78
```

To display the classification results as a heatmap, we call the function get.heatmap() with the resulted list as input.

```
> get.heatmap(res1.list)
```



In order to compare the classification predictions for microarray query profiles to those of Support Vector Machines (SVM), the function classifyProfile.svm() was implemented based on functions from the R-package e1071.

```
> res1.svm.df <- classifyProfile.svm(ref_matrix=micro_refmat, query_mat=micro_testmat,
+ chip1="hgu133plus2",chip2="hgu133a")
The reference matrix and the query are from different platforms...
Collapse rows ...
detecting marker genes...
building an SVM model...
16 profiles to be classified...
done!
> res1.svm.df
                                      predicted_class
                   query_name
             GSM44671 : Heart
1
                                         heart_atrium
2
            GSM44673 : Spleen adipose_tissue_omental
3
             GSM44674 : Ovary
                                                ovary
4
            GSM44675 : Kidney
                                        kidney\_cortex
```

```
GSM44676 : Skeletal Muscle
                                      skeletal_muscle
6
          GSM44678 : Prostate
                                              urethra
7
        GSM44689 : Cerebellum
                                           cerebellum
8
       GSM44693 : Bone Marrow
                                          bone_marrow
9
          GSM44698 : Thalamus
                                             thalamus
10 GSM44699 : Pituitary Gland
                                      pituitary_gland
11
       GSM44700 : Spinal Cord
                                          spinal_cord
12
            GSM44701 : Testis
                                               testes
13
             GSM44702 : Liver
                                                 liver
14
              GSM44704 : Lung
                                                  luna
15
        GSM44705 : Fetal Lung
                                                  lung
       GSM44706 : Fetal Liver
                                          bone_marrow
```

Our method classifyProfile() classified 14 samples correctly, whereas SVM classified 13 of the 16 query samples correctly. The sample 'GSM44678' was classified correctly by classifyProfile() as prostate and miclassified by SVM as urethra. The sample 'GSM44673' (tissue: spleen) was misclassified as bone marrow or adipose tissue omental by classifyProfile() or classifyProfile.svm(), respectively.

The sample 'GSM44706' (tissue: fetal liver) was misclassified as bone marrow by SVM. classifyProfile() predicted both bone marrow and liver as top hits with the same similarity score.

4.2 Classification of RNA-seq data

To classify RNA-seq gene expression profiles, we use the function classifyProfile.rnaseq(). It expects a reference and a test matrix. As a reference matrix we use an RNA-seq dataset derived from the study E-MTAB-1733 [7], which is available from ArrayExpress [8]. This dataset is named se_rnaseq_refmat and can be loaded with the following code:

```
> library(sampleClassifierData)
> data("se_rnaseq_refmat")
> rnaseq_refmat <- assay(se_rnaseq_refmat)
> dim(rnaseq_refmat)
[1] 43819 71
```

As a test matrix we use an RNA-seq dataset derived from the study E-MTAB-513 [9], which is available from ArrayExpress. This dataset is named $se_rnaseq_testmat$ and can be loaded with the following code:

```
> data("se_rnaseq_testmat")
> rnaseq_testmat <- assay(se_rnaseq_testmat)
> dim(rnaseq_testmat)
[1] 43819 12
```

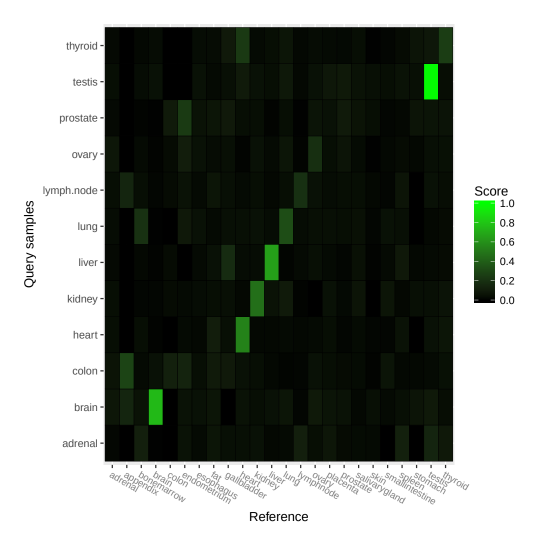
Now, we can call the function classifyProfile.rnaseq() to predict the classes of the samples in the test matrix:

```
> res2.list <- classifyProfile.rnaseq(ref_matrix=rnaseq_refmat, query_mat=rnaseq_testmat,
+ gene.ids.type="ensembl",write2File=FALSE)</pre>
```

```
Detecting marker genes...
Done!
12 profiles to be classified...
Done!
For simplicity, we show only the two top hits for each query sample:
> lapply(res2.list,"[",c(1,2),,drop=FALSE)
$adrenal
   Hits Score
                 Ratio
1 testis 0.141 26 / 184
2 spleen 0.121 12 / 99
$brain
     Hits Score Ratio
1 brain 0.75 138 / 184
2 appendix 0.143 1 / 7
$colon
        Hits Score Ratio
1 appendix 0.286 2 / 7
2 endometrium 0.135 5 / 37
$heart
  Hits Score Ratio
1 heart 0.543 100 / 184
2 fat 0.103 19 / 184
```

To display the classification results as a heatmap, we call the function get.heatmap() with the resulted list as input.

```
> get.heatmap(res2.list)
```



In order to compare the classification predictions for RNA-seq query profiles to those of SVM, the function classifyProfile.rnaseq.svm() was implemented based on functions from the R-package e1071.

```
> res2.svm.df <- classifyProfile.rnaseq.svm(ref_matrix=rnaseq_refmat, query_mat=rnaseq_testmat,
+ gene.ids.type="ensembl")
Detecting marker genes...
Done!
building an SVM model...
12 profiles to be classified...
done!
> res2.svm.df
   query_name predicted_class
1
      adrenal
                      appendix
2
        brain
                         brain
                  gallbladder
3
        colon
4
        heart
                         heart
5
       kidney
                        kidney
```

```
6
        liver
                          liver
7
          lung
                           lung
   lymph.node
                       appendix
8
9
        ovary
                    endometrium
                    endometrium
10
     prostate
11
       testis
                         testis
12
                        thyroid
      thyroid
```

Our method classifyProfile.rnaseq() classified 9 samples correctly, whereas SVM classified 7 of the 12 query samples correctly. To show the query samples that were misclassified by our method or SVM, we run the following code:

```
> misclas.inds <- which(as.character(res2.svm.df[,1])!=as.character(res2.svm.df[,2]))</pre>
> colnames(res2.svm.df) <- c("query_real_class","predicted_class_by_SVM")</pre>
> pred.classifyProfile.rnaseg <- as.character(unlist(lapply(res2.list[which(names(res2.list) %in%)</pre>
+ as.character(res2.svm.df[misclas.inds,1]))],"[",1,1,drop=TRUE)))
> comp.df <- cbind(res2.svm.df[misclas.inds,],</pre>
+ predicted_by_classifyProfile.rnaseq=pred.classifyProfile.rnaseq)
   query_real_class predicted_class_by_SVM predicted_by_classifyProfile.rnaseq
                                    appendix
1
            adrenal
                                                                            testis
3
               colon
                                 gallbladder
                                                                          appendix
8
         lymph.node
                                    appendix
                                                                         lymphnode
                                 endometrium
               ovary
                                                                             ovary
10
                                 endometrium
                                                                       endometrium
           prostate
```

5 sampleClassifier algorithm details

The algorithm used in sampleClassifier is a simple algorithm called Shared Marker Genes (SMG). As the name suggests, the number of shared marker genes between a query and a reference is used as similarity measure. The tool requires a reference matrix with at least three replicates for each sample type. This matrix is used for marker gene detection using MGFM or MGFR. Since the number of detected markers differs depending on the sample types, we filter the list of marker genes of each sample type. Using the complete list of markers of each sample type, will result in a bias towards the sample type with the most marker genes. For example, if testis is the tissue with the most marker genes, using all marker genes for classification will result in classifying query samples often as testis. Suppose the reference matrix contains four tissues: liver, lung, kidney and midbrain, and X=(16, 20, 100)100, 500) is the vector of lengths of predicted marker genes for these tissues. The filtering is based on the median number of marker genes, in this case median(X) = 60. If the number of predicted markers for a tissue > 60, then only the top 60 marker genes will be used for classification. If the number of predicted markers for a tissue \leq 60, then all markers are used for classification. After the filtering step, each query sample will be compared to all sample types in the reference and the number of marker genes shared between the query and each sample type in the reference is calculated. A query shares a marker gene with a reference sample if this marker gene is highly expressed in the query sample compared to all other sample types in the reference. The ratio of the number of shared marker genes and the total number of markers used for classification is used as a similarity score. This score has a

value between 0 and 1. A value of 1 means that the query shares all marker genes with the reference, and a value of 0 means that no marker genes are shared between the query and the reference. For each query, the hits are sorted according to this score. The class of the first top hit is predicted as a class for the query.

6 R sessionInfo

The results in this file were generated using the following packages:

```
> sessionInfo()
R version 3.4.2 (2017-09-28)
Platform: x86_64-apple-darwin15.6.0 (64-bit)
Running under: macOS High Sierra 10.13.1
Matrix products: default
BLAS: /Library/Frameworks/R.framework/Versions/3.4/Resources/lib/libRblas.0.dylib
LAPACK: /Library/Frameworks/R.framework/Versions/3.4/Resources/lib/libRlapack.dylib
locale:
[1] de_DE.UTF-8/de_DE.UTF-8/de_DE.UTF-8/c/de_DE.UTF-8/de_DE.UTF-8
attached base packages:
[1] parallel stats4
                        stats
                                   graphics grDevices utils
                                                                  datasets
[8] methods
              base
other attached packages:
 [1] hqu133a.db_3.2.3
                                 hgu133plus2.db_3.2.3
 [3] org.Hs.eg.db_3.5.0
                                 sampleClassifierData_1.2.0
 [5] SummarizedExperiment_1.8.0 DelayedArray_0.4.1
 [7] matrixStats_0.52.2
                                 GenomicRanges_1.30.0
 [9] GenomeInfoDb_1.14.0
                                 sampleClassifier_1.2.0
[11] MGFR_1.4.0
                                 MGFM 1.12.0
[13] annotate_1.56.1
                                 XML_3.98-1.9
[15] AnnotationDbi_1.40.0
                                 IRanges_2.12.0
[17] S4Vectors_0.16.0
                                 Biobase_2.38.0
[19] BiocGenerics_0.24.0
loaded via a namespace (and not attached):
 [1] progress_1.1.2
                             lattice_0.20-35
                                                      colorspace_1.3-2
 [4] htmltools_0.3.6
                             {\tt yaml\_2.1.14}
                                                      blob_1.1.0
 [7] rlang_0.1.4
                              e1071_1.6-8
                                                      DBI_0.7
[10] bit64_0.9-7
                              GenomeInfoDbData_0.99.1 plyr_1.8.4
[13] stringr_1.2.0
                              zlibbioc_1.24.0
                                                      munsell_0.4.3
                              memoise_1.1.0
                                                      evaluate_0.10.1
[16] gtable_0.2.0
[19] knitr_1.17
                              biomaRt_2.34.0
                                                      class_7.3-14
[22] Rcpp_0.12.13
                              xtable_1.8-2
                                                      backports_1.1.1
[25] scales_0.5.0
                             XVector_0.18.0
                                                      bit_1.1-12
[28] BiocStyle_2.6.0
                              ggplot2_2.2.1
                                                      digest_0.6.12
[31] stringi_1.1.5
                                                      grid_3.4.2
                              rprojroot_1.2
```

[34] tools_3.4.2	bitops_1.0-6	${\sf magrittr}_{-}$ 1.5
[37] RCurl_1.95-4.8	lazyeval_0.2.1	$tibble_1.3.4$
[40] RSQLite_2.0	pkgconfig $_2.0.1$	Matrix_1.2-11
[43] prettyunits_1.0.2	$assertthat_0.2.0$	$rmarkdown_{-}1.7$
[46] R6_2.2.2	compiler_3.4.2	

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http://www.ncbi.nlm.nih.gov/pubmed/24309898http://www.pubmedcentral.nih.gov/articlerender.fcgi?artid=PMC3916642,doi:10.1074/mcp.M113.035600.

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