Annotation-Driven Class Discovery

User’s Guide to the Bioconductor package *adSplit*

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Abstract

This is the vignette of the Bioconductor compliant package *adSplit*. We describe our implementation of *annotation-driven clustering* for microarray gene expression profiling studies.

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Chapter 1

Introduction

Background: Clustering algorithms are widely used in the analysis of microarrays. In clinical studies, they are not only used to cluster genes into groups of co-regulated genes, but also for clustering patients, and thereby defining novel disease entities based on gene expression profiles. Several distance based cluster algorithms have been suggested, but little attention has been paid to the choice of the metric on patient space. Even when using the Euclidean metric, including and excluding genes from the analysis leads to different distances between the same objects, and consequently to different clustering results.

Methodology: In this package, we implement a novel algorithm for investigating the dependency of clustering results on the choice of metric supporting genes. Our method combines expression data and functional annotation data. According to gene annotations, a list of candidate gene sets with a unique functional characterization is generated. Each set defines a metric on patient space, and consequently a clustering of patients. Based on a novel significance measure for clusterings, this list is filtered. Significant clusterings are reported together with the underlying gene sets and their functional definition.

Intended results: Our method reports clusterings defined by biologically focussed sets of genes. In our annotation driven clusterings, we have observed rediscoveries of clinically relevant patient subgroups through biologically plausible sets of genes. Hence, we conjecture that our method has the potential to reveal clinically relevant classes of patients in an unsupervised manner.

The algorithmic concept: We suggest a systematic approach to gene selection in an unsupervised setting. We describe an algorithm that produces a list of alternative clusterings using a variety of different metrics on patient space. While all metrics are of the Euclidean type, they differ in the set of genes used for characterizing the patient profiles. We derive candidate gene sets from functional annotation data, and filter the list by a novel significance measure for clustering strength. For practical use, it is desirable to have functional rationales characterizing clusterings. For instance, clusterings related to proliferation or apoptosis. To this end, we define candidate gene sets using functional
annotations from the Gene Ontology [1] and from the Kyoto Encyclopedia of Genes and Genomes [7].

The algorithm in a nutshell:

Annotation-driven splits specific to the corresponding gene set are computed using the k-means algorithm [5]. We use functional annotations from chip-specific meta-data packages provided in Bioconductor [3]. The quality of any clusterings is assessed using the diagonal linear discriminant (DLD) score [8]. In order to determine the statistical significance of a score, we also compute DLD scores for restricted metrics resulting from randomly chosen gene sets. Empirical p-values are calculated and false discovery rates (FDR) computed according to Benjamini Hochber [2]. Finally, we filter the list of clusterings for minimal subgroup size and to control the FDR. In a nutshell, the algorithm consists of the following steps:

For each biological term / pathway of interest, denoted $B_i$:

1. Find all $n_{B_i}$ genes annotated to $B_i$ and discard all others.
2. Perform 2-means clustering of reduced expression matrix. This yields an annotation-driven clustering $C_{B_i}$.
3. Compute DLD score $S(C_{B_i})$ for this clustering.
4. Draw 10000 random gene sets of size $n_{B_i}$ from the set of all measured genes. For each of these random gene sets, compute steps 2 and 3. This yields a vector $r_{n_{B_i}}$ of 10000 scores.
5. Assign an empirical p-value to the original clustering, denoting the proportion of entries of $r_{n_{B_i}}$ being greater or equal than $S(C_{B_i})$.
6. Correct the empirical p-values for multiple testing according to Benjamini-Hochberg [2]. The resulting q-values are used to control the false discovery rate of a splitSet.

The structure of the vignette: The described package contains functions to facilitate these steps. The first chapter describes, how to compute a split for a given term of interest. the second shows how to compute a random distribution of DLD-scores to judge the significance of an annotation-driven split. Finally, the last chapter describes how to collect results for many terms of interest and illustrate the achieved results.
Chapter 2

Annotation-Driven Splits

Before starting any analysis you have to load the adSplit package in your R session as follows:

```r
> library(adSplit)
```

For illustration of adSplit usage we use the Golub data set on acute leukemia [4] as it is stored in the golubEsets experimental data package. For preparing the data, issue the following commands:

```r
> library(golubEsets)
> data(Golub_Merge)
```

2.1 Initialize k-means with divisive hierarchical cluster centroids

K-means clustering critically depends on its initialization step. We derive an initialization based on the first split of a divisive hierarchical clustering (Chapter 6 in [9]). Of the resulting two clusters, we compute centroids which provide the starting points for the k-means algorithm [10]. This has been shown to outperform standard k-means with random starting points [11]. In fact, k-means is used to refine individual clusters and to correct inappropriate assignments made by the hierarchical method.

We have packed this procedure into the function diana2means of the here described package. In addition this function computes DLD-scores for the generated splits using the implementation taken from the ISIS package [8]. Actually, the determined split is only returned when return.cut is set to true. The following snippet of code uses the 10% most variable genes in the Golub-dataset to generate a split.

```r
> e <- exprs(Golub_Merge)
> vars <- apply(e, 1, var)
```
> e <- e[vars > quantile(vars, 0.9), ]
> diana2means(e)

[1] 9.983408

> diana2means(e, return.cut = TRUE)

2-means split holding 72 elements
  distribution is: 52 20
  DLD-score: 9.983408

This function returns a single number representing the splits DLD-score as default, when the argument return.cut is set to FALSE. Otherwise an object of class split holding the list elements cut and score is returned. For instance, the split attributions can be extracted as follows:

> x <- diana2means(e, return.cut = TRUE)
> x$cut

[1,] 1 0 0 1 0 0 0 1 1 0 1 1
[1,] 0 0 0 0 0 1 1 1 0 0
[1,] 1 0 1 1 1 1 1 0 1 0
[1,] 1 1 0 0 0 0 0 0 1 0
[1,] 0 0 0 0 0 0 0 0 0 0
[1,] 0 0 0 0 0 0 0 0 0 0
[1,] 1 0 0 0 0 0 0 0 0 0

For the computation of the DLD-score, this function takes into account the best scoring ngenes genes (rows). In order to avoid excessive influence of single strikingly well separating genes, the ignore.genes argument can be used to dismiss the strongest genes from the score.
2.2 Annotation-driven splits

The central function for the generation of annotation driven splits is called \texttt{adSplit}. It calls the above described function \texttt{diana2means} on a restricted set of genes. This function has a series of arguments, but its basic call is as follows:

\begin{verbatim}
> adSplit(Golub_Merge, "GO:0006915", "hu6800")
\end{verbatim}

Evaluating identifier GO:0006915 with 769 probesets...
-> skipped, too many probes associated (769)
Empty split set

This command generates an annotation-driven split for the term "apoptosis" encoded by the identifier "GO:0006915". The string given as the chip's name is used to load the annotation meta-data. Thus \texttt{adSplit} expects a library of the same name to be installed, where it looks for the hashes \texttt{<chip-name>GO2ALLPROBES} and \texttt{<chip-name>PATH2PROBE} as they are provided by Bioconductor meta data packages. If we issue a similar command for the term "signal transduction" (GO:0007165), the following happens:

\begin{verbatim}
> adSplit(Golub_Merge, "GO:0007165", "hu6800")
\end{verbatim}

Evaluating identifier GO:0007165 with 2692 probesets...
-> skipped, too many probes associated (2692)
Empty split set

This term has too many associated genes. Hence, it is skipped. In order to generate splits related to more generic terms, we can provide an explicit maximum limit for the amount of annotated genes to terms of interest. For instance:

\begin{verbatim}
> adSplit(Golub_Merge, "GO:0007165", "hu6800", max.probes = 7000)
\end{verbatim}

Evaluating identifier GO:0007165 with 2692 probesets...
Annotation-driven split set
holds 1 split on 72 elements
associated to: signal transduction (GO:0007165)
object distribution is: 61 11
score is: 14.13467
no empirical p-values computed

This command generates the wanted split based on more than 2000 genes. \texttt{adSplit} returns an object of class \texttt{splitSet} with the following list elements:
1. **cuts**: a matrix of split attributions. One row per annotation identifier (GO term or KEGG pathway for which a split has been generated. One column per object in the dataset.

2. **score**: one score per generated split

3. **pvalue**: one empirical p-value per generated split, or **NULL**.

The object may also be empty in which case all elements are **NULL**.
Chapter 3

Empirical p-Values for Splits

In order to generate empirical p-values for a given annotation-driven gene set, we suggest to sample random gene sets of the same size and apply the split generation algorithm implemented in \texttt{diana2means} to all of these gene sets. DLD-scores computed for the random gene sets are used to approximate the score’s null distribution. The fraction of the scores from this null distribution, which are higher than the observed score for the gene set with common annotation, is used as empirical p-value. Some details on how to compute these p-values with \texttt{adSplit} are collected in this chapter.

3.1 Drawing random gene sets

The first step needed for the random sampling is drawing a given number of probe-sets measured in dataset. In order to reflect one obvious characteristic of annotation based gene sets, once we have drawn one probe set at random, we always include all other probe-sets representing the same gene into the random selection. In order to speed up the repeated drawing procedure, we use a hash containing one entry per EntrezGene identifier holding all associated probe-sets. This environment is generated from the meta-data package \texttt{ENTREZID-hash} as follows:

\begin{verbatim}
> EID2PSenv <- makeEID2PROBESenv(hu6800ENTREZID)
\end{verbatim}

The returned hash is used as follows to draw random sets of probe-sets:

\begin{verbatim}
> drawRandomPS(10, EID2PSenv, ls(EID2PSenv))
\end{verbatim}

\begin{verbatim}
  2713  6992  3706
"X78711_at" "X81003_at" "X54938_at"
  8128  901 10633
"HG3962-HT4232_at" "U47414_at" "Y07847_at"
\end{verbatim}
drawRandomPS returns a named vector holding random probe-set identifiers. The names of this vector corresponds to the associated LOCUSLINK identifiers.

### 3.2 Generating DLD-score distributions

The functions for drawing random sets of probe-sets described in the previous section are combined with `diana2means` to generate null-distributions of DLD-scores. This is implemented in the function `randomDiana2means` which is used as follows:

```R
> scores <- randomDiana2means(20, exprs(Golub_Merge), "hu6800",
+                           ndraws = 1000)

determining 1000 random DLD-scores with 20 probe sets each (wait for 10 dots)
...........
```

The form of this distribution is skewed. The parameter `ignore.genes` changes this shape towards a more symmetric shape as shown in the next figure. This is intuitive, since we remove genes which drive DLD-scores and thus approach more closely a random distribution.

```R
> scores2 <- randomDiana2means(20, exprs(Golub_Merge),
+                           "hu6800", ndraws = 1000, ignore.genes = 5)

determining 1000 random DLD-scores with 20 probe sets each (wait for 10 dots)
...........
```

```R
> par(mfrow = c(1, 2))
> hist(scores, nclass = 30, main = "", col = "grey")
> hist(scores2, nclass = 30, main = "", col = "grey")
```
The left histogram here is the score distribution when including all genes, the right one results from exclusion of best scoring genes.

### 3.3 Adding empirical p-values while generating annotation driven splits

Finally, we can use the distribution generated with random sets of probe sets to compute empirical p-values. This is done by the `adSplit` function when \( B \) is specified, the number of samplings to be used.

```r
> glutamSplits <- adSplit(Golub_Merge, "KEGG:00251", "hu6800",
+                        B = 1000)
```

Evaluating identifier KEGG:00251 with 22 probesets...

This call returns an object of class `splitSet` with an additional entry called `pvalue`. The print method for the `splitSet` gives a summary on sets of splits and some additional information, if a split set contains only 1 split:

```r
> print(glutamSplits)

Empty split set
NULL
```
Chapter 4

Working with Split-Sets

4.1 Generating split-sets

The function `adSplit` accepts more than one annotation identifier in one call. In this case it generates splits for each of the provided identifier and collects the results into one single return object. For instance:

```r
> x <- adSplit(Golub_Merge, c("GO:0007165", "GO:0006915"),
+ "hu6800", max.probes = 7000)
```

Evaluating identifier GO:0007165 with 2692 probesets...
Evaluating identifier GO:0006915 with 769 probesets...

```r
> print(x)
```

Annotation-driven split set
- holds 2 splits on 72 elements
  - scores range is: 12.37441 14.13467
  - no empirical p-values computed

If the user doesn’t want to specify few terms of interest in this fashion, he/she may also provide one of the following specially treated identifiers:

1. GO: all available GO terms are used as gene set candidates.
2. KEGG: all available KEGG pathways are used as gene set candidates.
3. all: both these sets of annotation identifiers are used.

For instance, the following command determines all splits driven by KEGG pathways without sampling random gene lists for significance analysis:
> x <- adSplit(Golub_Merge, "KEGG", "hu6800")

> print(x)

Annotation-driven split set
  holds 108 splits on 72 elements
  scores range is: 2.621651 13.26944
  no empirical p-values computed

However, if empirical p-values are computed by sampling random gene lists, multiple
testing is an issue to be considered. We use the `multtest` to correct our p-values and thus
computing false discovery rates by the method suggested by Benjamini-Hochberg. The
 corresponding q-values are stored in the result’s list element called `qvalues`. You may
consider for illustration the precomputed object `golubKEGGSplits`:

> data(golubKEGGSplits)
> print(golubKEGGSplits)

Annotation-driven split set
  holds 70 splits on 72 elements
  scores range is: 3.382672 17.31385
  empirical p-values range is: 0.005 0.955
  q-value range is: 0.1633333 0.955

> summary(golubKEGGSplits$qvalues)

    Min. 1st Qu.  Median    Mean  3rd Qu.   Max.
 0.1633  0.4375  0.6681  0.6098  0.7745  0.9550

This object has been precomputed with the following command:

> golubKEGGSplits <- adSplit(Golub_Merge, "KEGG", "hu6800", B=1000)

It thus contains all splits driven by KEGG pathways and holds empirical p-values
deduced from 1000 random gene lists. Pathways which have fewer than 20 genes associated
are thereby skipped from the analysis.

### 4.2 Graphical illustrations

For showing the illustration utilities implemented in `adSplit`, we use the precomputed
object `golubKEGGSplits` included in the `adSplit` package.
The `golubKEGGSplits` object contains 70 splits and corresponding empirical p-values and corrected q-values. The split set is ordered according to p-values such that the first entries are the most significant ones. In order to get an overview of whether significant splits and how many of them are generated, the package `adSplit` offers a histogram method for objects of class `splitSet` called as follows:

```r
data(golubKEGGSplits)
hist(golubKEGGSplits)
```

![Distribution of p-Values](image)

In this histogram the empirical p-values are drawn as usual in a histogram. In addition, the corresponding q-values corrected for multiple testing are plotted as a line into the histogram. The corresponding scale is shown to the left of the plot.

An `image` method on the same objects can be used to get an actual representation of all splits generated. A filter argument called `filter.fdr` is used to focus on splits with a low false discovery rate. The following call requires a set of splits with less than 30% expected false positives.

```r
image(golubKEGGSplits, filter.fdr=0.3)
```
In this image, each column corresponds to a patient and each row corresponds to an annotation. The colors represent to which group the corresponding patient is attributed with respect to the corresponding annotation. This image is clustered in both directions in order to bring similar splits as well as similar patients close together.

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Chapter 5

Bibliography


