Protein Microarray Data Analysis using the PAA Package

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1 Introduction

1.1 General information

Protein Array Analyzer (*PAA*, *Turewicz et al.* [1]) is a package for protein microarray data analysis (esp., *ProtoArray* data). It imports single color (protein) microarray data that has been saved in 'gpr' file format. After preprocessing (background correction, batch filtering, normalization) univariate feature preselection is performed (e.g., using the "minimum M statistic" approach - hereinafter referred to as "mMs", [2]). Subsequently, a multivariate feature selection is conducted to discover biomarker candidates. For this purpose, either a frequency-based backwards elimination approach or ensemble feature selection can be used. *PAA* provides a complete toolbox of analysis tools including several different plots for results examination and evaluation.

In this vignette the general workflow of *PAA* will be outlined by analyzing an exemplary data set that accompanies this package.

1.2 Installation

The recommended way to install PAA is to type the commands described below in the R console (note: an active internet connection is needed):

```
> # only if you install a Bioconductor package for the first time
> if (!requireNamespace("BiocManager", quietly=TRUE))
+ install.packages("BiocManager")
> # else
> library("BiocManager")
> BiocManager::install("PAA", dependencies=TRUE)
```

This will install PAA including all dependencies.

Furthermore, *PAA* has an external dependency that is needed to provide full functionality. This external dependency is the free C++ software package "*Random Jungle*" that can be downloaded from http://www.randomjungle.de/. Note: *PAA* will be usable without *Random Jungle*. However, it needs this package for random jungle recursive feature elimination (*RJ-RFE*) provided by the function selectFeatures(). Please follow the instructions for your OS in the README file to install *Random Jungle* properly on your machine.

2 Loading *PAA* and importing data

AD3 GSM734835_PA42006_-AD3.gpr

After launching R, the first step of the exemplary analysis is to load PAA.

> library(PAA)

3

New microarray data should be imported using the function loadGPR() which is mainly a wrapper to *limma*'s function read.maimages() featuring optional duplicate aggregation for *ProtoArray* data. *PAA* supports the import of files in 'gpr' file format. The imported data is stored in an expression list object (*EList*, respectively, *EListRaw*, see Bioconductor package *limma*). Paths to a targets file and to a folder containing 'gpr' files (all 'gpr' files in this folder that are listed in the targets file will be read) are mandatory arguments. The folder that can be obtained by the command system.file("extdata", package = "PAA") contains an exemplary targets file that can be used as a template. Below, the first 3 rows of this targets file are shown.

```
> targets <- read.table(file=list.files(system.file("extdata", package="PAA"),
+ pattern = "^targets", full.names = TRUE), header=TRUE)
> print(targets[1:3,])
ArrayID FileName Group Batch Date Array SerumID
1 AD1 GSM734833_PA41992_-_AD1.gpr AD Batch1 10.11.2010 41992 AD1
2 AD2 GSM734834_PA41994_-_AD2.gpr AD Batch2 10.11.2010 41994 AD2
```

The columns "ArrayID", "FileName", and "Group" are mandatory. "Batch" is mandatory for microarray data that has been processed in batches. The remaining three columns as well as custom columns containing further information (e.g., clinical data) are optional.

AD Batch1 12.11.2010 42006

AD3

If array.type is set to "ProtoArray" duplicate spots can be aggregated. After importing, the object can be saved in a '*.RData' file for further sessions. In the following code chunk, loadGPR() is demonstrated using an exemplary dummy data set that comes with PAA and has been created from the real data set analyzed in this vignette.

```
> gpr <- system.file("extdata", package="PAA")
> targets <- list.files(system.file("extdata", package="PAA"),
+ pattern = "dummy_targets", full.names=TRUE)
> dummy.elist <- loadGPR(gpr.path=gpr, targets.path=targets,
+ array.type="ProtoArray")
> save(dummy.elist, file=paste(gpr, "/DummyData.RData",
+ sep=""), compress="xz")
```

In many gpr files the mandatory column "Description" is not available and the needed information is stored in another column. In order to process such gpr files loadGPR() constructs a makeshift "Description" column. If array.type is set to "ProtoArray" this is performed automatically. Otherwise, loadGPR() provides the arguments description, de scription.features and description.discard to pass the information needed for the construction of the makeshift "Description" column. In the following code chunk these arguments are demonstrated.

```
> targets2 <- list.files(system.file("extdata", package="PAA"),
+ pattern = "dummy_no_descr_targets", full.names=TRUE)
> elist2 <- loadGPR(gpr.path=gpr, targets.path=targets2, array.type="other",</pre>
```

+ description="Name", description.features="^Hs~", description.discard="Empty")

PAA comes with an exemplary protein microarray data set. This 20 Alzheimer's disease serum samples vs. 20 controls data is a subset of a publicly available *ProtoArray* data set. It can be downloaded from the repository "*Gene Expression Omnibus*" (*GEO*, http://www.ncbi.nlm.nih.gov/geo/, record "GSE29676"). It has been contributed by *Nagele E et al.* [3] (note: Because a data set stored in 'gpr' files would be too large to accompany this package the exemplary data is stored as an '*.RData' file).

In the following code chunk, the *PAA* installation path (where exemplary data is located) is localized, the new folder 'demo_output' (where all output of the following analysis will be saved) is created, and the exemplary data set is loaded (note: here, exceptionally not via LoadGPR()).

- > cwd <- system.file(package="PAA")</pre>
- > dir.create(paste(cwd, "/demo/demo_output", sep=""))
- > output.path <- paste(cwd, "/demo/demo_output", sep="")</pre>
- > load(paste(cwd, "/extdata/Alzheimer.RData", sep=""))

3 Preprocessing

Before preprocessing the microarrays should be inspected visually for any strong spatial biases. In *PAA* this can be done using the function plotArray() which plots the imported microarray data in the original arrangement mimicking the original scan image. Thus, plotArray() is a visualization tool that can be used to visualize arrays for which the original scan image is not available. Visual inspection of the spatial expression pattern can then identify possible local tendencies and strong spatial biases. Moreover, the array can be inspected at all stages of the preprocessing workflow in order to check the impact of the particular methods that have been applied.

Consequently, as a first step, always the plots of the foreground signals (data.type="fg") should be compared with the plots of the background signals (data.type="bg").

> plotArray(elist=elist, idx=3, data.type="bg", log=FALSE, normalized=FALSE,

+ aggregation="min", colpal="topo.colors")



9

Value

AD3 Array Plot

- > plotArray(elist=elist, idx=3, data.type="fg", log=FALSE, normalized=FALSE,
- + aggregation="min", colpal="topo.colors")



In the exemplary plots shown above there is a relatively strong artifact in the background signal that has a slight impact on the foreground signal. In both figures an irregular stripe runs from the upper right part of the array to the bottom left corner. Although in the background plot this artifact is more visible than in the foreground plot the corresponding spatial bias is obvious.

For background correction *limma*'s function backgroundCorrect() can be used:

```
> library(limma)
```

```
> elist <- backgroundCorrect(elist, method="normexp",</pre>
```

```
+ normexp.method="saddle")
```

If the microarrays were manufactured or processed in lots/batches, data analysis will suffer from batch effects resulting in wrong results. Hence, the elimination of batch effects is a crucial step of data preprocessing. A simple method to remove the most obvious batch effects is to find features that are extremely differential in different batches. In *PAA* this can be done for two batches using the function **batchFilter()**. This function takes an *EList* or *EListRaw* object and the batch-specific column name vectors **lot1** and **lot2** to find differential features regarding the batches/lots. For this purpose, thresholds for p-values (Student's t-test) and fold changes can be defined. To visualize the differential features a volcano plot is drawn. Finally, the differential features are removed and the remaining data is returned.

```
> lot1 <- elist$targets[elist$targets$Batch=='Batch1','ArrayID']</pre>
```

```
> lot2 <- elist$targets[elist$targets$Batch=='Batch2','ArrayID']</pre>
```

```
> elist.bF <- batchFilter(elist=elist, lot1=lot1, lot2=lot2, log=FALSE,</pre>
```

```
+ p.thresh=0.001, fold.thresh=3)
```



For multi-batch scenarios (i.e., where the number of distinct batches is larger than two) *PAA* provides the function **batchFilter.anova()**. This function calculates p-values via an one-way analysis of variance (ANOVA) in order to identify features which are differential regarding at least two batches. Apart from that this function works analogously to **batchFilter()** which has been designed for the scenario of two batches.

```
> elist.bF.a <- batchFilter.anova(elist=elist, log=FALSE, p.thresh=0.001,
+ fold.thresh=3)
> elist <- elist.bF</pre>
```

Another important step in preprocessing is normalization. To assist in choosing an appropriate normalization method for a given data set, *PAA* provides two functions: plotNormMethods() and plotMAPlots(). plotNormMethods() draws sample-wise boxplots of raw data and data after all kinds of normalization provided by *PAA*. For each normalization approach a plot containing all sample-wise boxplots is created. All boxplots will be saved as a high-quality 'tiff' file, if an output path is specified.

> plotNormMethods(elist=elist)

plotMAPlots() draws MA plots of raw data and data after applying all kinds of normalization methods provided by *PAA*. If idx="all" and an output path is defined (default), for each microarray one 'tiff' file containing MA plots will be created. If idx is an integer indicating the column index of a particular sample, MA plots only for this sample will be created.

> plotMAPlots(elist=elist, idx=10)



After choosing a normalization method, the function normalizeArrays() can be used in order to normalize the data. normalizeArrays() takes an *EListRaw* object, normalizes the data, and returns an *EList* object containing normalized data in log2 scale. As normalization methods "cyclicloess", "quantile" or "vsn" can be chosen. Furthermore, for *ProtoArrays* robust linear normalization ("rlm", see *Sboner A. et al.* [4]) is provided.

```
> elist <- normalizeArrays(elist=elist, method="cyclicloess",
+ cyclicloess.method="fast")
```

In addition to batchFilter(), the function batchAdjust() can be used after normalization via normalizeArrays() to adjust the data for batch effects. This is a wrapper to *sva*'s function ComBat() for batch adjustment using the empirical Bayes approach [5]. To use batchAdjust() the targets file information of the *EList* object must contain the columns "Batch" and "Group".

```
> elist <- batchAdjust(elist=elist, log=TRUE)
Standardizing Data across genes</pre>
```

After preprocessing, the corrected data can be inspected via plotArray() again. E.g., now the plot of the ProtoArray already plotted shows that the revealed spatial bias is less obvious after preprocessing.

```
> plotArray(elist=elist, idx=3, data.type="fg", log=TRUE, normalized=TRUE,
+ aggregation="min", colpal="topo.colors")
```







Because for further analysis also data in original scale will be needed, a copy of the *EList* object containing unlogged data should be created.

> elist.unlog <- elist
> elist.unlog\$E <- 2^(elist\$E)</pre>

4 Differential analysis

The goal of univariate differential analysis is to detect relevant differential features. Therefore, statistical measures such as p-values or fold changes are considered. *PAA* provides plotting functions in order to depict the number and the quality of the differential features in the data set. Accordingly, the function volcanoPlot() draws a volcano plot to visualize differential features. For this purpose, thresholds for p-values and fold changes can be defined. Furthermore, the p-value computation method ("mMs" or "tTest") can be set. When an output path is defined (via output.path) the plot will be saved as a 'tiff' file. In the next code chunk, an example with method="tTest" is given.

```
> c1 <- paste(rep("AD",20), 1:20, sep="")</pre>
```

```
> c2 <- paste(rep("NDC",20), 1:20, sep="")</pre>
```

```
> #volcanoPlot(elist=elist.unlog, group1=c1, group2=c2, method="tTest",
```

```
> volcanoPlot(elist=elist, group1=c1, group2=c2, log=TRUE, method="tTest",
```

Volcano Plot

```
+ p.thresh=0.01, fold.thresh=2)
```

io 0 9 ω -log10(p-value) C 0 ശ С 4 O 0 \sim 0 -3 -2 0 2 -4 -1 1 log2(fold change)

Here, an example with method="mMs" is given:

- > mMs.matrix1 <- mMs.matrix2 <- mMsMatrix(x=20, y=20)</pre>
- > volcanoPlot(elist=elist.unlog, group1=c1, group2=c2, log=FALSE, method="mMs",
- + p.thresh=0.01, fold.thresh=2, mMs.matrix1=mMs.matrix1,
- + mMs.matrix2=mMs.matrix2, above=1500, between=400)

Another plotting function is pvaluePlot() which draws a plot of p-values for all features in the data set (sorted in increasing order and in log2 scale). The p-value computation method ("tTest" or "mMs") can be set via the argument method. Furthermore, when adjust=TRUE adjusted p-values (method: Benjamini & Hochberg, 1995, computed via p.adjust()) will be used. For a better orientation, horizontal dashed lines indicate which p-values are smaller than 0.05 and 0.01. If adjust=FALSE, additionally, the respective Bonferroni significance threshold (to show p-values that would be smaller than 0.05 after a possible Bonferroni correction) for the given data is indicated by a third dashed line. Note: Bonferroni is not used for the adjustment. The dashed line is for better orientation only. When an output path is defined (via output.path) the plot will be saved as a 'tiff' file. In the next code chunk, an example with method="tTest" is given.

> pvaluePlot(elist=elist, group1=c1, group2=c2, log=TRUE, method="tTest")



Here, an example with method="mMs" is given:

> mMs.matrix1 <- mMs.matrix2 <- mMsMatrix(x=20, y=20)</pre>

- > pvaluePlot(elist=elist.unlog, group1=c1, group2=c2, log=FALSE, method="mMs",
- + mMs.matrix1=mMs.matrix1, mMs.matrix2=mMs.matrix2, above=1500,
- + between=400)

Here, an example with method="tTest" and adjust=TRUE is given:

```
> pvaluePlot(elist=elist, group1=c1, group2=c2, log=TRUE, method="tTest",
+ adjust=TRUE)
```



Here, an example with method="mMs" and adjust=TRUE is given:

```
> pvaluePlot(elist=elist.unlog, group1=c1, group2=c2, log=FALSE, method="mMs",
```

```
+ mMs.matrix1=mMs.matrix1, mMs.matrix2=mMs.matrix2, above=1500,
```

```
+ between=400, adjust=TRUE)
```

Finally, diffAnalysis() performs a detailed univariate differential analysis. This function takes an EList\$E- or EListRaw\$E- matrix (e.g., temp <- elist\$E) extended by row names comprising "BRC"-IDs of the corresponding features. The BRC-IDs can be created via:

brc <- paste(elist\$genes[,1], elist\$genes[,3], elist\$genes[,2]).</pre>

Next, the row names can be assigned as follows: rownames(temp) <- brc. Furthermore, the corresponding column name vectors, group labels and mMs- parameters are needed to perform the univariate differential analysis. This analysis covers inter alia p-value computation, p-value adjustment (method: Benjamini & Hochberg, 1995), and fold change computation. Since the results table is usually large, a path for saving the results should be defined via out put.path. Optionally, a vector of row indices (features) and additionally (not mandatory for subset analysis) a vector of corresponding feature names (feature.names) can be forwarded to perform the analysis for a feature subset.

```
> E <- elist.unlog$E
> rownames(E) <- paste(elist.unlog$genes[,1], elist.unlog$genes[,3],
+ elist.unlog$genes[,2])
> write.table(x=cbind(rownames(E),E),
+ file=paste(cwd,"/demo/demo_output/data.txt", sep=""), sep="\t", eol="\n",
+ row.names=FALSE, quote=FALSE)
> mMs.matrix1 <- mMs.matrix2 <- mMsMatrix(x=20, y=20)</pre>
```

> diff.analysis.results <- diffAnalysis(input=E, label1=c1, label2=c2,</pre> class1="AD", class2="NDC", output.path=output.path, + + mMs.matrix1=mMs.matrix1, mMs.matrix2=mMs.matrix2, above=1500, between=400) + > print(diff.analysis.results[1:10,]) BRC FDR.t. min..M.stat...mMs. t.test 1 1 2 11 0.352751286646465 0.65394032025644 0.243589743589744 2 1 2 13 0.15129653649193 0.501988087542915 0.0241860325286354 3 1 2 15 0.320875589014851 0.634210502468438 1 1 2 17 0.178493270666023 0.52723150928707 0.150422391245528 4 5 1 2 19 0.271589613948768 0.597932135616545 0.243589743589744 6 1 2 21 0.070766604220371 0.394970369965038 0.0457380457380457 7 1 3 1 0.0284911744794212 0.268607060873835 1 8 1 3 3 0.00921968894879156 0.142280968989524 0.5 9 1 3 5 0.00592168460061224 0.106226964810384 0.053014553014553 10 1 3 7 0.806173147194643 0.916805614435089 0.302494802494803 FDR.mMs. fold.change mean.AD mean.NDC 1 0.835015538626552 1.36257910953312 1384.85432520349 1016.34783295481 2 0.330335726331277 0.260784177611614 2189.55822304562 8396.05471121231 3 1 1.10224097047813 450.550786587284 408.758881818594 4 0.835015538626552 0.595491172830502 1518.91515397688 2550.69297964089 0.835015538626552 0.454051309774644 2530.1344545838 5572.35360875748 5 6 0.484098604098604 0.759043204421858 2636.25422481143 3473.12802414112 7 1 1.26294406395723 484.784697258338 383.852864979107 8 0.910368401063426 1.48059072464481 692.098536568906 467.447570114244 9 0.484098604098604 1.36023092627671 1991.44021666422 1464.045683857 10 0.910368401063426 0.908397966805547 818.482441505168 901.017474074085 median.AD median.NDC sd.AD sd.NDC 1 839.980702981771 857.735001801976 1643.99677640781 564.430270630399 2 1305.46661974481 2548.69868040326 2970.84080119717 18364.0058765935 3 413.716205403229 417.428604869083 165.734824607583 82.0150145643133 4 1215.20762270891 1689.1444916955 1062.44868297829 3151.76500089474 5 1824.40003073847 1864.76465290003 2444.44412207378 11793.1087294661 6 2248.99036532219 2924.27002197574 1276.31835442119 1553.96096388883 7 446.558759285451 349.263285007632 154.993619177929 122.909766431916 556.15638977055 455.41188100012 339.123207021637 93.501996624578 8 9 1873.36646648453 1436.18751389589 718.441427077343 323.488045744026 10 730.232223314623 469.463840773123 433.002023903254 1422.22599366312

Subsequently, the most relevant differential features (i.e., features having low p-values and high absolute fold changes) can be extracted as a univariate feature selection. Nevertheless, it is recommended to perform also multivariate feature selection and to consider feature panels obtained from both approaches.

5 Feature preselection

Before multivariate feature selection will be performed, it is recommended to discard features that are obviously not differential. Discarding them will accelerate runtimes without any negative impact on results. In *PAA*, this task is called *"feature preselection"* and it is performed by the function preselect(). This function iterates all features of the data set to score them via *mMs*, *Student's t-test*, or *mRMR*. If discard.features is TRUE (default), all features that are considered as obviously not differential will be collected and returned for discarding. Which features are considered as not differential depends on the parameters method, discard.threshold, and fold.thresh.

- If method = "mMs" features having an *mMs* value larger than discard.threshold (here: numeric between 0.0 and 1.0) or do not satisfy the minimal absolute fold change fold.thresh will be considered as not differential.
- If method = "tTest" features having a p-value larger than discard.threshold (here: numeric between 0.0 and 1.0) or do not satisfy the minimal absolute fold change fold.thresh will be considered as not differential.
- If method = "mrmr" mRMR scores for all features will be computed as scoring method (using the function mRMR.classic() of the R package mRMRe). Subsequently, features that are not the discard.threshold (here: integer indicating a number of features) features having the best mRMR scores are considered as not differential.

```
> mMs.matrix1 <- mMs.matrix2 <- mMsMatrix(x=20, y=20)
> pre.sel.results <- preselect(elist=elist.unlog, columns1=c1, columns2=c2,
+ label1="AD", label2="NDC", log=FALSE, discard.threshold=0.5,
+ fold.thresh=1.5, discard.features=TRUE, mMs.above=1500, mMs.between=400,
+ mMs.matrix1=mMs.matrix1, mMs.matrix2=mMs.matrix2,
+ method="mMs")</pre>
```

```
> elist <- elist[-pre.sel.results$discard,]</pre>
```

6 Feature selection

For multivariate feature selection *PAA* provides the function **selectFeatures()**. It performs a multivariate feature selection using *"frequency-based"* feature selection (based on *RF-RFE*, *RJ-RFE* or *SVM-RFE*) or *"ensemble"* feature selection (based on *SVM-RFE*).

Frequency-based feature selection (method="frequency"): The whole data is splitted in k cross validation training and test set pairs. For each training set a multivariate feature selection procedure is performed. The resulting k feature subsets are tested using the corresponding test sets (via classification). As a result, **selectFeatures()** returns the average k-fold cross validation classification accuracy as well as the selected feature panel (i.e., the union set of the k particular feature subsets). As multivariate feature selection methods random forest recursive feature elimination (*RF-RFE*), random jungle recursive feature elimination (*RJ-RFE*) and support vector machine recursive feature elimination (*SVM-RFE*) are supported. To reduce running times, optionally, an additional univariate preselection methods mMs ("mMs"), Student's t-test ("tTest") and mRMR ("mrmr") are supported. Alternatively, no preselection can be chosen ("none"). This approach is similar to the method proposed in *Baek et al.* [6].

Ensemble feature selection (method="ensemble"): From the whole data a previously defined number of subsamples is drawn defining pairs of training and test sets. Moreover, for each training set a previously defined number of bootstrap samples is drawn. Then, for each bootstrap sample *SVM-RFE* is performed and a feature ranking is obtained. To obtain a final ranking for a particular training set, all associated bootstrap rankings are aggregated to a single ranking. To score the cutoff best features, for each subsample a classification of the test set is performed (using a svm trained with the cutoff best features from the training set) and the classification accuracy is determined. Finally, the stability of the subsample-specific panels is assessed (via Kuncheva index, *Kuncheva LI, 2007* [7]), all subsample-specific rankings are aggregated, the top n features (defined by cutoff) are selected, the average classification accuracy is computed, and all these results are returned in a list. This approach has been proposed in *Abeel et al.* [8].

selectFeatures() takes an EListRaw or EList object, group-specific sample numbers, group labels and parameters choosing and configuring a multivariate feature selection method (frequency-based or ensemble feature selection) to select a panel of differential features. When an output path is defined (via output.path) results will be saved on the hard disk and when verbose is TRUE additional information will be printed to the console. Depending on the selection method, one of two different results lists will be returned:

- 1. If method is "frequency", the results list contains the following elements:
 - accuracy: average k-fold cross validation accuracy.
 - sensitivity: average k-fold cross validation sensitivity.
 - specificity: average k-fold cross validation specificity.
 - features: selected feature panel.
 - all.results: complete cross validation results.
- 2. If method is "ensemble", the results list contains the following elements:
 - accuracy: average accuracy regarding all subsamples.

- sensitivity: average sensitivity regarding all subsamples.
- specificity: average specificity regarding all subsamples.
- features: selected feature panel.
- all.results: all feature ranking results.
- stability: stability of the feature panel (i.e., Kuncheva index for the subrun-specific panels).

In the following two code chunks first *"frequency-based"* feature selection and then *"ensemble"* feature selection is demonstrated.

> selectFeatures.results <- selectFeatures(elist,n1=20,n2=20,label1="AD",</pre>

- + label2="NDC",log=TRUE,selection.method="rf.rfe",subruns=2,
- + candidate.number=1000,method="frequency")

> selectFeatures.results <- selectFeatures(elist,n1=20,n2=20,label1="AD",</pre>

+ label2="NDC", log=TRUE, subsamples=10, bootstraps=10, method="ensemble")

Because runtimes would take too long for this vignette *PAA* comes with precomputed se lectFeatures.results objects stored in '*.RData' files. These objects can be loaded as follows:

- > # results of frequency-based feature selection:
- > load(paste(cwd, "/extdata/selectFeaturesResultsFreq.RData", sep=""))
- > # or results of ensemble feature selection:
- > load(paste(cwd, "/extdata/selectFeaturesResultsEns.RData", sep=""))

7 Results inspection

After the selection of a feature panel, these features should be validated by manual inspection and evaluation for further research. To aid results inspection, *PAA* provides several functions. The function plotFeatures() plots the intensities of all features (represented by BRC-IDs) that have been selected by selectFeatures() (one sub-plot per feature) in group-specific colors. All sub-plots are aggregated in one figure. If output.path is not NULL, this figure will be saved as a 'tiff' file in output.path.

> plotFeatures(features=selectFeatures.results\$features, elist=elist, n1=20, + n2=20, group1="AD", group2="NDC")



Alternatively, the function plotFeaturesHeatmap() plots intensities of all features given in the vector features (represented by BRC-IDs) as a heatmap. If description is TRUE (default: FALSE), features will be described via protein names instead of UniProtKB accessions. Again, if output.path is not NULL, the heatmap will be saved as a 'tiff' file in output.path.

- + n1=20, n2=20, description=TRUE)



As an alternative to plotFeaturesHeatmap() the function plotFeaturesHeatmap.2() which is based on the *gplots* function heatmap.2() gives similar plots with some additional information. Apart from that both functions plotFeaturesHeatmap() and plotFeaturesHeatmap.2() are analogous.

> plotFeaturesHeatmap.2(features=selectFeatures.results\$features, elist=elist,

+ n1=20, n2=20, description=TRUE)



Finally, the function printFeatures() creates a table containing the selected biomarker candidate panel as well as additional information for results inspection. If output.path is defined, this table will be saved in a 'txt' file ('candidates.txt').

```
> elist$E <- round(elist$E,2)</pre>
> printFeatures(features=selectFeatures.results$features, elist=elist)[,-2]
       BRC
             AD1
                   AD2
                         AD3
                               AD4
                                     AD5
                                           AD6
                                                 AD7
                                                      AD8
                                                            AD9
                                                                AD10 AD11
1
   3 14 13
           9.59
                 9.53 8.98
                             9.52 9.96
                                         8.95
                                               9.64
                                                    9.16
                                                           9.23
                                                                9.23 8.94
2
   11 3 15 10.73 10.55 10.57 10.64 10.85 10.93 10.53 10.81
                                                          11.1 11.22 10.54
3
    14 4 3 11.83 11.3 10.83 10.72 10.59 10.43 11.19 12.2 10.5 10.62 11.1
4
   15 12 9 9.68 10.04 10.57 10.4 10.03
                                           9.9 9.61 10.24 10.19 10.83 10.67
5
   17 9 11 12.52 12.49 12.6 12.68 12.05 12.04 14.02 13.22 12.19 12.35 12.84
   24 17 3 9.26 8.74 9.66 9.32
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    25 3 7 11.85 12.99 12.25 11.57 11.8 11.71 11.59 11.81 11.53 12.01 11.88
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  25 11 21 11.31 11.2 11.73 11.71 11.18 11.14 11.25 11.67 11.48 11.58 11.78
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   29 4 13 12.52 12.26 12.63 12.65 12.67
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References

- Turewicz M, Ahrens M, May C, Marcus K, Eisenacher M. PAA: an R/Bioconductor package for biomarker discovery with protein microarrays. Bioinformatics (2016) 32 (10): 1577-1579. doi:10.1093/bioinformatics/btw037, PubMed PMID: 26803161.
- [2] Love B: The Analysis of Protein Arrays. In: Functional Protein Microarrays in Drug Discovery. CRC Press; 2007: 381-402.
- [3] Nagele E, Han M, Demarshall C, Belinka B, Nagele R (2011): Diagnosis of Alzheimer's disease based on disease-specific autoantibody profiles in human sera. PLoS One 6: e23112.
- [4] Sboner A. et al., Robust-linear-model normalization to reduce technical variability in functional protein microarrays. J Proteome Res 2009, 8(12):5451-5464.
- [5] Johnson WE, Li C, and Rabinovic A (2007) Adjusting batch effects in microarray expression data using empirical Bayes methods. Biostatistics 8:118-27.
- [6] Baek S, Tsai CA, Chen JJ.: Development of biomarker classifiers from highdimensional data. Brief Bioinform. 2009 Sep;10(5):537-46.
- [7] Kuncheva, LI: A stability index for feature selection. Proceedings of the IASTED International Conference on Artificial Intelligence and Applications. February 12-14, 2007. Pages: 390-395.

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[8] Abeel T, Helleputte T, Van de Peer Y, Dupont P, Saeys Y: Robust biomarker identification for cancer diagnosis with ensemble feature selection methods. Bioinformatics. 2010 Feb 1;26(3):392-8.