biosigner: A new method for signature discovery from omics data

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1 Introduction

High-throughput, non-targeted, technologies such as **transcriptomics**, **proteomics** and **metabolomics**, are widely used to **discover molecules** which allow to efficiently discriminate between biological or clinical conditions of interest (e.g., disease vs control states). Powerful **machine learning** approaches such as **Partial Least Square Discriminant Analysis** (PLS-DA), **Random Forest** (RF) and **Support Vector Machines** (SVM) have been shown to achieve high levels of prediction accuracy. **Feature selection**, i.e., the selection of the few features (i.e., the molecular signature) which are of highest discriminating value, is a critical step in building a robust and relevant classifier (Guyon and Elisseeff 2003): First, dimension reduction is usefull to **limit the risk of overfitting** and **increase the prediction stability** of the model; second, **intrepretation** of the molecular signature is facilitated; third, in case of the development of diagnostic product, a restricted list is required for the **subsequent validation** steps (Rifai, Gillette, and Carr 2006).

Since the comprehensive analysis of all combinations of features is not computationally tractable, several **selection techniques** have been described, including *filter* (e.g., *p*-values thresholding), *wrapper* (e.g., recursive feature elimination), and *embedded* (e.g., sparse PLS) approaches (Saeys, Inza, and Larranaga 2007). The major challenge for such methods is to be **fast** and extract **restricted and stable molecular signatures** which still provide **high performance** of the classifier (Gromski et al. 2014; Determan 2015).

2 The biosigner package

The biosigner package implements a new **wrapper** feature selection algorithm:

- 1. the dataset is split into training and testing subsets (by bootstraping, controling class proportion),
- 2. model is trained on the training set and balanced accuracy is evaluated on the test set,
- 3. the features are ranked according to their importance in the model,
- 4. the relevant feature subset at level f is found by a binary search: a feature subset is considered relevant if and only if, when randomly permuting the intensities of other features in the test subsets, the proportion of increased or equal prediction accuracies is lower than a defined threshold f,
- 5. the dataset is restricted to the selected features and steps 1 to 4 are repeated until the selected list of features is stable.

Three binary classifiers have been included in biosigner, namely **PLS-DA**, **RF** and **SVM**, as the performances of each machine learning approach may vary depending on the structure of the dataset (Determan 2015). The algorithm returns the **tier** of each feature for the selected classifer(s): tier **S** corresponds to the **final signature**, i.e., features which have been found significant in all the selection steps; features with tier *A* have been found significant in all but the last selection, and so on for tier *B* to *D*. Tier *E* regroup all previous round of selection.

As for a classification algorithm, the biosign method takes as input the x samples times features data frame (or matrix) of intensities, and the y factor (or character vector) of class labels (note that only binary classification is currently available). It returns the signature (signatureLs: selected feature names) and the trained model (modelLs) for each of the selected classifier. The plot method for biosign objects enable to visualize the individual boxplots of the selected features. Finally, the predict method allows to apply the trained classifier(s) on new datasets.

The algorithm has been successfully applied to **transcriptomics** and **metabolomics** data [Rinaudo et al. (2016); see also the *Hands-on* section below).

3 Hands-on

3.1 Loading

We first load the biosigner package:

library(biosigner)

We then use the **diaplasma** metabolomics dataset (Rinaudo et al. 2016) which results from the analysis of **plasma samples from 69 diabetic patients** were analyzed by reversed-phase liquid chromatography coupled to high-resolution mass spectrometry (**LC-HRMS**; Orbitrap Exactive) in the negative ionization mode. The raw data were pre-processed with XCMS and CAMERA (5,501 features), corrected for signal drift, log10 transformed, and annotated with an in-house spectral database. The patient's **age**, **body mass index**, and **diabetic type** are recorded (Rinaudo et al. 2016).

data(diaplasma)

We attach *diaplasma* to the search path and display a summary of the content of the *dataMatrix*, *sampleMetadata* and *variableMetadata* with the strF function from the (imported) ropls package:

```
attach(diaplasma)
library(ropls)
strF(dataMatrix)
          dim class
##
                         mode typeof
                                       size NAs min mean median max
##
   69 x 5,501 matrix numeric double 3.2 Mb
                                              0
                                                  0
                                                    4.2
                                                             4.4 8.2
             m096.009t01.6
                              m096.922t00.8 ...
                                                   m995.603t10.2
##
## DIA001 2.98126177377087 6.08172882312848 ... 3.93442594703862
                         0 6.13671997362279 ... 3.74201112636229
## DIA002
```

... ## DIA077 0 6.12515971273103 ... 4.55458598372024 ## DIA078 4.69123816772499 6.134420482337 ... 4.1816445335704 ## m995.613t10.2 ## DIA001 3.96424920154706 ## DIA002 3.78128422428722 ## ## DIA077 4.57310800324247 ## DIA078 4.20696191303494 strF(sampleMetadata) ## type age bmi ## factor numeric numeric ## nRow nCol size NAs 69 30 Mb 0 ## ## type age bmi ## DIA001 T2 70 31.6 ## DIA002 T2 67 28 ## ## DIA077 T2 50 27 ## DIA078 T2 65 29 strF(variableMetadata) mzmed rtmed ... pcgroup ## spiDb ## numeric numeric ... numeric character ## nRow nCol size NAs ## 5,501 6 0.7 Mb 0 ## mzmed rtmed ... pcgroup ## m096.009t01.6 96.00899361 93.92633015 ... 1984 ## m096.922t00.8 96.92192011 48.93274877 ... 4 ## ## m995.603t10.2 995.6030195 613.4388762 ... 7160 ## m995.613t10.2 995.6134422 613.4446705 ... 7161 ## spiDb ## m096.009t01.6 N-Acetyl-L-aspartic acid HMDB00812 ## m096.922t00.8 ## ## m995.603t10.2 ## m995.613t10.2

We see that the **diaplasma** list contains three objects:

- 1. dataMatrix: 69 samples x 5,501 matrix of numeric type containing the intensity profiles (log10 transformed),
- 2. sampleMetadata: a 69×3 data frame, with the patients'
 - type: diabetic type, factor
 - age: numeric
 - bmi: body mass index, numeric
- 3. variableMetadata: a 5,501 x 8 data frame, with the median m/z ('mzmed', numeric) and the median retention time in seconds ('rtmed', numeric) from XCMS, the 'isotopes' (character), 'adduct' (character) and 'pcgroups' (numeric) annotations from CAMERA, the names of the m/z and RT matching compounds from an in-house spectra of commercial metabolites ('name_hmdb', character), and the *p*-values resulting from the non-parametric hypothesis testing of difference in medians between types ('type_wilcox_fdr', numeric), and correlation with age ('age_spearman_fdr', numeric) and body mass index ('bmi_spearman_fdr', numeric), all corrected for multiple testing (False Discovery Rate).

We can observe that the 3 clinical covariates (diabetic type, age, and bmi) are stronly associated:

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```
with(sampleMetadata,
plot(age, bmi, cex = 1.5, col = ifelse(type == "T1", "blue", "red"), pch = 16))
legend("topleft", cex = 1.5, legend = paste0("T", 1:2),
text.col = c("blue", "red"))
```

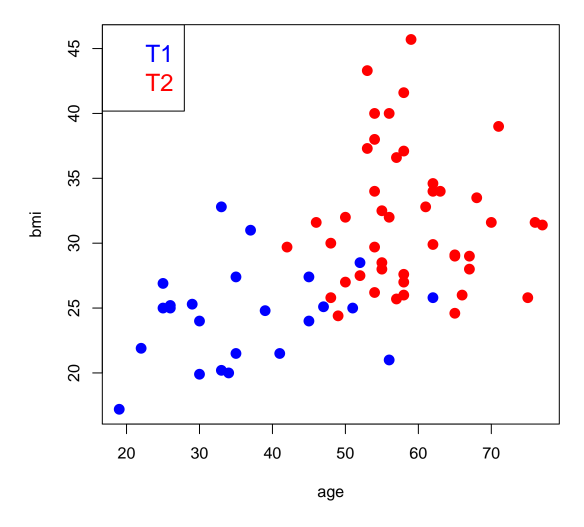


Figure 1: age, body mass index (bmi), and diabetic type of the patients from the diaplasma cohort.

3.2 Molecular signatures

Let us look for signatures of *type* in the *diaplasma* dataset by using the biosign method. To speed up computations in this demo vignette, we restrict the number of features (from 5,501 to about 500) and the number of bootstraps (5 instead of 50 [default]); the selection on the whole dataset, 50 bootstraps, and the 3 classifiers, takes around 10 min.

```
featureSelV1 <- variableMetadata[, "mzmed"] >= 450 &
variableMetadata[, "mzmed"] < 500
sum(featureSelV1)
## [1] 533</pre>
```

```
dataMatrix <- dataMatrix[, featureSelV1]</pre>
variableMetadata <- variableMetadata[featureSelV1, ]</pre>
set.seed(123)
diaSign <- biosign(dataMatrix, sampleMetadata[, "type"], bootI = 5)</pre>
set.seed(NULL)
## Significant features from 'S' groups:
##
                  plsda randomforest svm
## m471.241t07.6 "S"
                        "B"
                                      "A"
## m497.284t08.1 "S"
                        "S"
                                      "E"
## m495.261t08.7 "A"
                        "E"
                                      "S"
## m497.275t08.1 "S"
                         "A"
                                      "E"
## Accuracy:
        plsda randomforest
##
                               svm
## Full 0.759
                      0.742 0.808
## AS
        0.832
                      0.784 0.780
## S
        0.911
                      0.738 0.752
```

The arguments are:

- x: the numerical matrix (or data frame) of intensities (samples as rows, variables as columns),
- y: the factor (or character) specifying the sample labels from the 2 classes,
- methodVc: the classifier(s) to be used; here, the default all value means that all classifiers available (plsda, randomforest, and svm) are selected,
- bootI: the number of bootstraps is set to 5 to speed up computations when generating this vignette; we however
 recommend to keep the default 50 value for your analyzes (otherwise signatures may be less stable).

Note:

- If some features from the x matrix/data frame contain missing values (NA), these features will be removed prior to modeling with Random Forest and SVM (in contrast, the NIPALS algorithm from PLS-DA can handle missing values),
- 2. The set.seed command was used here to be sure that the results from this vignette can be reproduced exactly; by choosing alternative seeds (and the default bootI = 50), similar signatures are obtained, showing the stability of the selection.

The resulting signatures for the 3 selected classifiers are both printed and plotted as **tiers** from *S*, *A*, up to *E* by decreasing relevance. The (*S*) tier corresponds to the final signature, i.e. features which passed through all the backward selection steps. In contrast, features from the other tiers were discarded during the last (*A*) or previous (*B* to *E*) selection rounds.

Note that *tierMaxC* = 'A' argument in the *print* and *plot* methods can be used to view the features from the larger S+A signatures (especially when no S features have been found, or when the performance of the S model is much lower than the S+A model).

The **performance** of the model built with the input dataset (*balanced accuracy*: mean of the *sensitivity* and *specificity*), or the subset restricted to the S or S+A signatures are shown. We see that with 1 to 5 S feature signatures (i.e., less than 1% of the input), the 3 classifiers achieve good performances (even higher than the full Random Forest and SVM models). Furthermore, reducing the number of features decreases the risk of building non-significant models (i.e., models which do not perform significantly better than those built after randomly permuting the labels). The signatures from the 3 classifiers have some distinct features, which highlights the interest of comparing various machine learning approaches.

The **individual boxplots** of the features from the *complete* signature can be visualized with:

plot(diaSign, typeC = "boxplot")

Let us see the metadata of the *complete* signature:

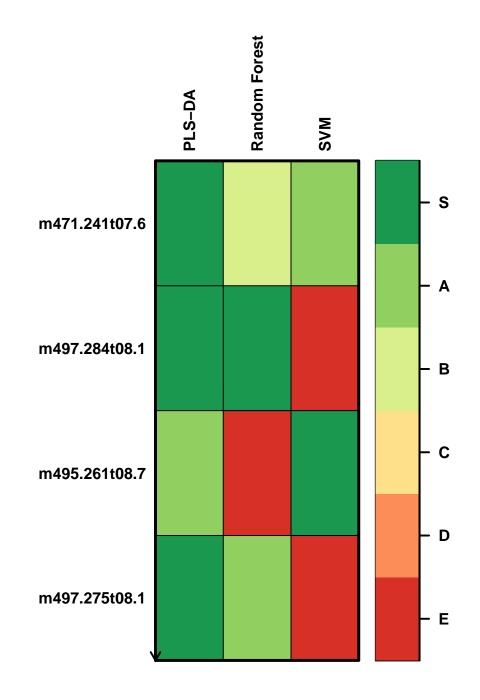
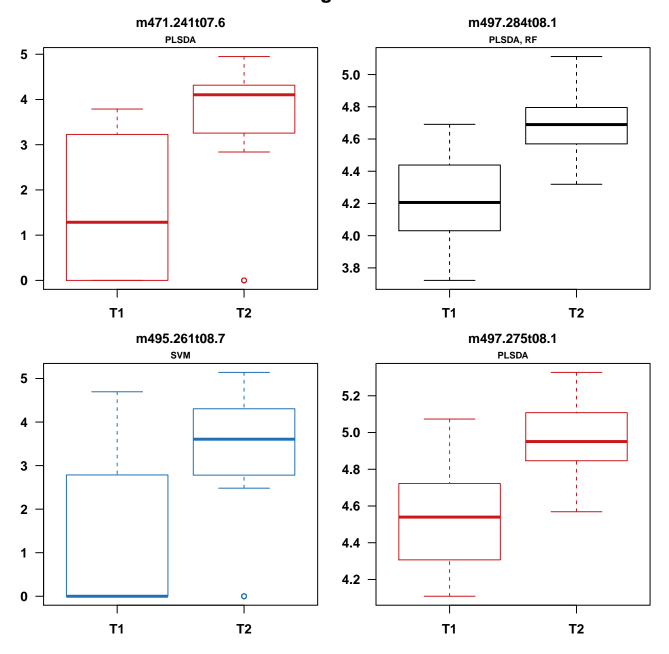


Figure 1: **Figure 2:** Relevant signatures for the *PLS-DA*, *Random Forest*, and *SVM* classifiers extracted from the diaplasma dataset. The *S* tier corresponds to the final metabolite signature, i.e., metabolites which passed through all the selection steps.

variableMetadata[getSignatureLs(diaSign)[["complete"]],]
mzmed rtmed isotopes adduct
m471.241t07.6 471.2408 455.5541
m497.284t08.1 497.2840 486.5338 [M+C1]- 462.31 [M-H]- 498.287
m495.261t08.7 495.2609 524.1249
m497.275t08.1 497.2755 486.5722 [M+C1]- 462.31 [M-H]- 498.287



'S' signature

Figure 2: Figure 3: Individual boxplots of the features selected for at least one of the classification methods. Features selected for a single classifier are colored (*red* for PLS-DA, *green* for Random Forest and *blue* for SVM).

##		pcgroup	spiDb
##	m471.241t07.6	10538	
##	m497.284t08.1	220	
##	m495.261t08.7	1655	
##	m497.275t08.1	220	Taurochenodeoxycholic acid_HMDB00951

We observe that the taurochenodeoxycholic acid has been annotated, in addition to another [M-H]- ion at 470.233 Da.

Six out of the 8 features are very significant by univariate hypothesis testings of difference between *type* medians, and to a lesser extent, of the correlation with *age* and *body mass index*.

3.3 Predictions

Let us split the dataset into a training (the first 4/5th of the 183 samples) and a testing subsets, and extract the relevant features from the training subset:

```
trainVi <- 1:floor(0.8 * nrow(dataMatrix))
testVi <- setdiff(1:nrow(dataMatrix), trainVi)</pre>
```

```
set.seed(123)
diaTrain <- biosign(dataMatrix[trainVi, ], sampleMetadata[trainVi, "type"],
bootI = 5)
set.seed(NULL)
```

```
## Significant features from 'S' groups:
##
                plsda randomforest svm
## m471.241t07.6 "B" "A"
                                  "S"
                    "S"
                                  "D"
## m497.284t08.1 "S"
                     "E"
                                  "S"
## m456.182t12.9 "E"
## m493.243t07.7 "E"
                     "E"
                                  "S"
## Accuracy:
##
       plsda randomforest
                           svm
## Full 0.794 0.777 0.793
## AS
       0.827
                   0.848 0.783
## S
              0.793 0.840
       0.843
```

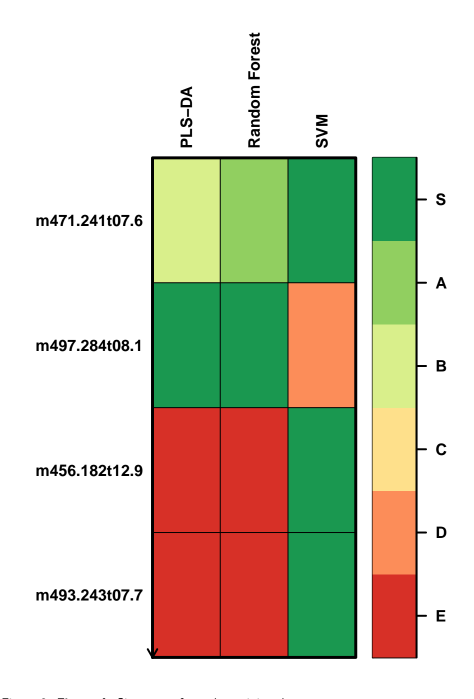
We extract the **fitted** types on the training dataset restricted to the *S* signatures:

```
diaFitDF <- predict(diaTrain)</pre>
```

We then print the confusion tables for each classifier:

```
lapply(diaFitDF, function(predFc) table(actual = sampleMetadata[trainVi,
"type"], predicted = predFc))
## $plsda
##
         predicted
## actual T1 T2
      T1 16 6
##
##
       T2 4 29
##
## $randomforest
##
        predicted
## actual T1 T2
       T1 14 8
##
       T2 7 26
##
##
## $svm
##
        predicted
## actual T1 T2
##
       T1 18 4
##
       T2 4 29
```

and the corresponding balanced accuracies:





```
sapply(diaFitDF, function(predFc) {
  conf <- table(sampleMetadata[trainVi, "type"], predFc)
  conf <- sweep(conf, 1, rowSums(conf), "/")
  round(mean(diag(conf)), 3)
})
##     plsda randomforest     svm
##     0.803     0.712     0.848</pre>
```

Note that these values are slightly different from the accuracies returned by biosign because the latter are computed by

using the resampling scheme selected by the bootl (or crossvall) arguments:

round(get	Accura	<pre>cyMN(diaTrain)["S",</pre>],	3)
##	plsda	randomforest		svm
##	0.843	0.793	0	.840

Finally, we can compute the performances on the test subset:

```
diaTestDF <- predict(diaTrain, newdata = dataMatrix[testVi, ])
sapply(diaTestDF, function(predFc) {
  conf <- table(sampleMetadata[testVi, "type"], predFc)
  conf <- sweep(conf, 1, rowSums(conf), "/")
  round(mean(diag(conf)), 3)
})
### plsda randomforest svm
## 0.750 0.667 0.750</pre>
```

3.4 Working on *ExpressionSet* omics objects from bioconductor

The **ExpressionSet** class from the Biobase bioconductor package has been developed to conveniently handle preprocessed omics objects, including the variables x samples matrix of intensities, and data frames containing the sample and variable metadata (Huber et al. 2015). The matrix and the two data frames can be accessed by the exprs, pData and fData respectively (note that the data matrix is stored in the object with samples in columns).

The biosign method can be applied to an *ExpressionSet* object, by using the object as the x argument, and by indicating as the y argument the name of the phenoData column to be used as the response.

In the example below, we will first build a minimal *ExpressionSet* object from the *diaplasma* data set, and we subsequently perform signature discovery:

```
library(Biobase)
diaSet <- ExpressionSet(assayData = t(dataMatrix),</pre>
phenoData = new("AnnotatedDataFrame", data = sampleMetadata))
set.seed(123)
biosign(diaSet, "type", bootI = 5)
set.seed(NULL)
## Significant features from 'S' groups:
##
                 plsda randomforest svm
## m471.241t07.6 "S" "B"
                                     "A"
                       "S"
                                     "E"
## m497.284t08.1 "S"
                       "E"
                                     "S"
## m495.261t08.7 "A"
## m497.275t08.1 "S"
                       "A"
                                     "E"
## Accuracy:
##
        plsda randomforest
                              svm
## Full 0.759
```

 ## Full 0.759
 0.742 0.808

 ## AS
 0.832
 0.784 0.780

 ## S
 0.911
 0.738 0.752

Note: Because of identical names in the two packages, the combine method from package Biobase is replaced by the combine method from package randomForest.

Before moving to new data sets, we detach *diaplasma* from the search path:

detach(diaplasma)

4 Extraction of biomarker signatures from other omics datasets

In this section, biosign is applied to two metabolomics and one transcriptomics data sets. Please refer to Rinaudo et al. (2016) for a full discussion of the methods and results.

4.1 Physiological variations of the human urine metabolome (metabolomics)

The **sacurine** LC-HRMS dataset from the dependent ropls package can also be used (Thevenot et al. 2015): Urine samples from a cohort of 183 adults were analyzed by using an LTQ Orbitrap in the negative ionization mode. A total of 109 metabolites were identified or annotated at the MSI level 1 or 2. Signal drift and batch effect were corrected, and each urine profile was normalized to the osmolality of the sample. Finally, the data were log10 transformed (see the ropls vignette for further details and examples).

We can for instance look for signatures of the gender:

```
data(sacurine)
set.seed(123) ##
sacSign <- biosign(sacurine[["dataMatrix"]],</pre>
sacurine[["sampleMetadata"]][, "gender"],
methodVc = "plsda")
set.seed(NULL)
## Significant features from 'S' groups:
##
                             plsda
                              "S"
## Malic acid
                              "S"
## p-Anisic acid
## Testosterone glucuronide "S"
## Accuracy:
##
        plsda
## Full 0.870
## AS
        0.861
## S
        0.879
```

4.2 Apples spikes with known compounds (metabolomics)

The **spikedApples** dataset was obtained by LC-HRMS analysis (SYNAPT Q-TOF, Waters) of one control and three spiked groups of 10 apples each. The spiked mixtures consists in 2 compounds which were not naturally present in the matrix and 7 compounds aimed at achieving a final increase of 20%, 40% or 100% of the endogeneous concentrations. The authors identified 22 features (out of the 1,632 detected in the positive ionization mode; i.e. 1.3%) which came from the spiked compounds. The dataset is included in the BioMark R package (Franceschi et al. 2012). Let us use the *control* and *group1* samples (20 in total) in this study.

```
library(BioMark)
data(SpikePos)
group1Vi <- which(SpikePos[["classes"]] %in% c("control", "group1"))
appleMN <- SpikePos[["data"]][group1Vi, ]
spikeFc <- factor(SpikePos[["classes"]][group1Vi])
annotDF <- SpikePos[["annotation"]]
rownames(annotDF) <- colnames(appleMN)</pre>
```

We can check, by using the opls method from the ropls package for multivariate analysis, that:

1. no clear separation can be observed by PCA:

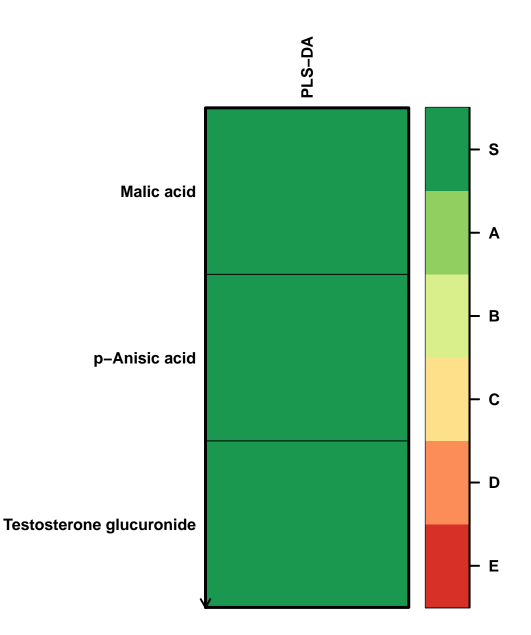
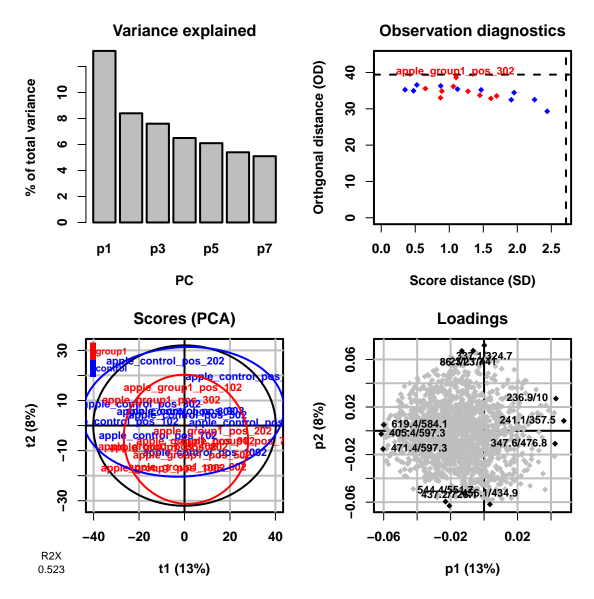


Figure 4: Figure 5: PLS-DA signature from the 'sacurine' data set.

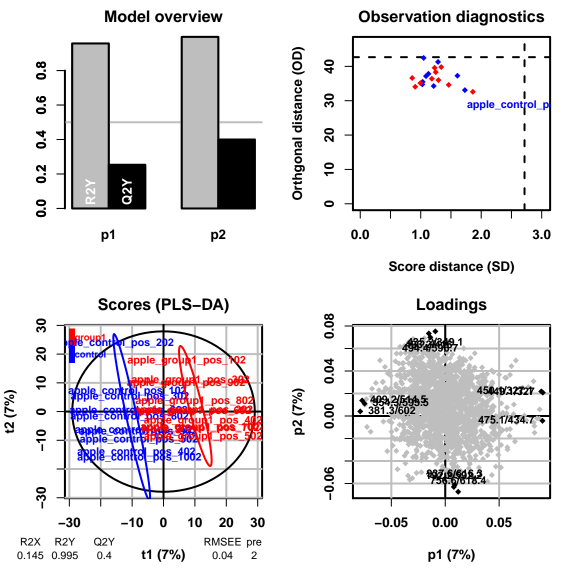
```
biomark.pca <- opls(appleMN, plotL = FALSE)
plot(biomark.pca, parAsColFcVn = spikeFc)
## PCA
## 20 samples x 1632 variables
## standard scaling of predictors
## R2X(cum) pre ort
## Total 0.523 7 0</pre>
```



2. PLS-DA modeling with the full dataset is not significant (as seen on the top left plot: 7 out of 20 models trained after random permutations of the labels have Q2 values higher than the model trained with the true labels):

```
biomark.pls <- opls(appleMN, spikeFc)</pre>
```

##	PLS-D	A							
##	20 sar	nples x 10	532 varial	bles and	1 respo	onse			
##	<pre>## standard scaling of predictors and response(s)</pre>								
##		R2X(cum)	R2Y(cum)	Q2(cum)	RMSEE	pre	ort	pR2Y	pQ2
##	Total	0.145	0.995	0.4	0.0396	2	0	0.15	0.5



Let us now extract the molecular signatures:

```
set.seed(123)
appleSign <- biosign(appleMN, spikeFc)
set.seed(NULL)</pre>
```

The 449.1/327 corresponds to the Cyanidin-3-galactoside (absent in the control) and the 475.1/434.7 is probably a potassium adduct of the Phloridzin (80% concentration increase in group1; Franceschi et al. (2012)).

```
annotDF <- SpikePos[["annotation"]]
rownames(annotDF) <- colnames(appleMN)
annotDF[getSignatureLs(appleSign)[["complete"]], c("adduct", "found.in.standards")]</pre>
```

4.3 Bone marrow from acute leukemia patients (transcriptomics)

Samples from 47 patients with acute lymphoblastic leukemia (ALL) and 25 patients with acute myeloid leukemia (AML) have been analyzed using Affymetrix Hgu6800 chips resulting in expression data of 7,129 gene probes (Golub et al. 1999).

The **golub** dataset is available in the golubEsets package from Bioconductor. Let us compute for example the SVM signature (to speed up this demo example, the number of features is restricted to 500):

```
library(golubEsets)
data(Golub Merge)
golubMN <- t(exprs(Golub Merge))</pre>
leukemiaFc <- pData(Golub_Merge)[["ALL.AML"]]</pre>
table(leukemiaFc)
varSubVi <- 1501:2000
set.seed(123)
golubSign <- biosign(golubMN[, varSubVi], leukemiaFc, methodVc = "svm")</pre>
set.seed(NULL)
## leukemiaFc
## ALL AML
## 47 25
## Significant features from 'S' groups:
##
              svm
## M19507_at "S"
## M27891_at "S"
## Accuracy:
##
           svm
## Full 0.950
## AS
        0.969
## S
        0.944
```

The computation results in a signature of 2 features only and a sparse SVM model performing almost as well (94.4% accuracy) as the model trained on the dataset of 500 variables (95.0% accuracy).

The hu6800.db bioconductor package can be used to get the annotation of the selected probes (Carlson 2016):

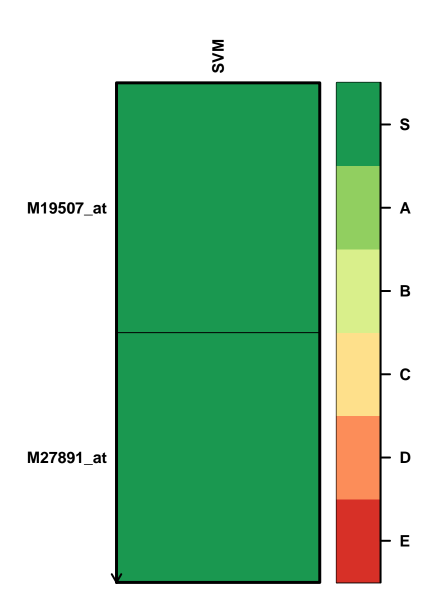
```
library(hu6800.db)
sapply(getSignatureLs(golubSign)[["complete"]],
    function(probeC)
    get(probeC, env = hu6800GENENAME))
## M19507_at M27891_at
## "myeloperoxidase" "cystatin C"
```

Cystatin C is part of the 50 gene signature selected by Golub and colleagues on the basis of a metric derived from the Student's statistic of mean differences between the AML and ALL groups (Golub et al. 1999). Interestingly, the second probe, myeloperoxidase, is a cytochemical marker for the diagnosis (and also potentially the prognosis) of acute myeloid leukemia (AML).

5 Session info

Here is the output of sessionInfo() on the system on which this document was compiled:

```
## R version 3.4.0 (2017-04-21)
## Platform: x86_64-pc-linux-gnu (64-bit)
## Running under: Ubuntu 16.04.2 LTS
##
## Matrix products: default
## BLAS: /home/biocbuild/bbs-3.5-bioc/R/lib/libRblas.so
## LAPACK: /home/biocbuild/bbs-3.5-bioc/R/lib/libRlapack.so
##
```





```
## locale:
## [1] LC_CTYPE=en_US.UTF-8
                                  LC_NUMERIC=C
## [3] LC_TIME=en_US.UTF-8
                                  LC_COLLATE=C
##
   [5] LC_MONETARY=en_US.UTF-8
                                  LC_MESSAGES=en_US.UTF-8
   [7] LC_PAPER=en_US.UTF-8
##
                                  LC_NAME=C
##
  [9] LC_ADDRESS=C
                                  LC_TELEPHONE=C
## [11] LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C
##
```

```
## attached base packages:
## [1] stats4
                 parallel stats
                                     graphics grDevices utils
                                                                    datasets
## [8] methods
                 base
##
## other attached packages:
   [1] hu6800.db_3.2.3
##
                             org.Hs.eg.db_3.4.1
                                                   AnnotationDbi_1.38.0
##
    [4] IRanges_2.10.0
                             S4Vectors_0.14.0
                                                   golubEsets_1.17.0
##
  [7] BioMark_0.4.5
                             st_1.2.5
                                                   sda_1.3.7
## [10] fdrtool_1.2.15
                             corpcor_1.6.9
                                                   entropy_1.2.1
## [13] MASS_7.3-47
                             glmnet_2.0-5
                                                   foreach_1.4.3
## [16] Matrix 1.2-9
                             pls 2.6-0
                                                   Biobase 2.36.0
## [19] BiocGenerics_0.22.0 ropls_1.8.0
                                                  biosigner_1.4.0
## [22] BiocStyle_2.4.0
##
## loaded via a namespace (and not attached):
## [1] Rcpp_0.12.10
                            compiler 3.4.0
                                                 class 7.3-14
## [4] iterators 1.0.8
                            tools_3.4.0
                                                 digest_0.6.12
## [7] memoise_1.1.0
                            RSQLite_1.1-2
                                                 evaluate_0.10
## [10] lattice_0.20-35
                            DBI_0.6-1
                                                 yaml_2.1.14
## [13] e1071_1.6-8
                            stringr_1.2.0
                                                knitr_1.15.1
## [16] rprojroot_1.2
                            grid_3.4.0
                                                 rmarkdown_1.4
## [19] magrittr_1.5
                            backports_1.0.5
                                                 codetools_0.2-15
## [22] htmltools_0.3.5
                            randomForest_4.6-12 stringi_1.1.5
```

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