

HOW TO use MCReestimate

Markus Ruschhaupt

March 31, 2012

Estimation the misclassification error

Every classification task starts with data. Here we choose the well known ALL/AML data set from T.Golub [1] available in the package *golubEsets*. Furthermore, we specify the name of the phenodata column that should be used for the classification.

```
> library(MCReestimate)
> library(randomForest)
> library(golubEsets)
> data(Golub_Train)
> class.column <- "ALL.AML"
```

Cross-validation is an appropriate instrument to estimate the misclassification rate of an algorithm that should be used for classification. Often a variable selection or aggregation is applied to the data set before the main classification procedure is started. But these steps must also be part of the cross-validation. In our example we want to perform a variable selection and only take the genes with the highest variance across all samples. Because we don't know exactly how many genes we want to have, we give two possible values (250 and 1000). The described methods is implemented in the functions `g.red.highest.var`. Further preprocessing functions are part of the package *MCReestimate* and also new functions can be implemented.

```
> Preprocessingfunctions      <- c("varSel.highest.var")
> list.of.poss.parameter      <- list(var.numbers=c(250,1000))
```

To use *MCReestimate* with a classification procedure a wrapper for this method must be available. The package *MCReestimate* includes wrapper for the following classification functions.

RF.wrap wrapper for random forest (based on the package *randomForest*)

PAM.wrap wrapper for PAM (based on the package *pamr*)

PLR.wrap wrapper for the penalised logistic regression (based on the package *MCRestimate*)

SVM.wrap wrapper for support vector machines (based on the package *e1071*)

GPLS.wrap wrapper for generalised partial least squares (based on the package *gpls*)

It is easy to write a wrapper for a new classification method. Here we want to use random forest for our classification task.

```
> class.function <- "RF.wrap"
```

Parameter to optimize: Most classification and preprocessing methods have parameters that must be chosen and/or optimized. In our example we choose two possible number of clusters for our preprocessing method. In each cross-validation step *MCRestimate* will choose the value that have the lowest misclassification error on the training set. This will be estimated through a second (inner) cross-validation with the function *tune* available in the package *e1071*.

Parameter for the plot of the results (see man page for details): We want to have the sample names as labels in the resulting plot. *Samples* is the name of the *phenoData* column the sample names are stored in.

```
> plot.label <- "Samples"
```

The Cross-validation parameter (see man page for details): For time reasons here we choose very small values. Normally the values for *cross.outer* and *cross.inner* should be at least 5.

```
> cross.outer <- 2
> cross.repeat <- 3
> cross.inner <- 2
```

Now we have specified all parameter and can run the function

```
> RF.estimate <- MCRestimate(Golub_Train,
+                             class.colum,
+                             classification.fun="RF.wrap",
+                             thePreprocessingMethods=Preprocessingfunctions,
+                             poss.parameters=list.of.poss.parameter,
+                             cross.outer=cross.outer,
+                             cross.inner=cross.inner,
+                             cross.repeat=cross.repeat,
+                             plot.label=plot.label)
```

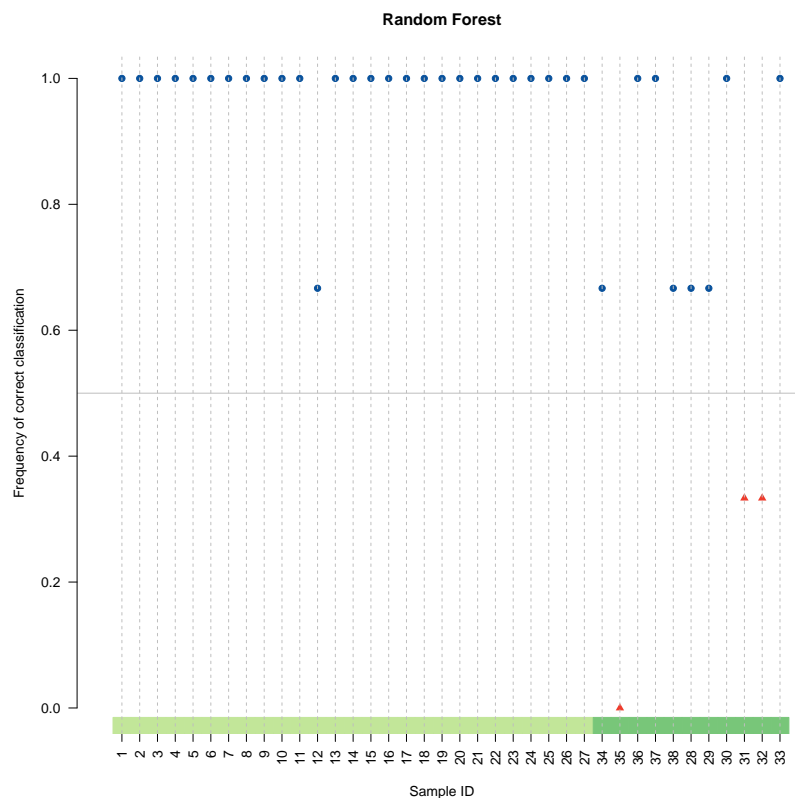
The result is an element of class *MCRestimate*

```
> class(RF.estimate)
```

```
[1] "MCReestimate"
```

For each group we see how many samples have been misclassified most of the time. We can also visualize the result. The plot shows for each sample the number of times it has been classified correctly.

```
> plot(RF.estimate,rownames.from.object=TRUE, main="Random Forest")
```



New data

We have estimated the misclassification rate of our complete algorithm. Because this seems to be a quit good result, we want to use this algorithm to classify new data. The function `ClassifierBuild` is used to build a classifier that can be used for new data.

```
> RF.classifier <- ClassifierBuild (Golub_Train,
+                                 class.colum,
+                                 classification.fun="RF.wrap",
+                                 thePreprocessingMethods=Preprocessingfunctions,
+                                 poss.parameters=list.of.poss.parameter,
+                                 cross.inner=cross.inner)
```

The result of the function is a list with various arguments.

```
> names(RF.classifier)
[1] "classifier.for.matrix"
[2] "classifier.for.exprSet"
[3] "classes"
[4] "parameter"
[5] "thePreprocessingMethods"
[6] "class.method"
[7] "cross.inner"
[8] "information"
```

The most important elements of this list are *classifier.for.matrix* and *classifier.for.exprSet*. These can now be used to classify new data. The first one can be used if the new data is given by a matrix and the second will be used if the new data is given by a *exprSet*. Here we want to classify the data from the *exprSet* *golubTest*, that is also part of the package *golubEsets*.

```
> data(Golub_Test)
> RF.classifier$classifier.for.exprSet(Golub_Test)

 39  40  42  47  48  49  41  43  44  45  46  70
ALL ALL ALL ALL ALL ALL ALL ALL ALL ALL ALL ALL
 71  72  68  69  67  55  56  59  52  53  51  50
ALL ALL ALL ALL ALL ALL ALL ALL AML AML AML AML
 54  57  58  60  61  65  66  63  64  62
ALL AML AML ALL ALL AML ALL AML AML AML
Levels: ALL AML
```

This work was supported by NGFN (Nationales Genomforschungsnetz).

References

- [1] Golub TR, Slonim DK, Tamayo P, Huard C, Gaasenbeek M, Mesirov JP, Coller H, Loh ML, Downing JR, Caligiuri MA, Bloomfield CD, Lander ES. Molecular classification of cancer: class discovery and class prediction by gene expression monitoring *Science* 286(5439): 531-7 (1999).