

# Package ‘SVM2CRM’

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**Type** Package

**Title** SVM2CRM: support vector machine for cis-regulatory elements detections

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**Description** Detection of cis-regulatory elements using svm implemented in LiblinearR.

**Author** Guidantonio Malagoli Tagliazucchi

<guidantonio.malagolitagliazucchi@unimore.it>

**License** GPL-3

**Depends** R (>= 3.2.0), LiblineaR, SVM2CRMdata

**Imports** AnnotationDbi, mclust, GenomicRanges, IRanges, zoo, squash, pls,rtracklayer,ROCR,verification

**biocViews** ChIPSeq, SupportVectorMachine, Software, Preprocessing, ChipOnChip

**Maintainer** Guidantonio Malagoli Tagliazucchi

<guidantonio.malagolitagliazucchi@unimore.it>

**NeedsCompilation** no

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**cisREfindbed***Create and preprocessing of the input for SVM2CRM analysis*

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**Description**

This function require as input a bed file in format chr, start, end, signalNorm. The genomic coordinates must be sort. The data should be normalized using others methods. For e.g the ChIP-seq data should be partitioned in 100bp and then normalize by the library size. This function simply load each bed file of one histone mark and then, partition the genome in n not overlapping windows of size (w). In particular, build a matrix where the number of columns for each histone mark depends on the size of the window and the bin size used during the preprocessing of the bed file. This function allow to smooth the signal of the bed file every n bin. The default function of smoothing is median. This is the suggested function to model the signal. Other function have not been tested.

**Usage**

```
cisREfindbed(list_file,chr="chr1", bin.size, windows, window.smooth=200,smoothing="FALSE",function.
```

**Arguments**

<code>list_file</code>	The list of bed files (e.g. character)
<code>chr</code>	A vector containing the list of chromosomes to consider during the analysis
<code>bin.size</code>	The size of bin used to preprocessed the bed files
<code>windows</code>	The size of the window (default=5000bp)
<code>window.smooth</code>	The size of the window to smooth the bin (default=200)
<code>smoothing</code>	logical, default is FALSE, if TRUE smooth the signal
<code>function.smoothing</code>	Set the function to smooth the signal (default is median)

**Details**

Some detailed description

**Value**

A data.frame where the number of columns depends on the windows size, bin.size and the number of histone marks considered for the prediction.

**Author(s)**

Guidantonio Malagoli Tagliazucchi `guidantonio.malagolitagliazucchi@unimore.it`

**See Also**

cisREfindbed, mclapply

**Examples**

```
library(rtracklayer)
chr<- "chr1"
#the values use to binarized the genome (in bp)
bin.size<-100
#the size of the windows (in bp)
windows<-500
#the step of smoothing (in bp): e.g. smooth the signal every 200bp
window.smooth<-200
#do you want apply a smoothing
smoothing<-"FALSE"
#what kind of function do you want use to smooth the signal
function.smoothing<-"median"
#list of file: format bed
list_file<-"CD4-H3K14ac.norm.w100.bed"

#completeTABLE<-cisREfindbed(list_file[1],chr=chr,bin.size=bin.size,windows=windows,window.smooth=window.smooth)
```

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createBed

*Create bed file of predictions using svm.*

---

**Description**

With this function the user can create a bed files with the regions of predicted cis-regulatory elements. Internal function.

**Usage**

```
createBed(test_set,label1,pred,outputfile)
```

**Arguments**

test_set	test_set produced for svm model
label1	class1
pred	p\$prediction object
outputfile	name of bed file

**Details**

Some detailed description

**Value**

A bed files with the genomic coordinates of the cis-regulatory elements predicted using SVM2CRM.

**Author(s)**

Guidantonio Malagoli Tagliazucchi guidantonio.malagolitagliazucchi@unimore.it

**See Also**

`cisREfindbed`

**Examples**

```

library("GenomicRanges")
library("SVM2CRMdata")

setwd(system.file("data", package="SVM2CRMdata"))
load("CD4_matrixInputSVMbin100window1000.rda")
completeTABLE<-CD4_matrixInputSVMbin100window1000

new.strings<-gsub(x=colnames(completeTABLE[,c(6:ncol(completeTABLE))]), pattern="CD4.", replacement="")
new.strings<-gsub(new.strings, pattern=".norm.w100.bed", replacement="")
colnames(completeTABLE)[c(6:ncol(completeTABLE))]<-new.strings

#list_file<-grep(dir(), pattern=".sort.txt", value=TRUE)

#train_positive<-getSignal(list_file, chr="chr1", reference="p300.distal.fromTSS.txt", win.size=500, bin.size=1)
#train_negative<-getSignal(list_file, chr="chr1", reference="random.region.hg18.nop300.txt", win.size=500, bin.size=1)
setwd(system.file("data", package="SVM2CRMdata"))
load("train_positive.rda")
load("train_negative.rda")

training_set<-rbind(train_positive, train_negative)
#the colnames of the training set should be the same of data_enhancer_svm
colnames(training_set)[c(5:ncol(training_set))]<-gsub(x=gsub(x=colnames(training_set[,c(5:ncol(training_set))]), pattern="CD4.", replacement=""), new.strings, ignore.case=TRUE)

setwd(system.file("extdata", package = "SVM2CRMdata"))
data_level2 <- read.table(file = "GSM393946.distal.p300fromTSS.txt", sep = "\t", stringsAsFactors = FALSE)
data_level2<-data_level2[data_level2[,1]=="chr1",]

DB <- data_level2[, c(1:3)]
colnames(DB)<-c("chromosome", "start", "end")

label <- "p300"

table.final.overlap<-findFeatureOverlap(query=completeTABLE, subject=DB, select="all")

data_enhancer_svm<-createSVMinput(inputpos=table.final.overlap, inputfull=completeTABLE, label1="enhancers", 1)
colnames(data_enhancer_svm)[c(5:ncol(data_enhancer_svm))]<-gsub(gsub(x=colnames(data_enhancer_svm[,c(5:ncol(data_enhancer_svm))]), pattern="CD4.", replacement=""), new.strings, ignore.case=TRUE)

```

```

listcolnames<-c("H2AK5ac","H2AK9ac","H3K23ac","H3K27ac","H3K27me3","H3K4me1","H3K4me3")

dftotann<-smoothInputFS(train_positive[,c(6:ncol(train_positive))],listcolnames,k=20)

results<-featSelectionWithKmeans(dftotann,5)

resultsFS<-results[[7]]

resultsFSfilter<-resultsFS[which(resultsFS[,2]>median(resultsFS[,2])),]

resultsFSfilterICRR<-resultsFSfilter[which(resultsFSfilter[,3]<0.50),]

listHM<-resultsFSfilterICRR[,1]
listHM<-gsub(gsub(listHM,pattern=". ",replacement=""),pattern="CD4.",replacement="")

selectFeature<-grep(x=colnames(training_set[,c(6:ncol(training_set))]),pattern=paste(listHM,collapse="|"),value=TRUE)

colSelect<-c("chromosome","start","end","label",selectFeature)
training_set<-training_set[,colSelect]

vecS <- c(2:length(listHM))
typeSVM <- c(0, 6, 7)[1]
costV <- c(0.001, 0.01, 0.1, 1, 10, 100, 1000)[6]
wlabel <- c("not_enhancer", "enhancer")
infofile<-data.frame(a=c(paste(listHM,"signal",sep=". ")))
infofile[,1]<-gsub(gsub(x=infofile[,1],pattern="CD4.",replacement=""),pattern=".sort.bed",replacement="")

tuningTAB <- tuningParametersCombROC(training_set = training_set, typeSVM = typeSVM, costV = costV,differentiation="enhancer")

tuningTABfilter<-tuningTAB[tuningTAB$fscore<0.95,]
#row_max_fscore<-which.max(tuningTABfilter[tuningTABfilter$nHM >2,"fscore"])
row_max_fscore<-which.max(tuningTABfilter[,"fscore"])
listHM_prediction<-gsub(tuningTABfilter[row_max_fscore,4],pattern="//",replacement="|")

columnPR<-grep(colnames(training_set),pattern=paste(listHM_prediction,collapse="|"),value=TRUE)

predictionGW(training_set=training_set,data_enhancer_svm=data_enhancer_svm, listHM=columnPR,pcClass.string="enhancer")

```

## createSVMinput

*Take the output of findFeatureOverlap and then create a positive and negative set of cis-regulatory elements*

**Description**

Take the output of findFeatureOverlap and then create a positive and negative set of cis-regulatory elements

**Usage**

```
createSVMinput(inputpos, inputfull, label1, label2)
```

**Arguments**

inputpos	output of findFeatureOverlap (see documentation findFeatureOverlap)
inputfull	output of cisREfindbed (see documentation cisREfinbed)
label1	a string to define the first class (e.g. enhancers)
label2	a string to define the second class (e.g. not_enhancers)

**Details**

Some detailed description

**Value**

A data.frame with the signals of the histone modifications for positive (e.g. enhancers) and negative (e.g. not\_enhancers) examples.

**Author(s)**

Guidantonio Malagoli Tagliazucchi guidantonio.malagolitagliazucchi@unimore.it

**See Also**

[findFeatureOverlap](#), [cisREfindbed](#)

**Examples**

```
library("SVM2CRMdata")
library("GenomicRanges")

setwd(system.file("data", package="SVM2CRMdata"))
load("CD4_matrixInputSVMbin100window1000.rda")
completeTABLE<-CD4_matrixInputSVMbin100window1000

new.strings<-gsub(x=colnames(completeTABLE[,c(6:ncol(completeTABLE))]), pattern="CD4.", replacement="")
new.strings<-gsub(new.strings, pattern=".norm.w100.bed", replacement="")
colnames(completeTABLE)[c(6:ncol(completeTABLE))]<-new.strings

#list_file<-grep(dir(), pattern=".sort.txt", value=TRUE)
#train_positive<-getSignal(list_file, chr="chr1", reference="p300.distal.fromTSS.txt", win.size=500, bin.size=1)
#train_negative<-getSignal(list_file, chr="chr1", reference="random.region.hg18.nop300.txt", win.size=500, bin.size=1)
#setwd(system.file("data", package="SVM2CRMdata"))
#load("train_positive.rda")
#load("train_negative.rda")

training_set<-rbind(train_positive, train_negative)
colnames(training_set)[c(5:ncol(training_set))]<-gsub(x=colnames(training_set[,c(5:ncol(training_set))]), pattern="CD4.", replacement="")
```

```

setwd(system.file("extdata", package="SVM2CRMdata"))
data_level2 <- read.table(file = "GSM393946.distal.p300fromTSS.txt", sep = "\t", stringsAsFactors = FALSE)
data_level2<-data_level2[data_level2[,1]=="chr1",]

DB <- data_level2[, c(1:3)]
colnames(DB)<-c("chromosome", "start", "end")

label <- "p300"

table.final.overlap<-findFeatureOverlap(query=completeTABLE, subject=DB)
data_enhancer_svm<-createSVMinput(inputpos=table.final.overlap, inputfull=completeTABLE, label1="enhancers", l

```

**featSelectionWithKmeans**

*This function select the most meaningful variables in a matrix of ChIP-seq data using k-means and ICRR.*

**Description**

In the research of promoters and enhancers the users start with large dataset of histone modification. The number of histone marks is a variable that influence the prediction of enhancers. Several papers investigated what is the best combination of histone marks to predict enhancers. There are not a consensus about the optimal number of histone mark in the prediction of cis-regulatory elements. Moreover, in the genome there are a number of histone marks that is greater than 50. The biological roles of all of these is not already clear. However from the computational aspect the admit of a huge number of variables (histone marks) can insert redundant information (signal of histone marks) during the step of prediction. Therefore here we introduced a step of features selection. The step of variables selection consist of two steps. In the first one the function perform a smoothing of the signals of each histone mark. For example, if the data were binned every 100bp, a window size of smoothing equal to 2 means that the signal of the histone mark is smoothed every 200 bp. In the second step, the function using a k number of clusters define by the user gather the signal of each histone mark and estimate the mean of signals inside each group. Finally is estimated the median of all means above computed and all histone marks with a signal less than of this are filtered out. We introduced also the concept of index coverage of the regulatory regions (ICRR). In the first step the function measure the total coverage of the cis-regulatory elements in the list of positive class and considering w, the windows size used to model the signal of the histone marks around a particular feature (positive class, negative class). Then it inspects in those enhancers the signals (S) of a particular histone marks is less or greater than the global mean (M) of the signal (S). Next the function compute the coverage only for these two fractions of enhancers and estimate the ICRR. This values assume values from 0 to 1. A value close to 0 means that the difference between the coverage of the two classes of enhancers is little, in contrast, if this value is close to 1 this means that there is a diversity between the two fraction of the cis-regulatory elements. The ICRR values can be used to select the histone marks to use during the analysis. Next using tuningParametersCombROC the user can tune the best optimal set of svm parameters and histone marks. The function featSelectionWithKmeans return a list containing the results of this analysis. The first element contain a matrix with the results of the selection analysis, the parameters used during the analysis and the filtered histone marks. Finally this function create a report with three

plots. In the first one the x-axis contain the labels of the histone marks while in y-axis reports the median of mean of each group. The second plot represent the same thing but using a scatterplot. The third page contain the same graph in the first page and a plot that contain the index of coverage between enhancers and not enhancers regions.

## Usage

```
featSelectionWithKmeans(dftotann, nk, autoK="no", w=1000, gmax=7, outputplot="feature.selection.pdf")
```

## Arguments

dftotann	a data.frame created with smoothInputFS
nk	number of cluster to use for k-means algorithm
autoK	logical, if autoK=yes, the function estimate automatically the number of cluster to use during k-means (autoK="no")
w	the windows size used to build the model (default=1000bp)
gmax	the parameter gmax used during the analysis (gmax=7)
outputplot	the name of the report pdf file (default="feature.selection.pdf")

## Details

input: the results of smoothinputFS. This function clusterized every column (histone marks) using a K number of cluster user defined. Next the mean of the signals in each group are estimated. The function *featSelectionWithKmeans* return a list containing the results of this analysis. The first element contain a matrix with the results of the selection analysis, the parameters used durint the analysis and the filtered histone marks. Finally this function create a report with three plots. In the first one the x-axis contain the labels of the histone marks while in y-axis reports the median of mean of each group. The second plot represent the same thing but using a scatterplot. The third page contain the same graph in the first page and a plot that contain the index of coverage between enhancers and not enhancers regions. Definition of k-clusters: The user can set manually the parameter k otherwise *featSelectionWithKmeans* implement automatically the investigation of k using a Bayesian Information Criterion for EM initialized.

## Value

- a list where the first element is the matrix of output of feature selection analysis - mom: the vertical mean of the matrix in the element 1 - nk: number of clusters used during the clustering - gmax: the parameter gmax used during the analysis - which histone marks have a the signal that is greater than the median global signal - which histone marks have a signal where the histone are greater than the mean of the global signal. - a data.frame where the first column contain the results of feature selection analysis with k-means and in the second column the results of ICRR (index coverage regulatory regions).

## Author(s)

Guidantonio Malagoli Tagliazucchi guidantonio.malagolitagliazucchi@unimore.it

**See Also**

`cisREfindbed`, `smoothInputFS`, `tuningParametersCombROC`

**Examples**

```

library("SVM2CRMdata")
library("GenomicRanges")

setwd(system.file("data", package="SVM2CRMdata"))
load("CD4_matrixInputSVMbin100window1000.rda")
completeTABLE<-CD4_matrixInputSVMbin100window1000

new.strings<-gsub(x=colnames(completeTABLE[,c(6:ncol(completeTABLE))]), pattern="CD4.", replacement="")
new.strings<-gsub(new.strings, pattern=".norm.w100.bed", replacement="")
colnames(completeTABLE)[c(6:ncol(completeTABLE))]<-new.strings

#list_file<-grep(dir(), pattern=".sort.txt", value=TRUE)
#train_positive<-getSignal(list_file, chr="chr1", reference="p300.distal.fromTSS.txt", win.size=500, bin.size=1)
#train_negative<-getSignal(list_file, chr="chr1", reference="random.region.hg18.nop300.txt", win.size=500, bin.size=1)
#load("train_positive.rda")
#load("train_negative.rda")

training_set<-rbind(train_positive, train_negative)

training_set<-rbind(train_positive, train_negative)
colnames(training_set)[c(5:ncol(training_set))]<-gsub(x=gsub(x=colnames(training_set[,c(5:ncol(training_set))]), pattern="CD4.", replacement=""), new.strings, ignore.case=TRUE)

setwd(system.file("extdata", package = "SVM2CRMdata"))
data_level2 <- read.table(file = "GSM393946.distal.p300fromTSS.txt", sep = "\t", stringsAsFactors = FALSE)
data_level2<-data_level2[data_level2[,1]=="chr1",]

DB <- data_level2[, c(1:3)]
colnames(DB)<-c("chromosome", "start", "end")

label <- "p300"

table.final.overlap<-findFeatureOverlap(query=completeTABLE, subject=DB, select="all")

data_enhancer_svm<-createSVMinput(inputpos=table.final.overlap, inputfull=completeTABLE, label1="enhancers", label2="p300")
colnames(data_enhancer_svm)[c(5:ncol(data_enhancer_svm))]<-gsub(gsub(x=colnames(data_enhancer_svm[,c(5:ncol(data_enhancer_svm))]), pattern="CD4.", replacement=""), new.strings, ignore.case=TRUE)

listcolnames<-c("H2AK5ac", "H2AK9ac", "H3K23ac", "H3K27ac", "H3K27me3", "H3K4me1", "H3K4me3")

dftotann<-smoothInputFS(train_positive[,c(6:ncol(train_positive))], listcolnames, k=20)

results<-featSelectionWithKmeans(dftotann, 5)

```

---

<b>findFeatureOverlap</b>	<i>Find the overlap of genomic regions between the output of cisREfind and a database of validated cis-regulatory elements.</i>
---------------------------	---

---

## Description

Find the overlap of genomic regions between the output of cisREfind and a database of validated cis-regulatory elements.

## Usage

```
findFeatureOverlap(query, subject, select)
```

## Arguments

query	database of validated cis-regulatory elements (e.g. p300 binding sites), GRanges class.
subject	output cisREfindbed function, GRanges class.
select	parameters of findOverlaps

## Details

See documentation of GenomicRanges for details.

## Value

A data frame with the genomic regions that overlap

## Author(s)

Guidantonio Malagoli Tagliazucchi guidantonio.malagolitagliazucchi@unimore.it

## See Also

cisREfind, findOverlaps, GenomicRanges

## Examples

```
library("SVM2CRMdata")
library("GenomicRanges")

setwd(system.file("data", package="SVM2CRMdata"))
load("CD4_matrixInputSVMbin100window1000.rda")
completeTABLE<-CD4_matrixInputSVMbin100window1000
new.strings<-gsub(x=colnames(completeTABLE[,c(6:ncol(completeTABLE))]), pattern="CD4.", replacement="")
new.strings<-gsub(new.strings, pattern=".norm.w100.bed", replacement="")
colnames(completeTABLE)[c(6:ncol(completeTABLE))]<-new.strings
```

```
setwd(system.file("extdata", package="SVM2CRMdata"))
data_level2 <- read.table(file = "GSM393946.distal.p300fromTSS.txt", sep = "\t", stringsAsFactors = FALSE)
data_level2<-data_level2[data_level2[,1]=="chr1",]

DB <- data_level2[, c(1:3)]
colnames(DB)<-c("chromosome", "start", "end")

label <- "p300"

table.final.overlap<-findFeatureOverlap(query=completeTABLE, subject=DB)
```

---

frequencyHM

*frequencyHM*

---

## Description

Compute the frequency of the histone marks after tuning of parameters (tuningParametersComb)

## Usage

```
frequencyHM(tuningTABfilter)
```

## Arguments

tuningTABfilter

A data.frame from tuningParametersComb

## Details

Some detailed description

## Value

A vector containing the frequencies of the histone marks

## Author(s)

Guidantonio Malagoli Tagliazucchi guidantonio.malagolitagliazucchi@unimore.it

## See Also

cisREfind, mclapply

## Examples

```

library("GenomicRanges")
library("SVM2CRMdata")

setwd(system.file("data", package="SVM2CRMdata"))
load("CD4_matrixInputSVMbin100window1000.rda")
completeTABLE<-CD4_matrixInputSVMbin100window1000

new.strings<-gsub(x=colnames(completeTABLE)[c(6:ncol(completeTABLE))], pattern="CD4.", replacement="")
new.strings<-gsub(new.strings, pattern=".norm.w100.bed", replacement="")
colnames(completeTABLE)[c(6:ncol(completeTABLE))]<-new.strings

#list_file<-grep(dir(), pattern=".sort.txt", value=TRUE)
#train_positive<-getSignal(list_file, chr="chr1", reference="p300.distal.fromTSS.txt", win.size=500, bin.size=100)
#train_negative<-getSignal(list_file, chr="chr1", reference="random.region.hg18.nop300.txt", win.size=500, bin.size=100)
#setwd(system.file("data", package="SVM2CRMdata"))
#load("train_positive.rda")
#load("train_negative.rda")

training_set<-rbind(train_positive, train_negative)
#the colnames of the training set should be the same of data_enhancer_svm
colnames(training_set)[c(5:ncol(training_set))]<-gsub(x=colnames(training_set), c(5:ncol(training_set)))

setwd(system.file("extdata", package = "SVM2CRMdata"))
data_level2 <- read.table(file = "GSM393946.distal.p300fromTSS.txt", sep = "\t", stringsAsFactors = FALSE)
data_level2<-data_level2[data_level2[,1]=="chr1",]

DB <- data_level2[, c(1:3)]
colnames(DB)<-c("chromosome", "start", "end")

label <- "p300"

table.final.overlap<-findFeatureOverlap(query=completeTABLE, subject=DB, select="all")

data_enhancer_svm<-createSVMinput(inputpos=table.final.overlap, inputfull=completeTABLE, label1="enhancers", label2="non_enhancers")
colnames(data_enhancer_svm)[c(5:ncol(data_enhancer_svm))]<-gsub(gsub(x=colnames(data_enhancer_svm), c(5:ncol(data_enhancer_svm)),))

listcolnames<-c("H2AK5ac", "H2AK9ac", "H3K23ac", "H3K27ac", "H3K27me3", "H3K4me1", "H3K4me3")

dftotann<-smoothInputFS(train_positive[,c(6:ncol(train_positive))], listcolnames, k=20)

results<-featSelectionWithKmeans(dftotann, 5)

resultsFS<-results[[7]]

resultsFSfilter<-resultsFS[which(resultsFS[,2]>median(resultsFS[,2])),]
resultsFSfilterICRR<-resultsFSfilter[which(resultsFSfilter[,3]<0.50),]

```

```

listHM<-resultsFSfilterICRR[,1]
listHM<-gsub(gsub(listHM,pattern="_.",replacement=""),pattern="CD4.",replacement="")

selectFeature<-grep(x=colnames(training_set[,c(6:ncol(training_set))]),pattern=paste(listHM,collapse="|"),va

colSelect<-c("chromosome","start","end","label",selectFeature)
training_set<-training_set[,colSelect]

vecS <- c(2:length(listHM))
typeSVM <- c(0, 6, 7)[1]
costV <- c(0.001, 0.01, 0.1, 1, 10, 100, 1000)[6]
wlabel <- c("not_enhancer", "enhancer")
infofile<-data.frame(a=c(paste(listHM,"signal",sep=".")))
infofile[,1]<-gsub(gsub(x=infofile[,1],pattern="CD4.",replacement=""),pattern=".sort.bed",replacement="")

tuningTAB <- tuningParametersCombROC(training_set = training_set, typeSVM = typeSVM, costV = costV,different.w

tuningTABfilter<-tuningTAB[tuningTAB$fscore<0.95,]

frequencyHM<-frequencyHM(tuningTABfilter)

```

**getSignal**

*Model the signals of each histone marks around genomic features (e.g. enhancers, not\_enhancers).*

**Description**

This function simply model the signal of each histone marks around the features used in the input files and considering the bin.size and window size defined during the pre-processing step.

**Usage**

```
getSignal(bedfilelist,chr,reference,win.size,bin.size,label1="enhancers")
```

**Arguments**

bedfilelist	test_set produced for svm model
chr	a vector containin the list of chromsome that you want use during the analysis (e.g."chr1")
reference	file with the reference position of the features. The genomic coordinates of positive and negative examples (e.g. enhancers, not_enhancers)
win.size	windows size used to smooth the signal
bin.size	original bin size used
label1	class of reference (e.g. enhancers or not_enhancers)

## Details

Some detailed description

## Value

A data.frame with the signals where in the column there are the signals of the histone marks and in the rows the cis-regulatory elements.

## Author(s)

Guidantonio Malagoli Tagliazucchi guidantonio.malagolitagliazucchi@unimore.it

## See Also

cisREfindbed

## Examples

```
library("SVM2CRMdata")
library("GenomicRanges")

setwd(system.file("data",package="SVM2CRMdata"))
load("CD4_matrixInputSVMbin100window1000.rda")
completeTABLE<-CD4_matrixInputSVMbin100window1000

new.strings<-gsub(x=colnames(completeTABLE[,c(6:ncol(completeTABLE))]),pattern="CD4.",replacement="")
new.strings<-gsub(new.strings,pattern=".norm.w100.bed",replacement="")
colnames(completeTABLE)[c(6:ncol(completeTABLE))]<-new.strings

#list_file<-grep(dir(),pattern=".sort.txt",value=TRUE)
#train_positive<-getSignal(list_file,chr="chr1",reference="p300.distal.fromTSS.txt",win.size=500,bin.size=1)
#train_negative<-getSignal(list_file,chr="chr1",reference="random.region.hg18.nop300.txt",win.size=500,bin.size=1)
#setwd(system.file("data",package="SVM2CRMdata"))
#load("train_positive.rda")
#load("train_negative.rda")
#training_set<-rbind(train_positive,train_negative)
#colnames(training_set)[c(5:ncol(training_set))]<-gsub(x=gsub(x=colnames(training_set[,c(5:ncol(training_set))]),pattern="CD4.",replacement=""),y=colnames(training_set[,c(5:ncol(training_set))]),y=colnames(training_set[,c(5:ncol(training_set))]))
```

## Description

This function use a contingency table and then estimate: sensitivity, false positive rate, accuracy, specificity, precision, positive predicted values, negative predicted values, false discovery rates, f-score. Useful to estimate the performance of a model. Internal function of plotROC.

**Usage**

```
performanceSVM(res)
```

**Arguments**

res	a confusion matrix
-----	--------------------

**Details**

Some detailed description

**Value**

A data.frame with tpr.sensitivity, fpr, acc, spc.specificity, precision, ppv, npv, fdr, fscore.

**Author(s)**

Guidantonio Malagoli Tagliazucchi guidantonio.malagolitagliazucchi@unimore.it

**See Also**

cisREfind, tuningParametersCOmbROC, predictionGW

**Examples**

```
library("GenomicRanges")
library("SVM2CRMdata")

setwd(system.file("data", package="SVM2CRMdata"))
load("CD4_matrixInputSVMbin100window1000.rda")
completeTABLE<-CD4_matrixInputSVMbin100window1000

new.strings<-gsub(x=colnames(completeTABLE[,c(6:ncol(completeTABLE))]), pattern="CD4.", replacement="")
new.strings<-gsub(new.strings, pattern=".norm.w100.bed", replacement="")
colnames(completeTABLE)[c(6:ncol(completeTABLE))]<-new.strings

#list_file<-grep(dir(), pattern=".sort.txt", value=TRUE)

#train_positive<-getSignal(list_file, chr="chr1", reference="p300.distal.fromTSS.txt", win.size=500, bin.size=1
#train_negative<-getSignal(list_file, chr="chr1", reference="random.region.hg18.nop300.txt", win.size=500, bin.
setwd(system.file("data", package="SVM2CRMdata"))
load("train_positive.rda")
load("train_negative.rda")

training_set<-rbind(train_positive, train_negative)
#the colnames of the training set should be the same of data_enhancer_svm
colnames(training_set)[c(5:ncol(training_set))]<-gsub(x=gsub(x=colnames(training_set[,c(5:ncol(training_set))]), pattern="CD4.", replacement=""), new.strings, ignore.case=TRUE)

setwd(system.file("extdata", package = "SVM2CRMdata"))
```

```

data_level2 <- read.table(file = "GSM393946.distal.p300fromTSS.txt", sep = "\t", stringsAsFactors = FALSE)
data_level2<-data_level2[data_level2[,1]=="chr1",]

DB <- data_level2[, c(1:3)]
colnames(DB)<-c("chromosome", "start", "end")

label <- "p300"

table.final.overlap<-findFeatureOverlap(query=completeTABLE,subject=DB,select="all")

data_enhancer_svm<-createSVMinput(inputpos=table.final.overlap,inputfull=completeTABLE,label1="enhancers",1
colnames(data_enhancer_svm)[c(5:ncol(data_enhancer_svm))]<-gsub(gsub(x=colnames(data_enhancer_svm[,c(5:ncol

listcolnames<-c("H2AK5ac", "H2AK9ac", "H3K23ac", "H3K27ac", "H3K27me3", "H3K4me1", "H3K4me3")

dftotann<-smoothInputFS(train_positive[,c(6:ncol(train_positive))],listcolnames,k=20)

results<-featSelectionWithKmeans(dftotann,5)

resultsFS<-results[[7]]

resultsFSfilter<-resultsFS[which(resultsFS[,2]>median(resultsFS[,2])),]

resultsFSfilterICRR<-resultsFSfilter[which(resultsFSfilter[,3]<0.50),]

listHM<-resultsFSfilterICRR[,1]
listHM<-gsub(gsub(listHM,pattern="_.",replacement=""),pattern="CD4.",replacement="")

selectFeature<-grep(x=colnames(training_set[,c(6:ncol(training_set))]),pattern=paste(listHM,collapse="|"),v

colSelect<-c("chromosome", "start", "end", "label", selectFeature)
training_set<-training_set[,colSelect]

vecS <- c(2:length(listHM))
typeSVM <- c(0, 6, 7)[1]
costV <- c(0.001, 0.01, 0.1, 1, 10, 100, 1000)[6]
wlabel <- c("not_enhancer", "enhancer")
infofile<-data.frame(a=c(paste(listHM,"signal",sep=". ")))
infofile[,1]<-gsub(gsub(x=infofile[,1],pattern="CD4.",replacement=""),pattern=".sort.bed",replacement="")

tuningTAB <- tuningParametersCombROC(training_set = training_set, typeSVM = typeSVM, costV = costV,different.

```

## Description

Plot the ROC curve of the best model using cross-fold validation

**Usage**

```
plotFscore(tuningTAB)
```

**Arguments**

tuningTAB	the output of tuningParametersComb.R
-----------	--------------------------------------

**Details**

See documentation tuningParametersCombROC, perfomanceSVM

**Value**

A pdf with the k-cross-correlation plots for: 1) k-cross-validation, 2) ROC horizontal 3) ROC vertical

**Author(s)**

Guidantonio Malagoli Tagliazucchi guidantonio.malagolitagliazucchi@unimore.it

**See Also**

cisREfindbed, tuningParametersCombROC, perfomanceSVM

**Examples**

```
library("GenomicRanges")
library("SVM2CRMdata")

setwd(system.file("data", package="SVM2CRMdata"))
load("CD4_matrixInputSVMbin100window1000.rda")
completeTABLE<-CD4_matrixInputSVMbin100window1000

new.strings<-gsub(x=colnames(completeTABLE[,c(6:ncol(completeTABLE))]), pattern="CD4.", replacement="")
new.strings<-gsub(new.strings, pattern=".norm.w100.bed", replacement="")
colnames(completeTABLE)[c(6:ncol(completeTABLE))]<-new.strings

list_file<-grep(dir(), pattern=".sort.txt", value=TRUE)

#train_positive<-getSignal(list_file, chr="chr1", reference="p300.distal.fromTSS.txt", win.size=500, bin.size=1
#train_negative<-getSignal(list_file, chr="chr1", reference="random.region.hg18.nop300.txt", win.size=500, bin.size=1
setwd(system.file("data", package="SVM2CRMdata"))
load("train_positive.rda")
load("train_negative.rda")
training_set<-rbind(train_positive, train_negative)
#the colnames of the training set should be the same of data_enhancer_svm
colnames(training_set)[c(5:ncol(training_set))]<-gsub(x=gsub(x=colnames(training_set[,c(5:ncol(training_set))]), pattern="CD4.", replacement=""), new.strings, ignore.case=TRUE)

setwd(system.file("extdata", package = "SVM2CRMdata"))
data_level2 <- read.table(file = "GSM393946.distal.p300fromTSS.txt", sep = "\t", stringsAsFactors = FALSE)
```

```

data_level2<-data_level2[data_level2[,1]=="chr1",]

DB <- data_level2[, c(1:3)]
colnames(DB)<-c("chromosome", "start", "end")

label <- "p300"

table.final.overlap<-findFeatureOverlap(query=completeTABLE,subject=DB,select="all")

data_enhancer_svm<-createSVMinput(inputpos=table.final.overlap,inputfull=completeTABLE,label1="enhancers",1
colnames(data_enhancer_svm)[c(5:ncol(data_enhancer_svm))]<-gsub(gsub(x=colnames(data_enhancer_svm[,c(5:ncol

listcolnames<-c("H2AK5ac", "H2AK9ac", "H3K23ac", "H3K27ac", "H3K27me3", "H3K4me1", "H3K4me3")

dftotann<-smoothInputFS(train_positive[,c(6:ncol(train_positive))],listcolnames,k=20)

results<-featSelectionWithKmeans(dftotann,5)

resultsFS<-results[[7]]

resultsFSfilter<-resultsFS[which(resultsFS[,2]>median(resultsFS[,2])),]

resultsFSfilterICRR<-resultsFSfilter[which(resultsFSfilter[,3]<0.50),]

listHM<-resultsFSfilterICRR[,1]
listHM<-gsub(gsub(listHM,pattern="_.",replacement=""),pattern="CD4.",replacement="")

selectFeature<-grep(x=colnames(training_set[,c(6:ncol(training_set))]),pattern=paste(listHM,collapse="|"),v

colSelect<-c("chromosome", "start", "end", "label", selectFeature)
training_set<-training_set[,colSelect]

vecS <- c(2:length(listHM))
typeSVM <- c(0, 6, 7)[1]
costV <- c(0.001, 0.01, 0.1, 1, 10, 100, 1000)[6]
wlabel <- c("not_enhancer", "enhancer")
infofile<-data.frame(a=c(paste(listHM,"signal",sep=". ")))
infofile[,1]<-gsub(gsub(x=infofile[,1],pattern="CD4.",replacement=""),pattern=".sort.bed",replacement="")

tuningTAB <- tuningParametersCombROC(training_set = training_set, typeSVM = typeSVM, costV = costV,different.

pdf("FSCORE_distribution.pdf")
plotFscore(tuningTAB)
dev.off()

```

## Description

Plot the ROC curve of the best model using cross-fold validation. Internal function of tuningParametersCombROC and predictionGW.

## Usage

```
plotROC(datatrain,k,y,different.weight,type,cost,output)
```

## Arguments

datatrain	training set (data.frame)
y	a vector with the list of labels
k	number of iteration for k-fold cross validation default values is 5.
output	name of output file.
different.weight	specify if the classes are unbalanced.TRUE
type	type of kernel to use
cost	parameter cost of SVM

## Details

Some detailed description

## Value

A pdf with the k-cross-correlation plots for: 1) k-cross-validation, 2) ROC horizontal 3) ROC vertical

## Author(s)

Guidantonio Malagoli Tagliazucchi guidantonio.malagolitagliazucchi@unimore.it

## See Also

cisREfindbed, tuningParametersCombROC, perfTuningParametersCombROC, performanceSVM

## Examples

```
library("GenomicRanges")
library("SVM2CRMdata")

setwd(system.file("data", package="SVM2CRMdata"))
load("CD4_matrixInputSVMbin100window1000.rda")
completeTABLE<-CD4_matrixInputSVMbin100window1000

new.strings<-gsub(x=colnames(completeTABLE[,c(6:ncol(completeTABLE))]), pattern="CD4.", replacement="")
new.strings<-gsub(new.strings, pattern=".norm.w100.bed", replacement="")
colnames(completeTABLE)[c(6:ncol(completeTABLE))]<-new.strings
```

```

#list_file<-grep(dir(),pattern=".sort.txt",value=TRUE)

#train_positive<-getSignal(list_file,chr="chr1",reference="p300.distal.fromTSS.txt",win.size=500,bin.size=1)
#train_negative<-getSignal(list_file,chr="chr1",reference="random.region.hg18.nop300.txt",win.size=500,bin.size=1)
  setwd(system.file("data",package="SVM2CRMdata"))
  load("train_positive.rda")
  load("train_negative.rda")

  training_set<-rbind(train_positive,train_negative)
  #the colnames of the training set should be the same of data_enhancer_svm
  colnames(training_set)[c(5:ncol(training_set))]<-gsub(x=gsub(x=colnames(training_set[,c(5:ncol(training_set))]),y=colnames(DB),fixed=TRUE),y=colnames(DB),fixed=TRUE)

  setwd(system.file("extdata", package = "SVM2CRMdata"))
  data_level2 <- read.table(file = "GSM393946.distal.p300fromTSS.txt",sep = "\t", stringsAsFactors = FALSE)
  data_level2<-data_level2[data_level2[,1]=="chr1",]

  DB <- data_level2[, c(1:3)]
  colnames(DB)<-c("chromosome","start","end")

  label <- "p300"

  table.final.overlap<-findFeatureOverlap(query=completeTABLE,subject=DB,select="all")

  data_enhancer_svm<-createSVMinput(inputpos=table.final.overlap,inputfull=completeTABLE,label1="enhancers",label2="not_enhancers")
  colnames(data_enhancer_svm)[c(5:ncol(data_enhancer_svm))]<-gsub(gsub(x=colnames(data_enhancer_svm[,c(5:ncol(data_enhancer_svm))]),y=colnames(DB),fixed=TRUE),y=colnames(DB),fixed=TRUE)

  listcolnames<-c("H2AK5ac","H2AK9ac","H3K23ac","H3K27ac","H3K27me3","H3K4me1","H3K4me3")

  dftotann<-smoothInputFS(train_positive[,c(6:ncol(train_positive))],listcolnames,k=20)

  results<-featSelectionWithKmeans(dftotann,5)
  resultsFS<-results[[7]]

  resultsFSfilter<-resultsFS[which(resultsFS[,2]>median(resultsFS[,2])),]
  resultsFSfilterICRR<-resultsFSfilter[which(resultsFSfilter[,3]<0.50),]

  listHM<-resultsFSfilterICRR[,1]
  listHM<-gsub(gsub(listHM,pattern="_.",replacement=""),pattern="CD4.",replacement="")

  selectFeature<-grep(x=colnames(training_set[,c(6:ncol(training_set))]),pattern=paste(listHM,collapse="|"))
  colSelect<-c("chromosome","start","end","label",selectFeature)
  training_set<-training_set[,colSelect]

  vecS <- c(2:length(listHM))
  typeSVM <- c(0, 6, 7)[1]
  costV <- c(0.001, 0.01, 0.1, 1, 10, 100, 1000)[6]
  wlabel <- c("not_enhancer", "enhancer")
  infofile<-data.frame(a=c(paste(listHM,"signal",sep=".")))

```

```

infofile[,1]<-gsub(gsub(x=infofile[,1],pattern="CD4.",replacement=""),pattern=".sort.bed",replacement="")

tuningTAB <- tuningParametersCombROC(training_set = training_set, typeSVM = typeSVM, costV = costV,differenceV=0.01)

tuningTABfilter<-tuningTAB[tuningTAB$fscore<0.95,]
#row_max_fscores<-which.max(tuningTABfilter[tuningTABfilter$nHM >2,"fscore"])
row_max_fscores<-which.max(tuningTABfilter[,"fscore"])
listHM_prediction<-gsub(tuningTABfilter[row_max_fscores,4],pattern="//",replacement="|")

columnPR<-grep(colnames(training_set),pattern=paste(listHM_prediction,collapse="|"),value=TRUE)

predictionGW(training_set=training_set,data_enhancer_svm=data_enhancer_svm, listHM=columnPR,pcClass.string="enhancer",nClass.string="not_enhancers",cost=100,type="linear",output.file="enhancers.bed")

```

**`predictionGW`***Perform prediction of cis-regulatory elements genome-wide***Description**

This function perform the prediction genome-wide of the cis-regulatory elements. The function return the output of performanceSVM and a bed file with the position of enhancers and not enhancers regions.

**Usage**

```
predictionGW(training_set,data_enhancer_svm,listHM,pcClass.string="enhancer",nClass.string="not_enhancers",cost=100,type="linear",output.file="enhancers.bed")
```

**Arguments**

<code>training_set</code>	training set (data.frame)
<code>data_enhancer_svm</code>	the signals of all histone marks along the genome
<code>listHM</code>	a vector with the histone marks that you want to use perform the prediction
<code>pcClass.string</code>	label of the first class (e.g. "enhancer")
<code>nClass.string</code>	label of the second class (e.g. "not_enhancers")
<code>pcClass</code>	number of positive class in the test set
<code>ncClass</code>	number of negative class in the test set
<code>cost</code>	parameter of svm (default=100)
<code>type</code>	type of kernel (default=0)
<code>output.file</code>	name of the bed file of output

**Details**

The ratio between the positive and negative regions usually is 1:10. However this ratio depends on your experimental design and your data. See documentation `cisREfindbed`, `tuningParamtersCombROC`, `featSelectionWithKmeans`.

**Value**

The performance of prediction and a bed file with the coordinates of genomic regions that contain the enhancers. The bed file is saved in the directory selected by the user.

**Author(s)**

Guidantonio Malagoli Tagliazucchi guidantonio.malagolitagliazucchi@unimore.it

**See Also**

cisREfindbed, mclapply

**Examples**

```

library("GenomicRanges")
library("SVM2CRMdata")

setwd(system.file("data", package="SVM2CRMdata"))
load("CD4_matrixInputSVMbin100window1000.rda")
completeTABLE<-CD4_matrixInputSVMbin100window1000

new.strings<-gsub(x=colnames(completeTABLE)[c(6:ncol(completeTABLE))], pattern="CD4.", replacement="")
new.strings<-gsub(new.strings, pattern=".norm.w100.bed", replacement="")
colnames(completeTABLE)[c(6:ncol(completeTABLE))]<-new.strings

#list_file<-grep(dir(), pattern=".sort.txt", value=TRUE)

#train_positive<-getSignal(list_file, chr="chr1", reference="p300.distal.fromTSS.txt", win.size=500, bin.size=1)
#train_negative<-getSignal(list_file, chr="chr1", reference="random.region.hg18.nop300.txt", win.size=500, bin.size=1)
setwd(system.file("data", package="SVM2CRMdata"))
load("train_positive.rda")
load("train_negative.rda")

training_set<-rbind(train_positive, train_negative)
#the colnames of the training set should be the same of data_enhancer_svm
colnames(training_set)[c(5:ncol(training_set))]<-gsub(x=gsub(x=colnames(training_set), c(5:ncol(training_set)), pattern=""))

setwd(system.file("extdata", package = "SVM2CRMdata"))
data_level2 <- read.table(file = "GSM393946.distal.p300fromTSS.txt", sep = "\t", stringsAsFactors = FALSE)
data_level2<-data_level2[data_level2[,1]=="chr1",]

DB <- data_level2[, c(1:3)]
colnames(DB)<-c("chromosome", "start", "end")

label <- "p300"

table.final.overlap<-findFeatureOverlap(query=completeTABLE, subject=DB, select="all")

data_enhancer_svm<-createSVMinput(inputpos=table.final.overlap, inputfull=completeTABLE, label1="enhancers", 1)
colnames(data_enhancer_svm)[c(5:ncol(data_enhancer_svm))]<-gsub(gsub(x=colnames(data_enhancer_svm), c(5:ncol(data_enhancer_svm)), pattern=""), pattern="")
```

```

listcolnames<-c("H2AK5ac","H2AK9ac","H3K23ac","H3K27ac","H3K27me3","H3K4me1","H3K4me3")

dftotann<-smoothInputFS(train_positive[,c(6:ncol(train_positive))],listcolnames,k=20)

results<-featSelectionWithKmeans(dftotann,5)

resultsFS<-results[[7]]

resultsFSfilter<-resultsFS[which(resultsFS[,2]>median(resultsFS[,2])),]

resultsFSfilterICRR<-resultsFSfilter[which(resultsFSfilter[,3]<0.50),]

listHM<-resultsFSfilterICRR[,1]
listHM<-gsub(gsub(listHM,pattern="_.",replacement=""),pattern="CD4.",replacement="")

selectFeature<-grep(x=colnames(training_set[,c(6:ncol(training_set))]),pattern=paste(listHM,collapse="|"),value=TRUE)

colSelect<-c("chromosome","start","end","label",selectFeature)
training_set<-training_set[,colSelect]

vecS <- c(2:length(listHM))
typeSVM <- c(0, 6, 7)[1]
costV <- c(0.001, 0.01, 0.1, 1, 10, 100, 1000)[6]
wlabel <- c("not_enhancer", "enhancer")
infofile<-data.frame(a=c(paste(listHM,"signal",sep=".")))
infofile[,1]<-gsub(gsub(x=infofile[,1],pattern="CD4.",replacement=""),pattern=".sort.bed",replacement="")

tuningTAB <- tuningParametersCombROC(training_set = training_set, typeSVM = typeSVM, costV = costV,differentiation=1)

tuningTABfilter<-tuningTAB[tuningTAB$fscore<0.95,]
#row_max_fscore<-which.max(tuningTABfilter[tuningTABfilter$nHM >2,"fscore"])
row_max_fscore<-which.max(tuningTABfilter[, "fscore"])
listHM_prediction<-gsub(tuningTABfilter[row_max_fscore,4],pattern="//",replacement="|")

columnPR<-grep(colnames(training_set),pattern=paste(listHM_prediction,collapse="|"),value=TRUE)

predictionGW(training_set=training_set,data_enhancer_svm=data_enhancer_svm, listHM=columnPR,pcClass.string=TRUE)

```

**smoothInputFS**

*Smooth the signals of the histone marks to prepare the input for feature selection analysis*

**Description**

Give the matrix obtained using getSignal this functions smooth the signals of each histone marks using a particular window (if bin=100).To size of smooth is bin\*k (e.g. a parameter k equal to 2 means that the signal is smooth every 200bp).

**Usage**

```
smoothInputFS(input_ann,k,listcolnames)
```

**Arguments**

<code>input_ann</code>	the data.frame with the training set
<code>k</code>	the size of smooth in bp
<code>listcolnames</code>	the names of column in which perform the smoothing. A vector with the list of histone marks.

**Details**

The smoothing is performed using the median

**Value**

A data.frame with the smoothed signals of histone marks

**Author(s)**

Guidantonio Malagoli Tagliazucchi guidantonio.malagolitagliazucchi@unimore.it

**See Also**

`cisREfindbed`, `featSelectionWithKmeans`, `tuningParametersCombROC`

**Examples**

```
library("GenomicRanges")
library("SVM2CRMdata")

setwd(system.file("data",package="SVM2CRMdata"))
load("CD4_matrixInputSVMbin100window100.rda")
completeTABLE<-CD4_matrixInputSVMbin100window100

new.strings<-gsub(x=colnames(completeTABLE[,c(6:ncol(completeTABLE))]),pattern="CD4.",replacement="")
new.strings<-gsub(new.strings,pattern=".norm.w100.bed",replacement="")
colnames(completeTABLE)[c(6:ncol(completeTABLE))]<-new.strings

#list_file<-grep(dir(),pattern=".sort.txt",value=TRUE)

#train_positive<-getSignal(list_file,chr="chr1",reference="p300.distal.fromTSS.txt",win.size=500,bin.size=10)
#train_negative<-getSignal(list_file,chr="chr1",reference="random.region.hg18.nop300.txt",win.size=500,bin.size=10)
setwd(system.file("data",package="SVM2CRMdata"))
load("train_positive.rda")
load("train_negative.rda")
```

```

  training_set<-rbind(train_positive,train_negative)
  colnames(training_set)[c(5:ncol(training_set))]<-gsub(x=colnames(training_set[,c(5:ncol(training_set))],)

  setwd(system.file("extdata", package = "SVM2CRMdata"))
  data_level2 <- read.table(file = "GSM393946.distal.p300fromTSS.txt",sep = "\t", stringsAsFactors = FALSE)
  data_level2<-data_level2[data_level2[,1]=="chr1",]

  DB <- data_level2[, c(1:3)]
  colnames(DB)<-c("chromosome","start","end")

  label <- "p300"

  table.final.overlap<-findFeatureOverlap(query=completeTABLE,subject=DB,select="all")

  data_enhancer_svm<-createSVMinput(inputpos=table.final.overlap,inputfull=completeTABLE,label1="enhancers",label2="p300")
  colnames(data_enhancer_svm)[c(5:ncol(data_enhancer_svm))]<-gsub(gsub(x=colnames(data_enhancer_svm[,c(5:ncol(data_enhancer_svm))],))

  listcolnames<-c("H2AK5ac","H2AK9ac","H3K23ac","H3K27ac","H3K27me3","H3K4me1","H3K4me3")

  dftotann<-smoothInputFS(train_positive[,c(6:ncol(train_positive))],listcolnames,k=20)

```

**tuningParametersCombROC**

*Test different models using different kernel of SVM, values of cost functions, the number of histone marks.*

**Description**

The function tuningParametersCombROC allow high flexibility: the user can set the type of kernel, the cost parameter of SVM, the number of histone marks. tuningParametersCombROC use performanceSVM and the output is a data.frame where for each model there are the parameters compute with performanceSVM. To help the user to discriminate how to discriminate the best model SVM2CRM implement several function to plot the results of performanceSVM.

**Usage**

```
tuningParametersCombROC(training_set, typeSVM, costV,different.weight=TRUE, vecS,infofile,pcClass.s)
```

**Arguments**

training_set	training set (data.frame)
typeSVM	a vector with the kinds of Kernel
costV	a vector with a list of cost parameters
different.weight	the data are unbalanced (default TRUE)
vecS	a vector with the number of histone marks (e.g. from 2 to x)

<code>infofile</code>	a data.frame where in the column "a" there are the histone marks, while in the column "b" a vectors of letters.
<code>pcClass.string</code>	label of the first class (e.g. "enhancers")
<code>nClass.string</code>	label of the second class (e.g. "not_enhancers")
<code>pcClass</code>	number of positive class in the training set
<code>ncClass</code>	number of negative class in the training set

## Details

Some detailed description

## Value

A data.frame with the values from perfomanceSVM for each trained model.

## Author(s)

Guidantonio Malagoli Tagliazucchi guidantonio.malagolitagliazucchi@unimore.it

## See Also

`cisREfindbed`, `perfomanceSVM`, `plotFscore`, `plotROC`

## Examples

```

library("GenomicRanges")
library("SVM2CRMdata")

setwd(system.file("data", package="SVM2CRMdata"))
load("CD4_matrixInputSVMbin100window1000.rda")
completeTABLE<-CD4_matrixInputSVMbin100window1000

new.strings<-gsub(x=colnames(completeTABLE[,c(6:ncol(completeTABLE))]), pattern="CD4.", replacement="")
new.strings<-gsub(new.strings, pattern=".norm.w100.bed", replacement="")
colnames(completeTABLE)[c(6:ncol(completeTABLE))]<-new.strings

#Create a vector that contain the list of the bed files that you want use during the analysis
#list_file<-grep(dir(), pattern=".sort.txt", value=TRUE)
#print(list_file)
#Here we used a data.frame that contain the genomic coordinates of p300 binding sites
#train_positive<-getSignal(list_file, chr="chr1", reference="p300.distal.fromTSS.txt", win.size=500, bin.size=100)
#train_negative<-getSignal(list_file, chr="chr1", reference="random.region.hg18.nop300.txt", win.size=500, bin.size=100)
#setwd(system.file("data", package="SVM2CRMdata"))
#load("train_positive.rda")
#load("train_negative.rda")

training_set<-rbind(train_positive, train_negative)
colnames(training_set)[c(5:ncol(training_set))]<-gsub(x=colnames(training_set[,c(5:ncol(training_set))]), pattern="CD4.", replacement="")

setwd(system.file("extdata", package = "SVM2CRMdata"))

```

```

data_level2 <- read.table(file = "GSM393946.distal.p300fromTSS.txt", sep = "\t", stringsAsFactors = FALSE)
data_level2<-data_level2[data_level2[,1]=="chr1",]

DB <- data_level2[, c(1:3)]
colnames(DB)<-c("chromosome", "start", "end")

label <- "p300"

table.final.overlap<-findFeatureOverlap(query=completeTABLE, subject=DB, select="all")

data_enhancer_svm<-createSVMinput(inputpos=table.final.overlap, inputfull=completeTABLE, label1="enhancers", la
colnames(data_enhancer_svm)[c(5:ncol(data_enhancer_svm))]<-gsub(gsub(x=colnames(data_enhancer_svm[,c(5:ncol(
listcolnames<-c("H2AK5ac", "H2AK9ac", "H3K23ac", "H3K27ac", "H3K27me3", "H3K4me1", "H3K4me3")

dftotann<-smoothInputFS(train_positive[,c(6:ncol(train_positive))],listcolnames,k=20)

results<-featSelectionWithKmeans(dftotann,5)

resultsFS<-results[[7]]

resultsFSfilter<-resultsFS[which(resultsFS[,2]>median(resultsFS[,2])),]

resultsFSfilterICRR<-resultsFSfilter[which(resultsFSfilter[,3]<0.50),]

listHM<-resultsFSfilterICRR[,1]
listHM<-gsub(gsub(listHM,pattern="_.",replacement=""),pattern="CD4.",replacement="")

selectFeature<-grep(x=colnames(training_set[,c(6:ncol(training_set))]),pattern=paste(listHM,collapse="|"),va
colSelect<-c("chromosome", "start", "end", "label", selectFeature)
training_set<-training_set[,colSelect]

vecS <- c(2:length(listHM))
typeSVM <- c(0, 6, 7)[1]
costV <- c(0.001, 0.01, 0.1, 1, 10, 100, 1000)[6]
wlabel <- c("not_enhancer", "enhancer")
infofile<-data.frame(a=c(paste(listHM,"signal",sep=".")))
infofile[,1]<-gsub(gsub(x=infofile[,1],pattern="CD4.",replacement=""),pattern=".sort.bed",replacement="")

tuningTAB <- tuningParametersCombROC(training_set = training_set, typeSVM = typeSVM, costV = costV,different.w

```

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