

# Package ‘RUVnormalize’

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**Title** RUV for normalization of expression array data

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**Description** RUVnormalize is meant to remove unwanted variation from gene expression data when the factor of interest is not defined, e.g., to clean up a dataset for general use or to do any kind of unsupervised analysis.

**License** GPL-3

**LazyLoad** yes

**Imports** RUVnormalizeData, Biobase

**Enhances** spams

**Depends** R (>= 2.10.0)

**NeedsCompilation** no

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## Contents

clScore . . . . .	2
iterativeRUV . . . . .	4
naiveRandRUV . . . . .	9
naiveReplicateRUV . . . . .	12
svdPlot . . . . .	15

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cIScore

*Computes a distance between two partitions of the same data*


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### Description

The function takes as input two partitions of a dataset into clusters, and returns a number which is small if the two partitions are close, large otherwise.

### Usage

```
cIScore(c1, c2)
```

### Arguments

c1                    A [vector](#) giving the assignment of the samples to cluster for the first partition  
c2                    A [vector](#) giving the assignment of the samples to cluster for the second partition

### Value

A number corresponding to the distance between c1 and c2

### Examples

```
if(require('RUVnormalizeData')){
  ## Load the data
  data('gender', package='RUVnormalizeData')

  Y <- t(exprs(gender))
  X <- as.numeric(phenoData(gender)$gender == 'M')
  X <- X - mean(X)
  X <- cbind(X/(sqrt(sum(X^2))))
  chip <- annotation(gender)

  ## Extract regions and labs for plotting purposes
  lregions <- sapply(rownames(Y),FUN=function(s) strsplit(s,'_')[[1]][2])
  llabs <- sapply(rownames(Y),FUN=function(s) strsplit(s,'_')[[1]][3])

  ## Dimension of the factors
  m <- nrow(Y)
  n <- ncol(Y)
  p <- ncol(X)

  Y <- scale(Y, scale=FALSE) # Center gene expressions

  cIdx <- which(featureData(gender)$isNegativeControl) # Negative control genes

  ## Prepare plots
```

```

annot <- cbind(as.character(sign(X)))
colnames(annot) <- 'gender'
plAnnots <- list('gender'='categorical')
lab.and.region <- apply(rbind(lregions, llabs),2,FUN=function(v) paste(v,collapse='_'))
gender.col <- c('-1' = "deppink3", '1' = "blue")

## Remove platform effect by centering.

Y[chip=='hgu95a.db',] <- scale(Y[chip=='hgu95a.db',], scale=FALSE)
Y[chip=='hgu95av2.db',] <- scale(Y[chip=='hgu95av2.db',], scale=FALSE)

## Number of genes kept for clustering, based on their variance
nKeep <- 1260

##-----
## Naive RUV-2 no shrinkage
##-----

k <- 20
nu <- 0

## Correction
nsY <- naiveRandRUV(Y, cIdx, nu.coeff=0, k=k)

## Clustering of the corrected data
sdY <- apply(nsY, 2, sd)
ssd <- sort(sdY,decreasing=TRUE,index.return=TRUE)$ix
kmres2ns <- kmeans(nsY[,ssd[1:nKeep],drop=FALSE],centers=2,nstart=200)
vclust2ns <- kmres2ns$cluster
nsScore <- cIScore(vclust2ns, X)

## Plot of the corrected data
svdRes2ns <- NULL
svdRes2ns <- svdPlot(nsY[, ssd[1:nKeep], drop=FALSE],
                    annot=annot,
                    labels=lab.and.region,
                    svdRes=svdRes2ns,
                    plAnnots=plAnnots,
                    kColors=gender.col, file=NULL)

##-----
## Naive RUV-2 + shrinkage
##-----

k <- m
nu.coeff <- 1e-2

## Correction
nY <- naiveRandRUV(Y, cIdx, nu.coeff=nu.coeff, k=k)

## Clustering of the corrected data
sdY <- apply(nY, 2, sd)
ssd <- sort(sdY,decreasing=TRUE,index.return=TRUE)$ix

```

```

kmres2 <- kmeans(nY[,ssd[1:nKeep],drop=FALSE],centers=2,nstart=200)
vclust2 <- kmres2$cluster
nScore <- c1Score(vclust2,X)

## Plot of the corrected data
svdRes2 <- NULL
svdRes2 <- svdPlot(nY[, ssd[1:nKeep], drop=FALSE],
                  annot=annot,
                  labels=lab.and.region,
                  svdRes=svdRes2,
                  plAnnots=plAnnots,
                  kColors=gender.col, file=NULL)
}

```

---

iterativeRUV	<i>Remove unwanted variation from a gene expression matrix using control genes, optionally replicate samples, and iterative estimates of the factor of interest</i>
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## Description

The function takes as input a gene expression matrix as well as the index of negative control genes and replicate samples. It estimates and remove unwanted variation from the gene expression. The major difference with naiveRandRUV and naiveReplicateRUV is that iterativeRUV jointly estimates the factor of interest and the unwanted variation term. It does so iteratively, by estimating each term using the current estimate of the other one.

## Usage

```
iterativeRUV(Y, cIdx, scIdx=NULL, paramXb, k, nu.coeff=0, cEps=1e-08, maxIter=30,
             Wmethod="svd", Winit=NULL, wUpdate=maxIter + 1, tol=1e-6)
```

## Arguments

Y	Expression matrix where the rows are the samples and the columns are the genes.
cIdx	Column index of the negative control genes in Y, for estimation of unwanted variation.
scIdx	Matrix giving the set of replicates. Each row is a set of arrays corresponding to replicates of the same sample. The number of columns is the size of the largest set of replicates, and the smaller sets are padded with -1 values. For example if the sets of replicates are (1,11,21), (2,3), (4,5), (6,7,8), the scIdx should be 1 11 21 2 3 -1 4 5 -1 6 7 8
paramXb	A <a href="#">list</a> containing parameters for the estimation of the term of interest: K corresponds to the rank of X. lambda is the regularization parameter. Large values of lambda lead to sparser, more shrunk estimates of beta. D, batch, iter and mode should not be modified unless you are familiar with sparse dictionary learning algorithms.

<code>k</code>	Desired rank for the estimated unwanted variation term. The returned rank may be lower if the replicate arrays and control genes did not contain a signal of rank <code>k</code> .
<code>nu.coeff</code>	Regularization parameter for the unwanted variation.
<code>cEps</code>	tolerance for relative changes of $W_a$ and $X_b$ estimators at each step. When both get smaller than <code>cEps</code> , the iterations stop.
<code>maxIter</code>	Maximum number of iterations.
<code>Wmethod</code>	'svd' or 'rep', depending whether $W$ is estimated from control genes or replicate samples.
<code>Winit</code>	Optionally provides an initial value for $W$ .
<code>wUpdate</code>	Number of iterations between two updates of $W$ . By default, $W$ is never updated. Make sure that enough iterations are done after the last update of $W$ . E.g, setting $W$ to <code>maxIter</code> will only allow for one iteration of estimating $\alpha$ given $(X_b, W)$ and no re-estimation of $X_b$ .
<code>tol</code>	Smallest ratio allowed between a squared singular value of $Y[, cIdx]$ and the largest of these squared singular values. All smaller singular values are discarded.

## Details

In terms of model, the rank `k` can be thought of as the number of independent sources of unwanted variation in the data (i.e., if one source is a linear combination of other sources, it does not increase the rank). The ridge `nu.coeff` should be inversely proportional to the (expected) magnitude of the unwanted variation.

In practice, even if the real number of independent sources of unwanted variation (resp. their magnitude) is known, using a smaller `k` (resp., larger ridge) could yield better corrections because one may not have enough samples to effectively estimate all the effects.

More intuition and guidance on the practical choice of these parameters are available in the paper (<http://biostatistics.oxfordjournals.org/content/17/1/16.full>) and its supplement (<http://biostatistics.oxfordjournals.org/content/suppl/2015/08/17/kxv026.DC1/kxv026supp.pdf>). In particular: - Equation 2.3 in the manuscript gives an interpretation of the ridge parameter in terms of a probabilistic model. - Section 5.1 of the manuscript provides guidelines to select both parameters on real data. - Section 3 of the supplement compares the effect of reducing the rank and increasing the ridge. - Section 4 of the supplement gives a detailed discussion of how to select the ridge parameter on a real example.

## Value

A `list` containing the following terms:

<code>X, b</code>	if <code>p</code> is not <code>NULL</code> , contains an estimate of the factor of interest ( $X$ ) and its effect ( $\beta$ ) obtained using rank- <code>p</code> restriction of the SVD of $Y - W\alpha$ .
<code>W, a</code>	Estimates of the unwanted variation factors ( $W$ ) and their effect ( $\alpha$ ).
<code>cY</code>	The corrected expression matrix $Y - W\alpha$ .

**Examples**

```

if(require('RUVnormalizeData') && require('spams')){
  ## Load the spams library
  library(spams)

  ## Load the data
  data('gender', package='RUVnormalizeData')

  Y <- t(exprs(gender))
  X <- as.numeric(phenoData(gender)$gender == 'M')
  X <- X - mean(X)
  X <- cbind(X/(sqrt(sum(X^2))))
  chip <- annotation(gender)

  ## Extract regions and labs for plotting purposes
  lregions <- sapply(rownames(Y),FUN=function(s) strsplit(s,'_')[[1]][2])
  llabs <- sapply(rownames(Y),FUN=function(s) strsplit(s,'_')[[1]][3])

  ## Dimension of the factors
  m <- nrow(Y)
  n <- ncol(Y)
  p <- ncol(X)

  Y <- scale(Y, scale=FALSE) # Center gene expressions

  cIdx <- which(featureData(gender)$isNegativeControl) # Negative control genes

  ## Prepare plots
  annot <- cbind(as.character(sign(X)))
  colnames(annot) <- 'gender'
  plAnnots <- list('gender'='categorical')
  lab.and.region <- apply(rbind(lregions, llabs),2,FUN=function(v) paste(v,collapse='_'))
  gender.col <- c('-1' = "deppink3", '1' = "blue")

  ## Remove platform effect by centering.

  Y[chip=='hgu95a.db',] <- scale(Y[chip=='hgu95a.db',], scale=FALSE)
  Y[chip=='hgu95av2.db',] <- scale(Y[chip=='hgu95av2.db',], scale=FALSE)

  ## Number of genes kept for clustering, based on their variance
  nKeep <- 1260

  ## Prepare control samples

  scIdx <- matrix(-1,84,3)
  rny <- rownames(Y)
  added <- c()
  c <- 0

  ## Replicates by lab
  for(r in 1:(length(rny) - 1)){
    if(r %in% added)

```

```

        next
      c <- c+1
      scIdx[c,1] <- r
      cc <- 2
      for(rr in seq(along=rny[(r+1):length(rny)])){
        if(all(strsplit(rny[r], '_')[[1]][-3] == strsplit(rny[r+rr], '_')[[1]][-3])){
          scIdx[c,cc] <- r+rr
          cc <- cc+1
          added <- c(added,r+rr)
        }
      }
    }
  }
  scIdxLab <- scIdx

  scIdx <- matrix(-1,84,3)
  rny <- rownames(Y)
  added <- c()
  c <- 0

  ## Replicates by region
  for(r in 1:(length(rny) - 1)){
    if(r %in% added)
      next
    c <- c+1
    scIdx[c,1] <- r
    cc <- 2
    for(rr in seq(along=rny[(r+1):length(rny)])){
      if(all(strsplit(rny[r], '_')[[1]][-2] == strsplit(rny[r+rr], '_')[[1]][-2])){
        scIdx[c,cc] <- r+rr
        cc <- cc+1
        added <- c(added,r+rr)
      }
    }
  }
  scIdx <- rbind(scIdxLab,scIdx)

  ## Number of genes kept for clustering, based on their variance
  nKeep <- 1260

  ## Prepare plots
  annot <- cbind(as.character(sign(X)))
  colnames(annot) <- 'gender'
  plAnnots <- list('gender'='categorical')
  lab.and.region <- apply(rbind(lregions, llabs),2,FUN=function(v) paste(v,collapse='_'))
  gender.col <- c('-1' = "deppink3", '1' = "blue")

  ##-----
  ## Iterative replicate-based
  ##-----

  cEps <- 1e-6
  maxIter <- 30
  p <- 20

```

```

paramXb <- list()
paramXb$K <- p
paramXb$D <- matrix(c(0.),nrow = 0,ncol=0)
paramXb$batch <- TRUE
paramXb$iter <- 1
paramXb$mode <- 'PENALTY'
paramXb$lambda <- 0.25

## Correction
iRes <- iterativeRUV(Y, cIdx, scIdx, paramXb, k=20, nu.coeff=0,
                    cEps, maxIter,
                    Wmethod='rep', wUpdate=11)

ucY <- iRes$cY

## Cluster the corrected data
sdY <- apply(ucY, 2, sd)
ssd <- sort(sdY,decreasing=TRUE,index.return=TRUE)$ix
kmresIter <- kmeans(ucY[,ssd[1:nKeep]],centers=2,nstart=200)
vclustIter <- kmresIter$cluster
IterScore <- clScore(vclustIter,X)

## Plot the corrected data
svdResIter <- NULL
svdResIter <- svdPlot(ucY[, ssd[1:nKeep], drop=FALSE],
                    annot=annot,
                    labels=lab.and.region,
                    svdRes=svdResIter,
                    plAnnots=plAnnots,
                    kColors=gender.col, file=NULL)

##-----
## Iterated ridge
##-----

paramXb <- list()
paramXb$K <- p
paramXb$D <- matrix(c(0.),nrow = 0,ncol=0)
paramXb$batch <- TRUE
paramXb$iter <- 1
paramXb$mode <- 'PENALTY' #2
paramXb$lambda <- 1
paramXb$lambda2 <- 0

## Correction
iRes <- iterativeRUV(Y, cIdx, scIdx=NULL, paramXb, k=nrow(Y), nu.coeff=1e-2/2,
                    cEps, maxIter,
                    Wmethod='svd', wUpdate=11)

nrcY <- iRes$cY

## Cluster the corrected data

```



```

sdY <- apply(nrcY, 2, sd)
ssd <- sort(sdY,decreasing=TRUE,index.return=TRUE)$ix
kmresIter <- kmeans(nrcY[,ssd[1:nKeep]],centers=2,nstart=200)
vclustIter <- kmresIter$cluster
IterRandScore <- c1Score(vclustIter,X)

## Plot the corrected data
svdResIterRand <- NULL
svdResIterRand <- svdPlot(nrcY[, ssd[1:nKeep], drop=FALSE],
                        annot=annot,
                        labels=lab.and.region,
                        svdRes=svdResIterRand,
                        plAnnots=plAnnots,
                        kColors=gender.col, file=NULL)
}

```

---

naiveRandRUV	<i>Remove unwanted variation from a gene expression matrix using negative control genes</i>
--------------	---

---

## Description

The function takes as input a gene expression matrix as well as the index of negative control genes. It estimates unwanted variation from these control genes, and removes them by regression, using ridge and/or rank regularization.

## Usage

```
naiveRandRUV(Y, cIdx, nu.coeff=0.001, k=min(nrow(Y), length(cIdx)), tol=1e-6)
```

## Arguments

Y	Expression matrix where the rows are the samples and the columns are the genes.
cIdx	Column index of the negative control genes in Y, for estimation of unwanted variation.
nu.coeff	Regularization parameter for the unwanted variation.
k	Desired rank for the estimated unwanted variation term.
tol	Smallest ratio allowed between a squared singular value of Y[, cIdx] and the largest of these squared singular values. All smaller singular values are discarded.

## Details

In terms of model, the rank k can be thought of as the number of independent sources of unwanted variation in the data (i.e., if one source is a linear combination of other sources, it does not increase the rank). The ridge nu.coeff should be inversely proportional to the (expected) magnitude of the unwanted variation.

In practice, even if the real number of independent sources of unwanted variation (resp. their magnitude) is known, using a smaller  $k$  (resp., larger ridge) could yield better corrections because one may not have enough samples to effectively estimate all the effects.

More intuition and guidance on the practical choice of these parameters are available in the paper (<http://biostatistics.oxfordjournals.org/content/17/1/16.full>) and its supplement (<http://biostatistics.oxfordjournals.org/content/suppl/2015/08/17/kxv026.DC1/kxv026supp.pdf>). In particular: - Equation 2.3 in the manuscript gives an interpretation of the ridge parameter in terms of a probabilistic model. - Section 5.1 of the manuscript provides guidelines to select both parameters on real data. - Section 3 of the supplement compares the effect of reducing the rank and increasing the ridge. - Section 4 of the supplement gives a detailed discussion of how to select the ridge parameter on a real example.

## Value

A *matrix* corresponding to the gene expression after subtraction of the estimated unwanted variation term.

## Examples

```
if(require('RUVnormalizeData')){

  ## Load the data
  data('gender', package='RUVnormalizeData')

  Y <- t(exprs(gender))
  X <- as.numeric(phenoData(gender)$gender == 'M')
  X <- X - mean(X)
  X <- cbind(X/(sqrt(sum(X^2))))
  chip <- annotation(gender)

  ## Extract regions and labs for plotting purposes
  lregions <- sapply(rownames(Y),FUN=function(s) strsplit(s,'_')[[1]][2])
  llabs <- sapply(rownames(Y),FUN=function(s) strsplit(s,'_')[[1]][3])

  ## Dimension of the factors
  m <- nrow(Y)
  n <- ncol(Y)
  p <- ncol(X)

  Y <- scale(Y, scale=FALSE) # Center gene expressions

  cIdx <- which(featureData(gender)$isNegativeControl) # Negative control genes

  ## Prepare plots
  annot <- cbind(as.character(sign(X)))
  colnames(annot) <- 'gender'
  plAnnots <- list('gender'='categorical')
  lab.and.region <- apply(rbind(lregions, llabs),2,FUN=function(v) paste(v,collapse='_'))
  gender.col <- c('-1' = "deeppink3", '1' = "blue")

  ## Remove platform effect by centering.
```

```

Y[chip=='hgu95a.db',] <- scale(Y[chip=='hgu95a.db',], scale=FALSE)
Y[chip=='hgu95av2.db',] <- scale(Y[chip=='hgu95av2.db',], scale=FALSE)

## Number of genes kept for clustering, based on their variance
nKeep <- 1260

##-----
## Naive RUV-2 no shrinkage
##-----

k <- 20
nu <- 0

## Correction
nsY <- naiveRandRUV(Y, cIdx, nu.coeff=0, k=k)

## Clustering of the corrected data
sdY <- apply(nsY, 2, sd)
ssd <- sort(sdY,decreasing=TRUE,index.return=TRUE)$ix
kmres2ns <- kmeans(nsY[,ssd[1:nKeep],drop=FALSE],centers=2,nstart=200)
vclust2ns <- kmres2ns$cluster
nsScore <- clScore(vclust2ns, X)

## Plot of the corrected data
svdRes2ns <- NULL
svdRes2ns <- svdPlot(nsY[, ssd[1:nKeep], drop=FALSE],
                    annot=annot,
                    labels=lab.and.region,
                    svdRes=svdRes2ns,
                    plAnnots=plAnnots,
                    kColors=gender.col, file=NULL)

##-----
## Naive RUV-2 + shrinkage
##-----

k <- m
nu.coeff <- 1e-2

## Correction
nY <- naiveRandRUV(Y, cIdx, nu.coeff=nu.coeff, k=k)

## Clustering of the corrected data
sdY <- apply(nY, 2, sd)
ssd <- sort(sdY,decreasing=TRUE,index.return=TRUE)$ix
kmres2 <- kmeans(nY[,ssd[1:nKeep],drop=FALSE],centers=2,nstart=200)
vclust2 <- kmres2$cluster
nScore <- clScore(vclust2,X)

## Plot of the corrected data
svdRes2 <- NULL
svdRes2 <- svdPlot(nY[, ssd[1:nKeep], drop=FALSE],

```

```

        annot=annot,
        labels=lab.and.region,
        svdRes=svdRes2,
        plAnnots=plAnnots,
        kColors=gender.col, file=NULL)
    }

```

---

naiveReplicateRUV      *Remove unwanted variation from a gene expression matrix using replicate samples*

---

### Description

The function takes as input a gene expression matrix as well as the index of negative control genes and replicate samples. It estimates and remove unwanted variation from the gene expression.

### Usage

```
naiveReplicateRUV(Y, cIdx, scIdx, k, rrem=NULL, p=NULL, tol=1e-6)
```

### Arguments

Y	Expression matrix where the rows are the samples and the columns are the genes.
cIdx	Column index of the negative control genes in Y, for estimation of unwanted variation.
scIdx	Matrix giving the set of replicates. Each row is a set of arrays corresponding to replicates of the same sample. The number of columns is the size of the largest set of replicates, and the smaller sets are padded with -1 values. For example if the sets of replicates are (1,11,21), (2,3), (4,5), (6,7,8), the scIdx should be 1 11 21 2 3 -1 4 5 -1 6 7 8
k	Desired rank for the estimated unwanted variation term. The returned rank may be lower if the replicate arrays and control genes did not contain a signal of rank k.
rrem	Optional, indicates which arrays should be removed from the returned result. Useful if the replicate arrays were not actual samples but mixtures of RNA which are only useful to estimate UV but which should not be included in the analysis.
p	Optional. If given, the function returns an estimate of the term of interest using rank-p restriction of the SVD of the corrected matrix.
tol	Directions of variance lower than this value in the replicate samples are dropped (which may result in an estimated unwanted variation term of rank smaller than k).

**Details**

In terms of model, the rank  $k$  can be thought of as the number of independent sources of unwanted variation in the data (i.e., if one source is a linear combination of other sources, it does not increase the rank).

In practice, even if the real number of independent sources of unwanted variation is known, using a smaller  $k$  (resp., larger ridge) could yield better corrections because one may not have enough samples to effectively estimate all the effects.

**Value**

A `list` containing the following terms:

<code>X, b</code>	if <code>p</code> is not <code>NULL</code> , contains an estimate of the factor of interest ( <code>X</code> ) and its effect (beta) obtained using rank- <code>p</code> restriction of the SVD of <code>Y - W alpha</code> .
<code>W, a</code>	Estimates of the unwanted variation factors ( <code>W</code> ) and their effect ( <code>alpha</code> ).
<code>cY</code>	The corrected expression matrix <code>Y - W alpha</code>
<code>Yctls</code>	The differences of replicate arrays which were used to estimate <code>W</code> and <code>alpha</code> .

**Examples**

```
if(require('RUVnormalizeData')){
  ## Load the data
  data('gender', package='RUVnormalizeData')

  Y <- t(exprs(gender))
  X <- as.numeric(phenoData(gender)$gender == 'M')
  X <- X - mean(X)
  X <- cbind(X/(sqrt(sum(X^2))))
  chip <- annotation(gender)

  ## Extract regions and labs for plotting purposes
  lregions <- sapply(rownames(Y),FUN=function(s) strsplit(s,'_')[[1]][2])
  llabs <- sapply(rownames(Y),FUN=function(s) strsplit(s,'_')[[1]][3])

  ## Dimension of the factors
  m <- nrow(Y)
  n <- ncol(Y)
  p <- ncol(X)

  Y <- scale(Y, scale=FALSE) # Center gene expressions

  cIdx <- which(featureData(gender)$isNegativeControl) # Negative control genes

  ## Prepare plots
  annot <- cbind(as.character(sign(X)))
  colnames(annot) <- 'gender'
  plAnnots <- list('gender'='categorical')
  lab.and.region <- apply(rbind(lregions, llabs),2,FUN=function(v) paste(v,collapse='_'))
  gender.col <- c('-1' = "deeppink3", '1' = "blue")
}
```

```

## Remove platform effect by centering.

Y[chip=='hgu95a.db',] <- scale(Y[chip=='hgu95a.db',], scale=FALSE)
Y[chip=='hgu95av2.db',] <- scale(Y[chip=='hgu95av2.db',], scale=FALSE)

## Prepare control samples

scIdx <- matrix(-1,84,3)
rny <- rownames(Y)
added <- c()
c <- 0

## Replicates by lab
for(r in 1:(length(rny) - 1)){
  if(r %in% added)
    next
  c <- c+1
  scIdx[c,1] <- r
  cc <- 2
  for(rr in seq(along=rny[(r+1):length(rny)])){
    if(all(strsplit(rny[r], '_')[[1]][-3] == strsplit(rny[r+rr], '_')[[1]][-3])){
      scIdx[c,cc] <- r+rr
      cc <- cc+1
      added <- c(added,r+rr)
    }
  }
}
scIdxLab <- scIdx

scIdx <- matrix(-1,84,3)
rny <- rownames(Y)
added <- c()
c <- 0

## Replicates by region
for(r in 1:(length(rny) - 1)){
  if(r %in% added)
    next
  c <- c+1
  scIdx[c,1] <- r
  cc <- 2
  for(rr in seq(along=rny[(r+1):length(rny)])){
    if(all(strsplit(rny[r], '_')[[1]][-2] == strsplit(rny[r+rr], '_')[[1]][-2])){
      scIdx[c,cc] <- r+rr
      cc <- cc+1
      added <- c(added,r+rr)
    }
  }
}
scIdx <- rbind(scIdxLab,scIdx)

## Number of genes kept for clustering, based on their variance

```

```

nKeep <- 1260

## Prepare plots
annot <- cbind(as.character(sign(X)))
colnames(annot) <- 'gender'
plAnnots <- list('gender'='categorical')
lab.and.region <- apply(rbind(lregions, llabs),2,FUN=function(v) paste(v,collapse='_'))
gender.col <- c('-1' = "deeppink3", '1' = "blue")

## Remove platform effect by centering.

## Correction
sRes <- naiveReplicateRUV(Y, cIdx, scIdx, k=20)

## Clustering on the corrected data
sdY <- apply(sRes$cY, 2, sd)
ssd <- sort(sdY,decreasing=TRUE,index.return=TRUE)$ix
kmresRep <- kmeans(sRes$cY[,ssd[1:nKeep],drop=FALSE],centers=2,nstart=200)
vclustRep <- kmresRep$cluster
RepScore <- clScore(vclustRep,X)

## Plot of the corrected data
svdResRep <- NULL
svdResRep <- svdPlot(sRes$cY[, ssd[1:nKeep], drop=FALSE],
                    annot=annot,
                    labels=lab.and.region,
                    svdRes=svdResRep,
                    plAnnots=plAnnots,
                    kColors=gender.col, file=NULL)
}

```

svdPlot

*Plot the data projected into the space spanned by their first two principal components*

### Description

The function takes as input a gene expression matrix and plots the data projected into the space spanned by their first two principal components.

### Usage

```
svdPlot(Y, annot=NULL, labels=NULL, svdRes=NULL, plAnnots=NULL, kColors=NULL, file=NULL)
```

### Arguments

Y Expression matrix where the rows are the samples and the columns are the genes.

annot	A matrix containing the annotation to be plotted. Each row must correspond to a sample (row) of argument Y, each column must be a categorical or continuous descriptor for the sample. Optional.
labels	A vector with one element per sample (row) of argument Y. If this argument is specified, each sample is represented by its label. Otherwise, it is represented by a dot (if no annotation is provided) or by the value of the annotation. Optional.
svdRes	A list containing the result of <code>svd(Y)</code> , possibly restricted to the first few singular values. Optional: if not provided, the function computes the SVD.
plAnnots	A list specifying whether each column of the <code>annot</code> argument corresponds to a categorical or continuous factor. Each element of the list is named after a column of <code>annot</code> , and contains a string 'categorical' or 'continuous'. For each element of this list, a plot is produced where the samples are represented by colors corresponding to their annotation. Optional.
kColors	A vector of colors to be used to represent categorical factors. Optional: a default value is provided. If a categorical factors has more levels than the number of colors provided, colors are not used and the factor is represented in black.
file	A string giving the path to a pdf file for the plot. Optional.

### Value

A [list](#) containing the result of `svd(Y, nu=2, nv=0)`.

### Examples

```
if(require('RUVnormalizeData')){

  ## Load the data
  data('gender', package='RUVnormalizeData')

  Y <- t(exprs(gender))
  X <- as.numeric(phenoData(gender)$gender == 'M')
  X <- X - mean(X)
  X <- cbind(X/(sqrt(sum(X^2))))
  chip <- annotation(gender)

  ## Extract regions and labs for plotting purposes
  lregions <- sapply(rownames(Y),FUN=function(s) strsplit(s,'_')[[1]][2])
  llabs <- sapply(rownames(Y),FUN=function(s) strsplit(s,'_')[[1]][3])

  ## Dimension of the factors
  m <- nrow(Y)
  n <- ncol(Y)
  p <- ncol(X)

  Y <- scale(Y, scale=FALSE) # Center gene expressions

  cIdx <- which(featureData(gender)$isNegativeControl) # Negative control genes

  ## Prepare plots
  annot <- cbind(as.character(sign(X)))
}
```



```

colnames(annot) <- 'gender'
plAnnots <- list('gender'='categorical')
lab.and.region <- apply(rbind(lregions, llabs),2,FUN=function(v) paste(v,collapse='_'))
gender.col <- c('-1' = "deeppink3", '1' = "blue")

## Remove platform effect by centering.

Y[chip=='hgu95a.db',] <- scale(Y[chip=='hgu95a.db',], scale=FALSE)
Y[chip=='hgu95av2.db',] <- scale(Y[chip=='hgu95av2.db',], scale=FALSE)

## Number of genes kept for clustering, based on their variance
nKeep <- 1260

##-----
## Naive RUV-2 no shrinkage
##-----

k <- 20
nu <- 0

## Correction
nsY <- naiveRandRUV(Y, cIdx, nu.coeff=0, k=k)

## Clustering of the corrected data
sdY <- apply(nsY, 2, sd)
ssd <- sort(sdY,decreasing=TRUE,index.return=TRUE)$ix
kmres2ns <- kmeans(nsY[,ssd[1:nKeep],drop=FALSE],centers=2,nstart=200)
vclust2ns <- kmres2ns$cluster
nsScore <- c1Score(vclust2ns, X)

## Plot of the corrected data
svdRes2ns <- NULL
svdRes2ns <- svdPlot(nsY[, ssd[1:nKeep], drop=FALSE],
                    annot=annot,
                    labels=lab.and.region,
                    svdRes=svdRes2ns,
                    plAnnots=plAnnots,
                    kColors=gender.col, file=NULL)

##-----
## Naive RUV-2 + shrinkage
##-----

k <- m
nu.coeff <- 1e-2

## Correction
nY <- naiveRandRUV(Y, cIdx, nu.coeff=nu.coeff, k=k)

## Clustering of the corrected data
sdY <- apply(nY, 2, sd)
ssd <- sort(sdY,decreasing=TRUE,index.return=TRUE)$ix
kmres2 <- kmeans(nY[,ssd[1:nKeep],drop=FALSE],centers=2,nstart=200)

```

```
vclust2 <- kmres2$cluster
nScore <- clScore(vclust2,X)

## Plot of the corrected data
svdRes2 <- NULL
svdRes2 <- svdPlot(nY[, ssd[1:nKeep], drop=FALSE],
                  annot=annot,
                  labels=lab.and.region,
                  svdRes=svdRes2,
                  plAnnots=plAnnots,
                  kColors=gender.col, file=NULL)
}
```

# Index

`clScore`, [2](#)

`iterativeRUV`, [4](#)

`list`, [4](#), [5](#), [13](#), [16](#)

`matrix`, [10](#)

`naiveRandRUV`, [9](#)

`naiveReplicateRUV`, [12](#)

`svdPlot`, [15](#)

`vector`, [2](#)