# Package 'pvca'

May 11, 2024

Type Package

Title Principal Variance Component Analysis (PVCA)

Version 1.44.0
<b>Date</b> 2013-09-25
Author Pierre Bushel <bushel@niehs.nih.gov></bushel@niehs.nih.gov>
Maintainer Jianying LI <1i11@niehs.nih.gov>
Description This package contains the function to assess the batch sources by fitting all ``sources" as random effects including two-way interaction terms in the Mixed Model(depends on lme4 package) to selected principal components, which were obtained from the original data correlation matrix. This package accompanies the book "Batch Effects and Noise in Microarray Experiements, chapter 12.
<b>Depends</b> R (>= $2.15.1$ )
Imports Matrix, Biobase, vsn, stats, lme4
Suggests golubEsets
<b>License</b> LGPL ( $>= 2.0$ )
LazyLoad yes
biocViews Microarray, BatchEffect
git_url https://git.bioconductor.org/packages/pvca
git_branch RELEASE_3_19
git_last_commit aa9b186
git_last_commit_date 2024-04-30
Repository Bioconductor 3.19
Date/Publication 2024-05-10
Contents
pvca-package
Index
1

2 pvca-package

pvca-package

A package that provides an approach to assess the source of batch effects in a microarray gene expression experiment

## **Description**

This package contains the function to assess the batch sources by fitting all "sources" as random effects including two-way interaction terms in the Mixed Model(depends on lme4 package) to selected principal components, which were obtained from the original data correlation matrix. This package accompanies the book "Batch Effects and Noise in Microarray Experiments, chapter 12.

#### **Details**

Package: pvca Type: Package Version: 1.0

Date: 2012-09-11 License: LGPL (>= 2.0)

library(golubEsets)

data(Golub\_Merge) pct\_threshold <- 0.6 batch.factors <- c("ALL.AML", "BM.PB", "Source")

pvcaObj <- pvcaBatchAssess (Golub\_Merge, batch.factors, pct\_threshold) bp <- barplot(pvcaObj\$dat, xlab = "Effects", ylab = "Weighted average proportion variance", ylim= c(0,1.1),col = c("blue"), las=2, main="PVCA estimation bar chart") axis(1, at = bp, labels = pvcaObj\$label, xlab = "Effects", cex.axis = 0.5, las=2) values = pvcaObj\$dat new\_values = round(values, 3) text(bp,pvcaObj\$dat,labels = new\_values, pos=3, cex = 0.8) print(sessionInfo())

## Author(s)

Pierre Bushel <bushel@niehs.nih.gov>

Maintainer: Jianying LI li11@niehs.nih.gov>

## References

Batch Effects and Noise in Microarray Experiments: Sources and Solutions. 2009 John Wiley & Sons, Ltd.

pvcaBatchAssess 3

pvcaBatchAssess	Principal Variance Component Analysis (PVCA)

#### **Description**

This package contains the function to assess the batch sources by fitting all "sources" as random effects including two-way interaction terms in the Mixed Model(depends on lme4 package) to selected principal components, which were obtained from the original data correlation matrix. This package accompanies the book "Batch Effects and Noise in Microarray Experiements, chapter 12.

## Usage

pvcaBatchAssess(abatch, batch.factors, threshold)

## **Arguments**

abatch an instance of ExpresseionSet which can be imported from Biobase

batch.factors A vector of factors that the mixed linear model will be fit on

threshold the percentile value of the minimum amount of the variabilities that the selected

principal components need to explain

## Details

Often times "batch effects" are present in microarray data due to any number of factors, including e.g. a poor experimental design or when the gene expression data is combined from different studies with limited standardization. To estimate the variability of experimental effects including batch, a novel hybrid approach known as principal variance component analysis (PVCA) has been developed. The approach leverages the strengths of two very popular data analysis methods: first, principal component analysis (PCA) is used to efficiently reduce data dimension while maintaining the majority of the variability in the data, and variance components analysis (VCA) fits a mixed linear model using factors of interest as random effects to estimate and partition the total variability. The PVCA approach can be used as a screening tool to determine which sources of variability (biological, technical or other) are most prominent in a given microarray data set. Using the eigenvalues associated with their corresponding eigenvectors as weights, associated variations of all factors are standardized and the magnitude of each source of variability (including each batch effect) is presented as a proportion of total variance. Although PVCA is a generic approach for quantifying the corresponding proportion of variation of each effect, it can be a handy assessment for estimating batch effect before and after batch normalization.

## Value

1 4			. 1 C	C 1 , 1 CC ,	C 1
dat A	A numerica vector	contains the	percentile of source	es of batch effect	for each

term

label A character vector containing the name for each term for plot label purpose

4 pvcaBatchAssess

## Note

Modified and maintained by Jianying Li

## Author(s)

Pierre Bushel

## **Examples**

## **Index**

```
* BatchEffect
pvcaBatchAssess, 3
* MixedModel
pvcaBatchAssess, 3
* PCA
pvcaBatchAssess, 3
* package
pvca-package, 2

pvca (pvca-package), 2
pvca-package, 2
pvcaBatchAssess, 3
```