

AC-PCA: simultaneous dimension reduction and adjustment for confounding variation

User's Guide

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AC-PCA: simultaneous dimension reduction and adjustment for confounding variation
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1 Introduction

1.1 Scope

Dimension reduction methods are commonly applied to visualize datapoints in a lower dimensional space and identify dominant patterns in the data. Confounding variation, technically and biologically originated, may affect the performance of these methods, and hence the visualization and interpretation of the results. AC-PCA simultaneously performs dimension reduction and adjustment for confounding variation. The intuition of AC-PCA is that it captures the variations that are invariant to the confounding factors. One major application area for AC-PCA is the transcriptome data, and it can be implemented to other data types as well.

1.2 How to get help

This user’s guide addresses many scenarios for confounding factors. Additional questions about AC-PCA can be sent to linzx06@gmail.com. Comments on possible improvements are very welcome.

1.3 Input for AC-PCA

AC-PCA requires two inputs: the data matrix X and the confounder matrix Y . X is of dimension $n \times p$, where n is the number of samples and p is the number of variables (genes). Y is of dimension $n \times q$, where n is the number of samples and q is the number of confounding factors. Missing data is allowed in both X and Y . The columns in X and Y are assumed to be centered.

Confounding factors depend on the experimental design and can also depend on the scientific question: confounding factors can be different for the same experiment. As an example, consider a transcriptome experiment where gene expression levels were measured in multiple tissues from multiple species. If one wants to capture the variation across tissues that is shared among species, then the species labels are the confounding factors. In contrast, if the variation across species but shared among tissues are desirable, the tissue labels are the confounding factors. AC-PCA is suitable for experiments with complex designs.

In the following sections, we implement AC-PCA and provide details on how to design Y for various types of confounders and various experimental designs.

2 Theory behind AC-PCA

2.1 AC-PCA in a general form

Let X be the $N \times p$ data matrix and Y be the $N \times l$ matrix for l confounding factors. Let $K = YY^T$. We propose the following objective function to adjust

for confounding variation:

$$\begin{aligned} & \underset{v \in \mathbb{R}^p}{\text{maximize}} && v^T X^T X v - \lambda v^T X^T K X v \\ & \text{subject to} && \|v\|_2^2 \leq 1, \end{aligned} \quad (1)$$

We can choose Y so that $v^T X^T K X v$ represents the confounding variation in the projected data. Formula 1 can be generalized to other kernels on Y . In the R package, we provide options for linear kernel (i.e. $Y Y^T$) and Gaussian kernel.

2.2 AC-PCA with sparse loading

A sparse solution for v can be achieved by adding ℓ_1 constraint:

$$\underset{v \in \mathbb{R}^p}{\text{maximize}} v^T X^T X v \quad \text{subject to } v^T X^T K X v \leq c_1, \|v\|_1 \leq c_2, \|v\|_2^2 \leq 1. \quad (2)$$

3 Specific experimental designs and case studies

3.1 Categorical confounding factors

Categorical confounding factors are commonly observed in biological data, it can be technical (different experimental batches, etc.) and biological (donor labels, different races, species, etc.). we treat samples with the same confounder label as a group. Here are the group labels for simulated example 1:

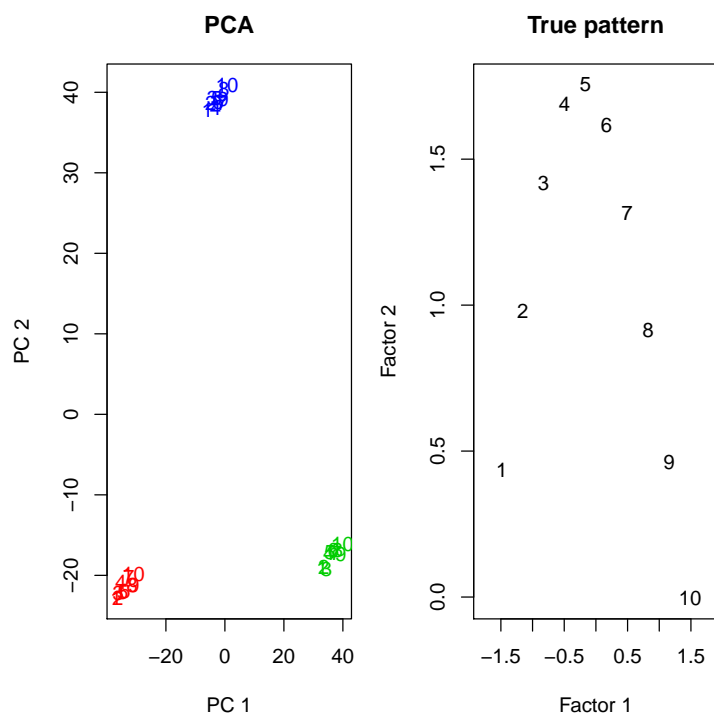
```
> library(acPCA)
> data(data_example1)
> data_example1$group ### the group labels

[1] 1 1 1 1 1 1 1 1 1 1 2 2 2 2 2 2 2 2 2 2 3 3 3 3 3 3 3 3 3 3

> X <- data_example1$X ### the data matrix
```

we perform regular PCA and compare the result with the true simulated pattern:

```
> pca <- prcomp(X, center=T) ###regular PCA
> par(mfrow=c(1,2), pin=c(2.5,2.5), mar=c(4.1, 3.9, 3.2, 1.1))
> plot(pca$x[,1], pca$x[,2], xlab="PC 1", ylab="PC 2", col="black", type="n", main="PCA")
> text(pca$x[,1], pca$x[,2], labels = data_example1$lab, col=data_example1$group+1)
> plot(data_example1$true_pattern[1,], data_example1$true_pattern[2,],
+       xlab="Factor 1", ylab="Factor 2", col="black", type="n", main="True pattern",
+       xlim=c(min(data_example1$true_pattern[1,])-0.3,
+               max(data_example1$true_pattern[1,])+0.3) )
> text(data_example1$true_pattern[1,], data_example1$true_pattern[2,], labels = 1:10)
```



In the PCA plot, each color represents a group of samples with the same confounder label. The confounding variation dominates variation of the true pattern.

To adjust for the confounding variation, the Y matrix is designed such that the penalty term equals the between groups sum of squares:

```
> Y <- data_example1$Y ### the confounder matrix
> Y
```

	[,1]	[,2]	[,3]
[1,]	0.6666667	-0.3333333	-0.3333333
[2,]	0.6666667	-0.3333333	-0.3333333
[3,]	0.6666667	-0.3333333	-0.3333333
[4,]	0.6666667	-0.3333333	-0.3333333
[5,]	0.6666667	-0.3333333	-0.3333333
[6,]	0.6666667	-0.3333333	-0.3333333
[7,]	0.6666667	-0.3333333	-0.3333333
[8,]	0.6666667	-0.3333333	-0.3333333
[9,]	0.6666667	-0.3333333	-0.3333333
[10,]	0.6666667	-0.3333333	-0.3333333
[11,]	-0.3333333	0.6666667	-0.3333333
[12,]	-0.3333333	0.6666667	-0.3333333

```

[13,] -0.3333333  0.6666667 -0.3333333
[14,] -0.3333333  0.6666667 -0.3333333
[15,] -0.3333333  0.6666667 -0.3333333
[16,] -0.3333333  0.6666667 -0.3333333
[17,] -0.3333333  0.6666667 -0.3333333
[18,] -0.3333333  0.6666667 -0.3333333
[19,] -0.3333333  0.6666667 -0.3333333
[20,] -0.3333333  0.6666667 -0.3333333
[21,] -0.3333333 -0.3333333  0.6666667
[22,] -0.3333333 -0.3333333  0.6666667
[23,] -0.3333333 -0.3333333  0.6666667
[24,] -0.3333333 -0.3333333  0.6666667
[25,] -0.3333333 -0.3333333  0.6666667
[26,] -0.3333333 -0.3333333  0.6666667
[27,] -0.3333333 -0.3333333  0.6666667
[28,] -0.3333333 -0.3333333  0.6666667
[29,] -0.3333333 -0.3333333  0.6666667
[30,] -0.3333333 -0.3333333  0.6666667

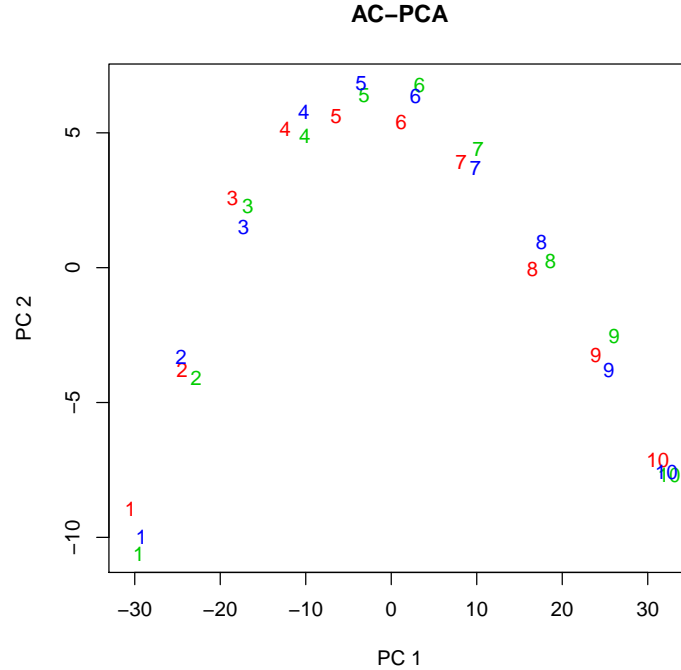
```

Here are the results when we implement AC-PCA:

```

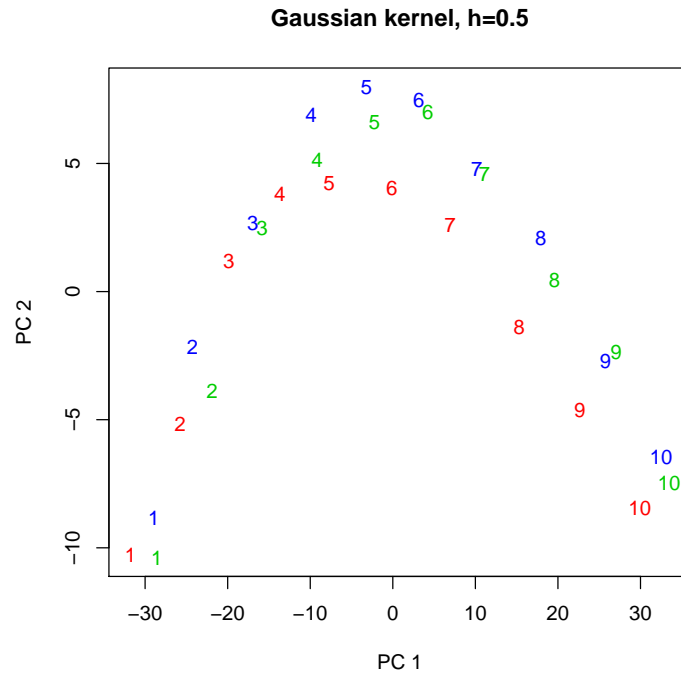
> par(mfrow=c(1,1))
> ### first tune lambda
> result_cv <- acPCAtuneLambda(X=X, Y=Y, nPC=2, lambdas=seq(0, 10, 0.05),
+                               anov=T, kernel = "linear", quiet=T)
> ###run with the best lambda
> result <- acPCA(X=X, Y=Y, lambda=result_cv$best_lambda,
+                 kernel="linear", nPC=2)
> ###the signs of the PCs are meaningless
> plot(result$Xv[,1], -result$Xv[,2], xlab="PC 1", ylab="PC 2",
+       type="n", main="AC-PCA")
> text(result$Xv[,1], -result$Xv[,2], labels = data_example1$lab,
+       col=data_example1$group+1)

```



We set $anov = T$ as the penalty term has the ANOVA interpretation. AC-PCA is able to recover the true latent structure. We can also implement AC-PCA with Gaussian kernel (bandwidth $h = 0.5$):

```
> h <- 0.5
> result_cv <- acPCAtuneLambda(X=X, Y=Y, nPC=2, lambdas=seq(0, 10, 0.05),
+                             anov=F, kernel = "gaussian",
+                             bandwidth=h, quiet=T)
> result <- acPCA(X=X, Y=Y, lambda=result_cv$best_lambda,
+                kernel="gaussian", bandwidth=h, nPC=2)
> ###the signs of the PCs are meaningless
> plot(result$Xv[,1], -result$Xv[,2], xlab="PC 1", ylab="PC 2",
+      col="black", type="n", main="Gaussian kernel, h=0.5")
> text(result$Xv[,1], -result$Xv[,2], labels = data_example1$lab,
+      col=data_example1$group+1)
```



Gaussian kernel also works well. For detailed discussion on bandwidth selection, please refer to Chapter 3 in [4].

3.2 Continuous confounding factors

Continuous confounding factors may be present in biological data, for example, age.

Simulated example 2: the confounder is assumed to be continuous and it contributes a global trend to the data.

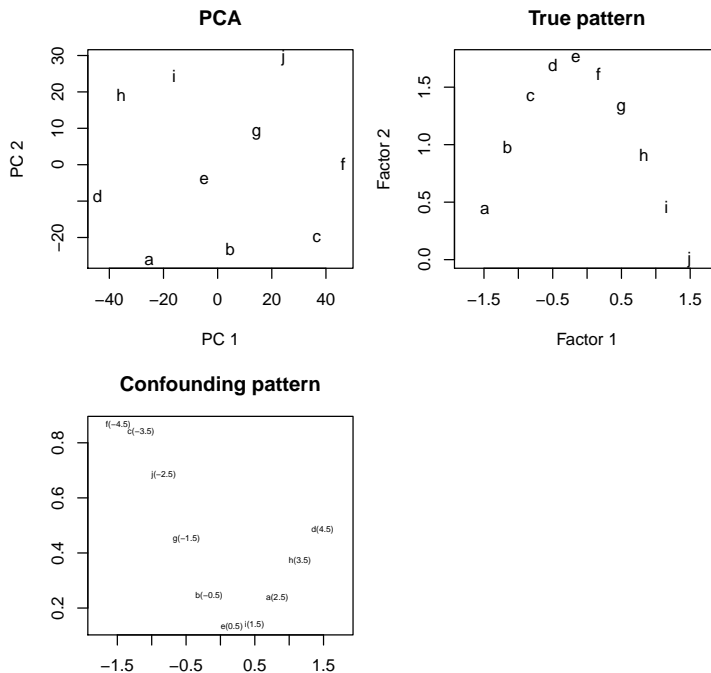
We first perform regular PCA:

```
> data(data_example2)
> X <- data_example2$X ###the data matrix
> Y <- data_example2$Y ###the confounder matrix
> Y
```

```
      [,1]
[1,]  2.5
[2,] -0.5
[3,] -3.5
[4,]  4.5
[5,]  0.5
[6,] -4.5
```

```
[7,] -1.5
[8,]  3.5
[9,]  1.5
[10,] -2.5
```

```
> pca <- prcomp(X, center=T) ###regular PCA
> par(mfrow=c(2,2), pin=c(2.5,2.5), mar=c(4.1, 3.9, 3.2, 1.1))
> plot(pca$x[,1], pca$x[,2], xlab="PC 1", ylab="PC 2", col="black", type="n", main="PCA")
> text(pca$x[,1], pca$x[,2], labels = data_example2$lab)
> plot(data_example2$true_pattern[1,], data_example2$true_pattern[2,],
+       xlab="Factor 1", ylab="Factor 2", col="black", type="n", main="True pattern",
+       xlim=c(min(data_example2$true_pattern[1,])-0.3,
+               max(data_example2$true_pattern[1,])+0.3) )
> text(data_example2$true_pattern[1,], data_example2$true_pattern[2,],
+       labels = data_example2$lab)
> plot(data_example2$confound_pattern[1,], data_example2$confound_pattern[2,],
+       xlab="", ylab="", col="black", type="n", main="Confounding pattern",
+       xlim=c(min(data_example2$confound_pattern[1,])-0.3,
+               max(data_example2$confound_pattern[1,])+0.3) )
> text(data_example2$confound_pattern[1,], data_example2$confound_pattern[2,], labels =
+       paste(data_example2$lab, '(', data_example2$Y, ')', sep=""), cex=0.5 )
```

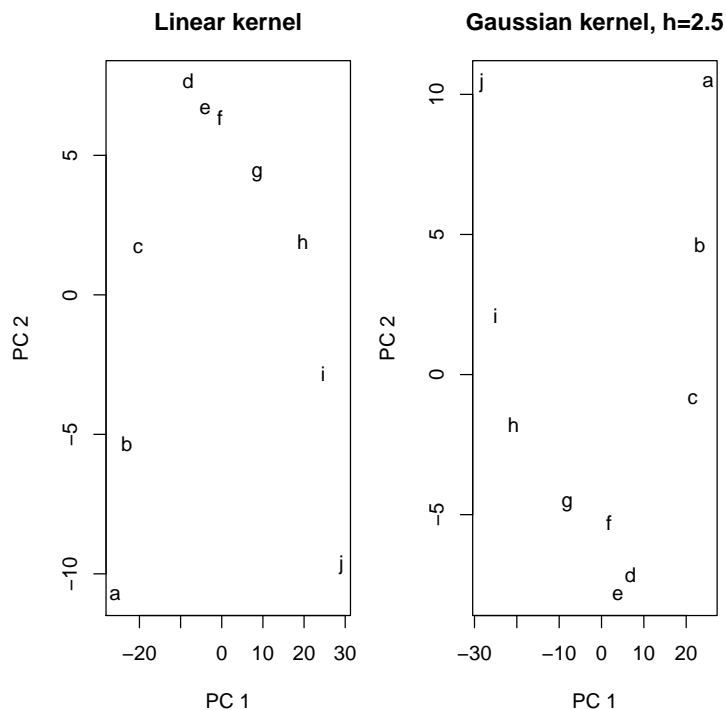


Next, we implement AC-PCA with linear and Gaussian kernels:


```

> ### linear kernel
> result_cv <- acPCAtuneLambda(X=X, Y=Y, nPC=2, lambdas=seq(0, 10, 0.05),
+                             anov=F, kernel = "linear", quiet=T)
> result <- acPCA(X=X, Y=Y, lambda=result_cv$best_lambda,
+                 kernel="linear", nPC=2)
> par(mfrow=c(1,2), pin=c(2.5,2.5), mar=c(4.1, 3.9, 3.2, 1.1))
> plot(result$Xv[,1], result$Xv[,2], xlab="PC 1", ylab="PC 2",
+       col="black", type="n", main="Linear kernel")
> text(result$Xv[,1], result$Xv[,2], labels = data_example2$lab)
> ### Gaussian kernel
> result_cv <- acPCAtuneLambda(X=X, Y=Y, nPC=2, lambdas=seq(0, 10, 0.05),
+                             anov=F, kernel = "gaussian",
+                             bandwidth=mean(abs(Y)), quiet=T)
> result <- acPCA(X=X, Y=Y, lambda=result_cv$best_lambda,
+                 kernel = "gaussian", bandwidth=mean(abs(Y)), nPC=2)
> plot(result$Xv[,1], result$Xv[,2], xlab="PC 1", ylab="PC 2",
+       col="black", type="n", main=paste("Gaussian kernel, h=", mean(abs(Y)), sep="" )
> text(result$Xv[,1], result$Xv[,2], labels = data_example2$lab)

```



The results of linear and Gaussian kernels are comparable.

3.3 Experiments with replicates

Consider an experiment where the gene expression levels are measured under multiple biological conditions with several replicates. It may be desirable to capture the variation across biological conditions but shared among replicates. Simulated example 3: there are 10 biological conditions, each with $n = 3$ replicates. The variation is shared among replicates for half of the genes and not shared for the other genes. Suppose we want to capture the variation shared among replicates, the confounder matrix can be chosen such that samples within the same biological condition are “pushed” together:

```
> data(data_example3)
> X <- data_example3$X ### the data matrix
> data_example3$lab ### the biological conditions

[1] 1 1 1 2 2 2 3 3 3 4 4 4 5 5 5 6 6 6 7 7 7 8 8 8 9
[26] 9 9 10 10 10

> Y <- data_example3$Y ### the confounder matrix
> dim(Y)

[1] 30 30

> Y[,1]

[1] 0.5773503 -0.5773503 0.0000000 0.0000000 0.0000000 0.0000000
[7] 0.0000000 0.0000000 0.0000000 0.0000000 0.0000000 0.0000000
[13] 0.0000000 0.0000000 0.0000000 0.0000000 0.0000000 0.0000000
[19] 0.0000000 0.0000000 0.0000000 0.0000000 0.0000000 0.0000000
[25] 0.0000000 0.0000000 0.0000000 0.0000000 0.0000000 0.0000000
```

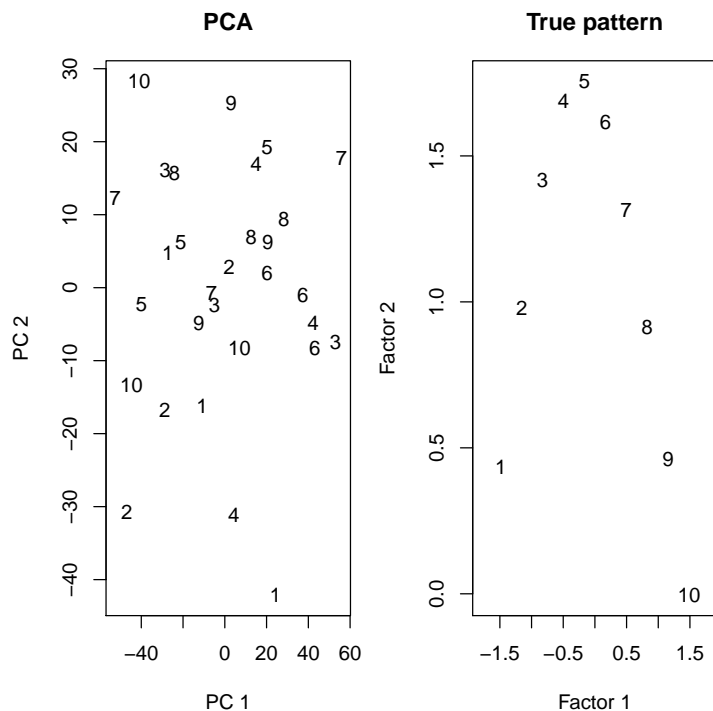
In each column of Y , there is one $1/\sqrt{n}$ and one $-1/\sqrt{n}$ corresponding to a pair of samples from the same biological condition. So the number of columns in Y is $10 \times 3 \times 2/2$. For a biological condition, treating the 3 replicates as 3 groups, it can be shown that the penalty term $v^T X^T Y Y^T X v$ equals the summation of the between groups sum of squares over the biological conditions. When the number of replicates across the biological conditions is not the same, to maintain the ANOVA interpretation, we can change the corresponding entries in Y to $1/\sqrt{n_i}$ and $-1/\sqrt{n_i}$, where n_i is the number of replicates in the i th biological condition. We first implement PCA and compare it with the true pattern:

```
> pca <- prcomp(X, center=T) ###regular PCA
> par(mfrow=c(1,2), pin=c(2.5,2.5), mar=c(4.1, 3.9, 3.2, 1.1))
> plot(pca$x[,1], pca$x[,2], xlab="PC 1", ylab="PC 2", col="black", type="n", main="PCA")
> text(pca$x[,1], pca$x[,2], labels = data_example3$lab)
> plot(data_example3$true_pattern[1,], data_example3$true_pattern[2,],
+       xlab="Factor 1", ylab="Factor 2", col="black", type="n", main="True pattern",
+       xlim=c(min(data_example3$true_pattern[1,])-0.3,
```

```

+           max(data_example3$true_pattern[1,])+0.3) )
> text(data_example3$true_pattern[1,], data_example3$true_pattern[2,],
+       labels = 1:10)

```

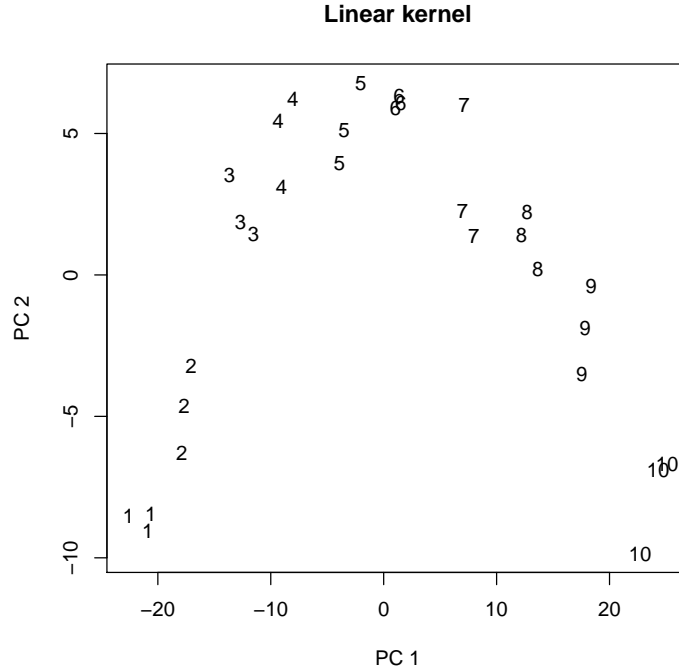


Next we implement AC-PCA:

```

> result_cv <- acPCAtuneLambda(X=X, Y=Y, nPC=2, lambdas=seq(0, 10, 0.05),
+                             anov=T, kernel = "linear", quiet=T)
> result <- acPCA(X=X, Y=Y, lambda=result_cv$best_lambda,
+                kernel="linear", nPC=2)
> par(mfrow=c(1,1))
> plot(-result$Xv[,1], result$Xv[,2], xlab="PC 1", ylab="PC 2",
+      col="black", type="n", main="Linear kernel")
> text(-result$Xv[,1], result$Xv[,2], labels = data_example3$lab)

```



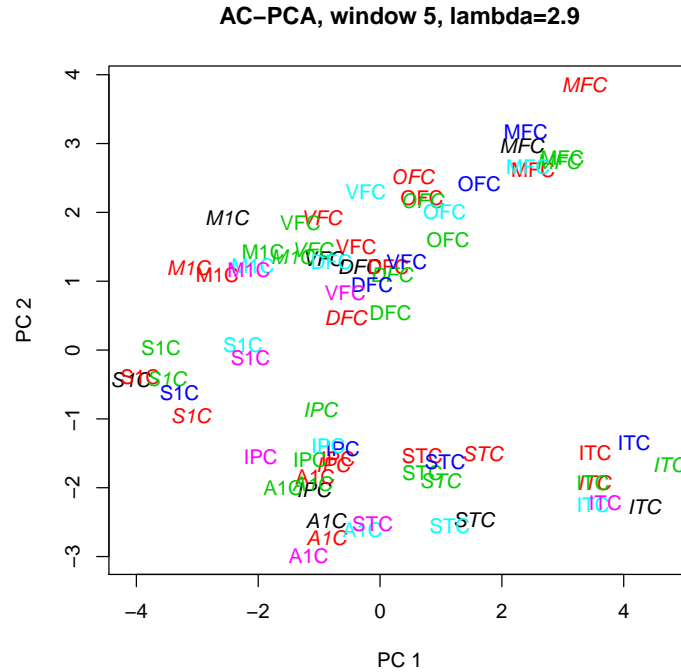
3.4 Application to a human brain exon array dataset

We analyze the human brain exon array dataset [3]. The dataset was downloaded from the Gene Expression Omnibus (GEO) database under the accession number GSE25219. The dataset was generated from 1,340 tissue samples collected from 57 developing and adult post-mortem brains. In the dataset, gene expression levels were measured in different brain regions in multiple time windows. We use a subset of 1,000 genes for demonstration purpose. For time window 5, we first perform PCA to visualize the brain regions:

```
> data(data_brain_w5)
> X <- data_brain_w5$X;
> table(data_brain_w5$hemispheres) ###1: left hemisphere, 3: right hemisphere

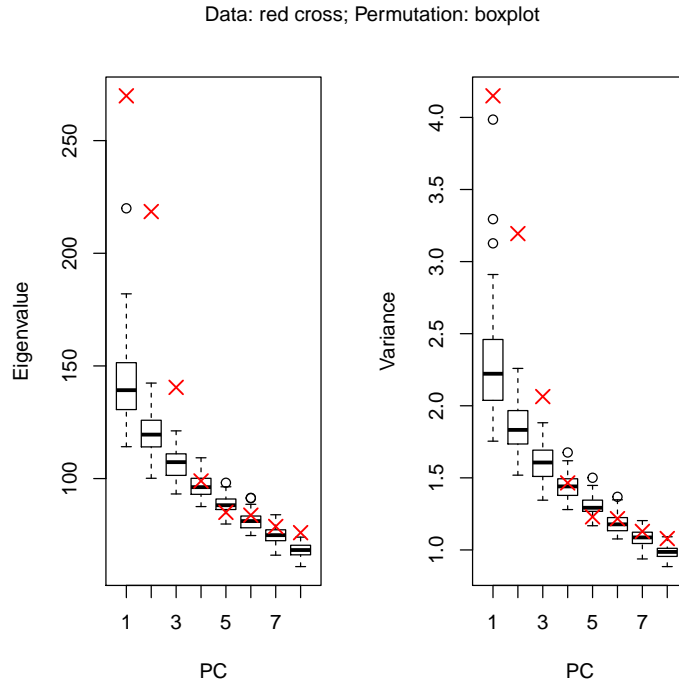
1 3
45 29

> pca <- prcomp(X, center=T) ###regular PCA
> plot(pca$x[,1], pca$x[,2], xlab="PC 1", ylab="PC 2",
+      col="black", type="n", main="PCA, window 5, brain data")
> text(pca$x[,1], pca$x[,2], labels = data_brain_w5$regions,
+      col=data_brain_w5$donor_labs, font=data_brain_w5$hemispheres)
```

In the manuscript, we used $\lambda = 5$:

```
> result2 <- acPCA(X=X, Y=Y, lambda=5, kernel="linear", nPC=2)
> Xv2 <- result2$Xv
> plot(-Xv2[,1], Xv2[,2], xlab="PC 1", ylab="PC 2",
+      col="black", type="n", main="AC-PCA, window 5, lambda=5")
> text(-Xv2[,1], Xv2[,2], labels = data_brain_w5$regions,
+      col=data_brain_w5$donor_labs, font=data_brain_w5$hemispheres)
```

The first 3 PCs are significant.

3.5 Application to a model organism ENCODE (modENCODE) RNA-Seq dataset

The modENCODE project generates the transcriptional landscapes for model organisms during development[1, 2]. In the analysis, we used the time-course RNA-Seq data for fly and worm embryonic development. The modENCODE RNA-Seq dataset was downloaded from <https://www.encodeproject.org/comparative/transcriptome/>. For the orthologs in fly that map to multiple orthologs in worm, we took median to get a one to one match, resulting in 4831 ortholog paris. We first perform some exploratory analysis:

```
> library(acPCA)
> data(data_fly_worm)
> data_fly <- data_fly_worm$data_fly ###the data matrix for fly
> data_worm <- data_fly_worm$data_worm ###the data matrix for fly
> dim(data_fly)

[1] 12 4831

> dim(data_worm)

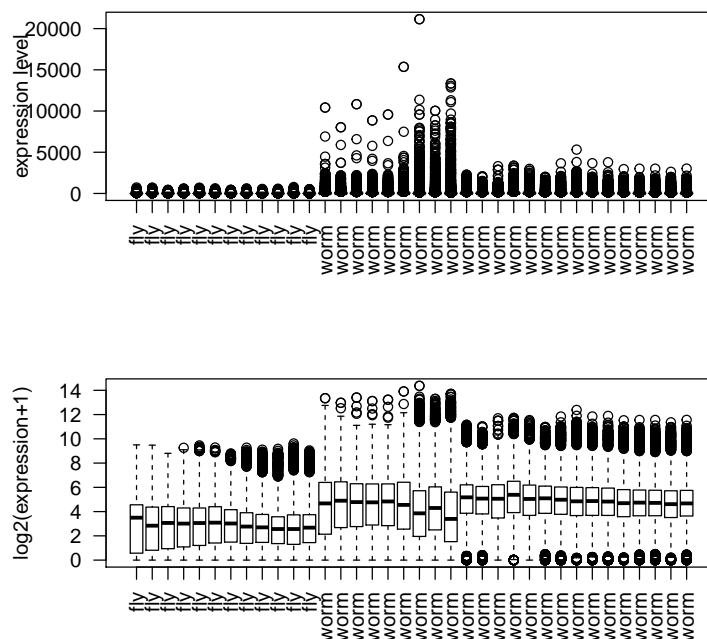
[1] 24 4831
```



```

> par(mfrow=c(2,1), pin=c(2.5,2.5), mar=c(4.1, 3.9, 3.2, 1.1))
> boxplot(t(rbind(data_fly, data_worm)),las = 2, names =
+         c(rep('fly', 12), rep('worm', 24)), ylab="expression level")
> boxplot(log2(t(rbind(data_fly, data_worm))+1),las = 2,
+         names = c(rep('fly', 12), rep('worm', 24)), ylab="log2(expression+1)")

```

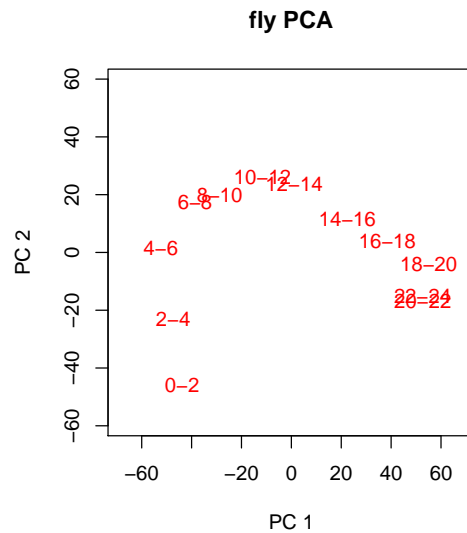


We can clearly see that the expression levels in worm samples tend to be higher. Next we perform PCA on fly and worm separately on the log2 scale:

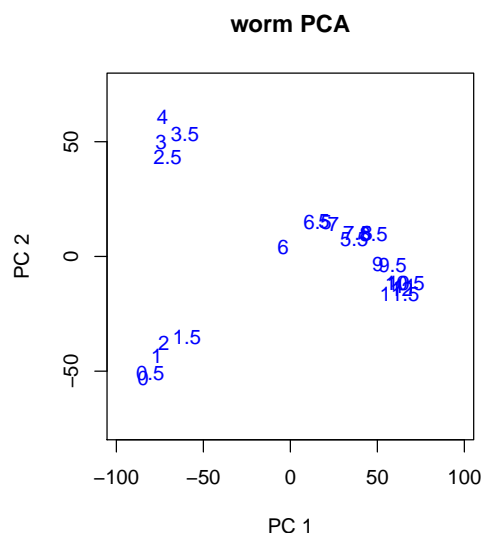
```

> pca_fly <- prcomp(log2(data_fly+1), center=T)
> pca_worm <- prcomp(log2(data_worm+1), center=T)
> par(mfrow=c(1,1), pin=c(3, 3))
> plot(pca_fly$x[,1], -pca_fly$x[,2], xlab="PC 1", ylab="PC 2",
+      col="red", type="n", main="fly PCA",
+      xlim=c(-(max(abs(pca_fly$x[,1]))+13), max(abs(pca_fly$x[,1]))+13),
+      ylim=c(-(max(abs(pca_fly$x[,2]))+13), max(abs(pca_fly$x[,2]))+13))
> text(pca_fly$x[,1], -pca_fly$x[,2], labels =
+      data_fly_worm$fly_time, col="red")

```

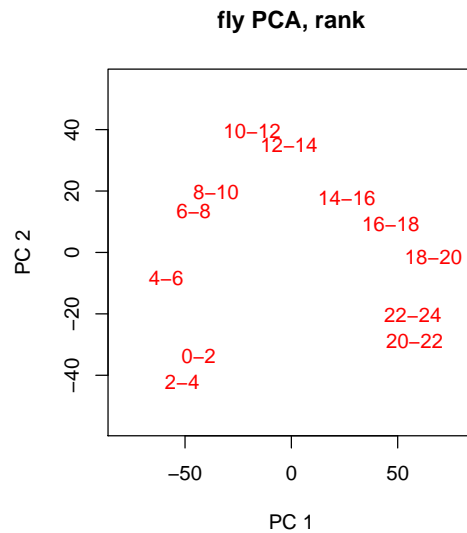


```
> par(mfrow=c(1,1), pin=c(3, 3))
> plot(pca_worm$x[,1], pca_worm$x[,2], xlab="PC 1", ylab="PC 2",
+      col="red", type="n", main="worm PCA",
+      xlim=c(-(max(abs(pca_worm$x[,1]))+13), max(abs(pca_worm$x[,1]))+13),
+      ylim=c(-(max(abs(pca_worm$x[,2]))+13), max(abs(pca_worm$x[,2]))+13))
> text(pca_worm$x[,1], pca_worm$x[,2], labels =
+      data_fly_worm$worm_time, col="blue")
```

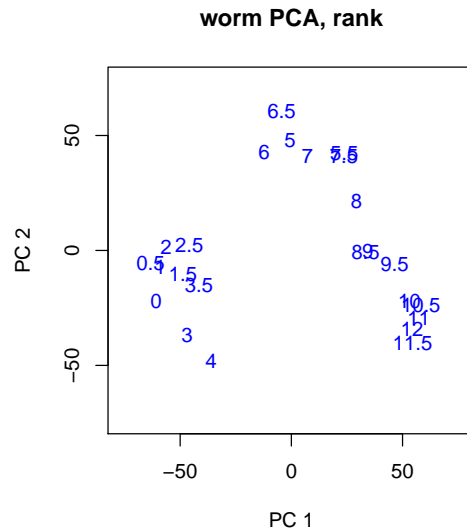


In the PCA plots, each data point represents a time window (fly) or a time point (worm), in the unit of hours(h). For fly, there is a smooth temporal pattern. For worm, there seems to be three clusters, which may originate from the dominated effect of a small subset of highly expressed genes. To adjust for the highly expressed genes, we used the rank across samples within the same species. The rank matrix was then scaled to have unit variance. Another benefit of using rank is that we simultaneously make adjustment for the observation that worm genes tend to have higher expression levels. For ties in the expression level, we used 'average' option in the R's rank function. Here is PCA on the rank matrix:

```
> X <- data_fly_worm$X ####this is the scaled rank matrix. Fly and worm are combined.
> pca_fly_rank <- prcomp(X[1:12,], center=T)
> pca_worm_rank <- prcomp(X[13:36,], center=T)
> par(mfrow=c(1,1), pin=c(3, 3))
> plot(pca_fly_rank$x[,1], -pca_fly_rank$x[,2], xlab="PC 1", ylab="PC 2",
+      col="red", type="n", main="fly PCA, rank",
+      xlim=c(-(max(abs(pca_fly_rank$x[,1]))+13), max(abs(pca_fly_rank$x[,1]))+13),
+      ylim=c(-(max(abs(pca_fly_rank$x[,2]))+13), max(abs(pca_fly_rank$x[,2]))+13))
> text(pca_fly_rank$x[,1], -pca_fly_rank$x[,2], labels =
+      data_fly_worm$fly_time, col="red")
```

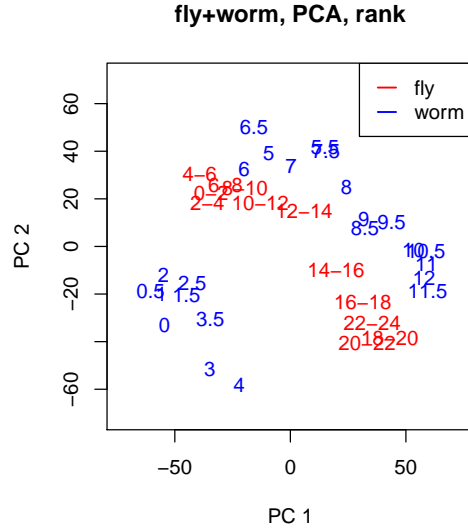


```
> par(mfrow=c(1,1), pin=c(3, 3))
> plot(pca_worm_rank$x[,1], pca_worm_rank$x[,2], xlab="PC 1", ylab="PC 2",
+      col="red", type="n", main="worm PCA, rank",
+      xlim=c(-(max(abs(pca_worm_rank$x[,1]))+13), max(abs(pca_worm_rank$x[,1]))+13),
+      ylim=c(-(max(abs(pca_worm_rank$x[,2]))+13), max(abs(pca_worm_rank$x[,2]))+13))
> text(pca_worm_rank$x[,1], pca_worm_rank$x[,2], labels =
+      data_fly_worm$worm_time, col="blue")
```



Using rank gives a better visualization, as the data points are more spread out. Next we implement PCA on fly and worm jointly, using the rank matrix:

```
> pca_rank <- prcomp(X, center=T)
> par(mfrow=c(1,1), pin=c(3, 3))
> plot(pca_rank$x[,1], pca_rank$x[,2], xlab="PC 1", ylab="PC 2",
+      col="red", type="n", main="fly+worm, PCA, rank",
+      xlim=c(-(max(abs(pca_rank$x[,1]))+13), max(abs(pca_rank$x[,1]))+13),
+      ylim=c(-(max(abs(pca_rank$x[,2]))+13), max(abs(pca_rank$x[,2]))+13))
> text(pca_rank$x[,1], pca_rank$x[,2],
+      labels=data_fly_worm$X_time,
+      col=c(rep("red", 12), rep("blue", 24)) )
> legend("topright", legend = c("fly", "worm"), lty=1,
+      col=c("red", "blue"), lwd=1.5, seg.len=1)
```



The variation of species confounds the PCA result, as we observed that samples from the same species tend to be close.

Next we implement AC-PCA. Without prior knowledge on the alignment of the developmental stages in fly and worm, we shrink the data points in fly towards the mean of the data points in worm: for example, time window 0-2h in fly is shrunk to the mean of time points 0h, 0.5h and 1h in worm, as the embryonic stage for fly is 24h, and for worm it is 12h. Time window 8-10h in fly is shrunk to the mean of 4h and 5h in worm, as the worm sample 4.5h is missing.

```
> Y <- data_fly_worm$Y
> Y[,1]

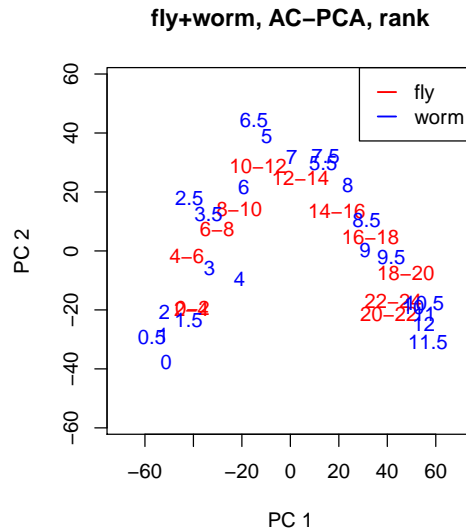
[1] 1.0000000 0.0000000 0.0000000 0.0000000 0.0000000 0.0000000
[7] 0.0000000 0.0000000 0.0000000 0.0000000 0.0000000 0.0000000
[13] -0.3333333 -0.3333333 -0.3333333 0.0000000 0.0000000 0.0000000
[19] 0.0000000 0.0000000 0.0000000 0.0000000 0.0000000 0.0000000
[25] 0.0000000 0.0000000 0.0000000 0.0000000 0.0000000 0.0000000
[31] 0.0000000 0.0000000 0.0000000 0.0000000 0.0000000 0.0000000

> result_cv <- acPCAtuneLambda(X=X, Y=Y, nPC=2, lambdas=seq(0, 10, 0.05),
+                               anov=F, kernel = "linear", quiet=T)
> result <- acPCA(X=X, Y=Y, lambda=result_cv$best_lambda,
```

```

+           kernel="linear", nPC=2)
> par(mfrow=c(1,1), pin=c(3, 3))
> plot(-result$Xv[,1], result$Xv[,2], xlab="PC 1", ylab="PC 2",
+      col="red", type="n", main="fly+worm, AC-PCA, rank",
+      xlim=c(-(max(abs(result$Xv[,1]))+13), max(abs(result$Xv[,1]))+13),
+      ylim=c(-(max(abs(result$Xv[,2]))+13), max(abs(result$Xv[,2]))+13))
> text(-result$Xv[,1], result$Xv[,2],
+      labels=data_fly_worm$X_time,
+      col=c(rep("red", 12), rep("blue", 24)) )
> legend("topright", legend = c("fly", "worm"),
+      lty=1, col=c("red","blue"),lwd=1.5,seg.len=1)

```



The variation of different species is adjusted. The genes with top loadings in AC-PCA tend to have consistent temporal pattern in fly and worm (see manuscript).

3.6 Some tips for implementation in RNA-Seq data

PCA generally works for “normal-like” data. For data that are far from “normal”, the result may be driven by genes with large variation.

Figure 2 compares the distribution of gene expression levels between RNA-Seq and microarray experiments for the same biological sample. Log transformation of RPKM makes the data more “normal-like”, but may not be sufficient

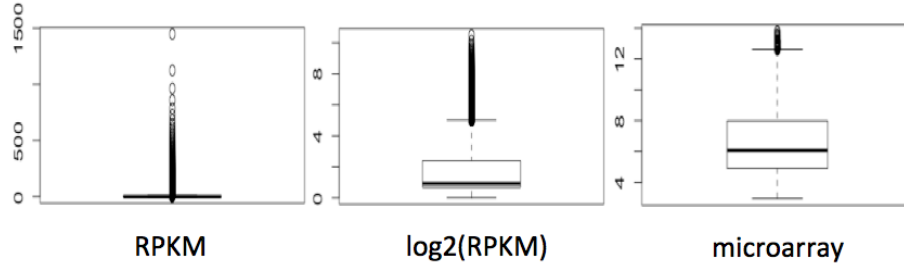


Figure 2: Boxplots comparing RNA-Seq and microarray, same biological sample

for some datasets. The highly expressed “outlier” genes may still affect the result. One way to get around this is to use some robust statistics, such as rank: (a) Rank among samples: for each gene, rank the expression levels among samples, then the expression levels will be 1,...,number of samples. We used this strategy for the modENCODE RNA-Seq data.

(b) Rank among genes: in each sample, rank the gene expression levels among all genes, then the expression level will be 1,...,number of genes.

(a) makes the distribution for every gene to be the same and (b) makes the distribution for every sample to be the same. Based on our experience, both approaches seem to work well for visualization purpose.

3.7 AC-PCA with sparsity

For AC-PCA with sparsity constraints, there are two tuning parameters, c_1 and c_2 : c_1 controls the strength of the penalty, and c_2 controls the sparsity of v . The parameters are tuned sequentially: c_1 is tuned without the sparsity constraint, and then c_2 is tuned with c_1 fixed. We implement the procedure on the brain exon array data, time window 2:

```
> set.seed(10)
> data(data_brain_w2)
> X <- data_brain_w2$X ###the data matrix
> Y <- data_brain_w2$Y
> result1cv <- acPCAtuneLambda(X=X, Y=Y, nPC=2, lambdas=seq(0, 20, 0.1),
+                               anov=T, kernel = "linear", quiet=T)
> result1 <- acPCA(X=X, Y=Y, lambda=result1cv$best_lambda, kernel="linear", nPC=2)
> v_ini <- as.matrix(result1$v[,1])
> par(mfrow=c(1,2), pin=c(2.5,2.5), mar=c(4.1, 3.9, 3.2, 1.1))
> c2s <- seq(1, 0, -0.1)*sum(abs(v_ini))
> resultcv_spc1_coarse <- acSPCcv( X=X, Y=Y, c2s=c2s, v_ini=v_ini,
+                               kernel="linear", quiet=T, fold=10)
> c2s <- seq(0.9, 0.7, -0.02)*sum(abs(v_ini))
> resultcv_spc1_fine <- acSPCcv( X=X, Y=Y, c2s=c2s, v_ini=v_ini,
```

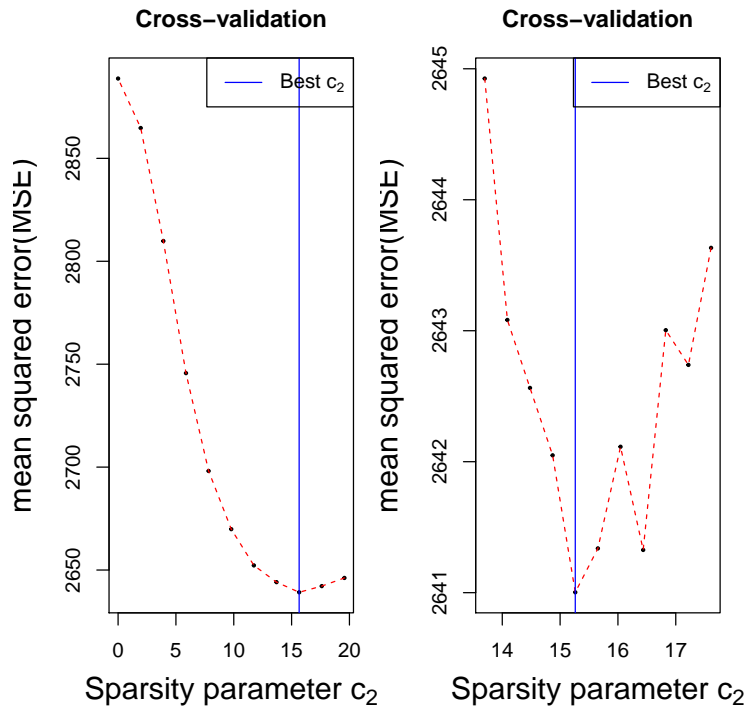


```

+                                     kernel="linear", quiet=T, fold=10)
> result_spc1 <- acSPC( X=X, Y=Y, c2=resultcv_spc1_fine$best_c2,
+                       v_ini=v_ini, kernel="linear")
> v1 <- result_spc1$v
> sum(v1!=0)

```

[1] 736



To speed-up the computational time, a coarse search for c_2 was performed first, followed by a finer grid.

Multiple sparse principal components can be obtained by subtracting the first several principal components, and then implement the algorithm:

```

> v_substract <- as.matrix(result1$v[,1])
> v_ini <- as.matrix(result1$v[,2])
> par(mfrow=c(1,2), pin=c(2.5,2.5), mar=c(4.1, 3.9, 3.2, 1.1))
> c2s <- seq(1, 0, -0.1)*sum(abs(v_ini))
> resultcv_spc2_coarse <- acSPCcv(X=X, Y=Y, c2s=c2s, v_ini=v_ini,
+                                v_substract=v_substract, kernel="linear",
+                                quiet=T, fold=10)
> c2s <- seq(0.5, 0.2, -0.02)*sum(abs(v_ini))
> resultcv_spc2_fine <- acSPCcv(X=X, Y=Y, c2s=c2s, v_ini=v_ini,
+                                v_substract=v_substract, kernel="linear",
+

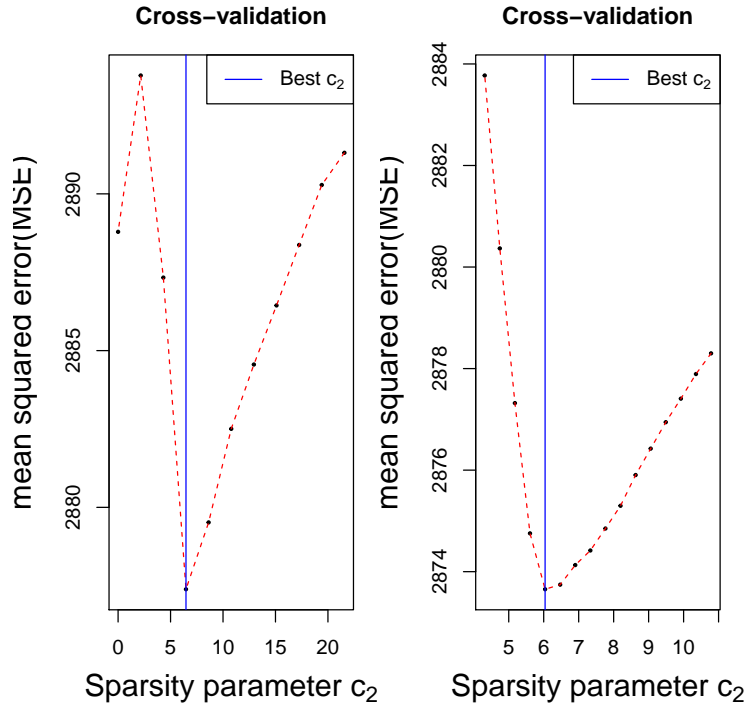
```

```

+                               quiet=T, fold=10)
> result_spc2 <- acSPC(X=X, Y=Y, c2=resultcv_spc2_fine$best_c2,
+                     v_ini=v_ini, v_substract=v_substract, kernel="linear")
> v2 <- result_spc2$v
> sum(v2!=0)

[1] 68

```



In the example, the non-sparse first PC was subtracted. We can also subtract the sparse PC:

```
> v_substract <- v1
```

References

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- [2] Mark B Gerstein, Joel Rozowsky, Koon-Kiu Yan, Daifeng Wang, Chao Cheng, James B Brown, Carrie A Davis, LaDeana Hillier, Cristina Sisú, Jingyi Jessica Li, et al. Comparative analysis of the transcriptome across distant species. *Nature*, 512(7515):445–448, 2014.

- [3] Hyo Jung Kang, Yuka Imamura Kawasaki, Feng Cheng, Ying Zhu, Xuming Xu, Mingfeng Li, André MM Sousa, Mihovil Pletikos, Kyle A Meyer, Goran Sedmak, et al. Spatio-temporal transcriptome of the human brain. *Nature*, 478(7370):483–489, 2011.
- [4] Matt P Wand and M Chris Jones. *Kernel smoothing*. Crc Press, 1994.