QCB 508 – Week 12

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EDA of HD Data

Rationale

Exploratory data analysis (EDA) of high-dimensional data adds the additional challenge that many variables must be examined simultaneously. Therefore, in addition to the EDA methods we discussed earlier, methods are often employed to organize, visualize, or numerically capture high-dimensional data into lower dimensions.

Examples of EDA approaches applied to HD data include:

- Traditional EDA methods covered earlier
- Cluster analysis
- Dimensionality reduction

Cluster Analysis

An overview of common cluster analysis methods can be found here:

http://sml201.github.io/lectures/week12/week12.html

These slides include:

- Distance measures
- Hierarchical clustering
- K-means clustering

Example: Cancer Subtypes

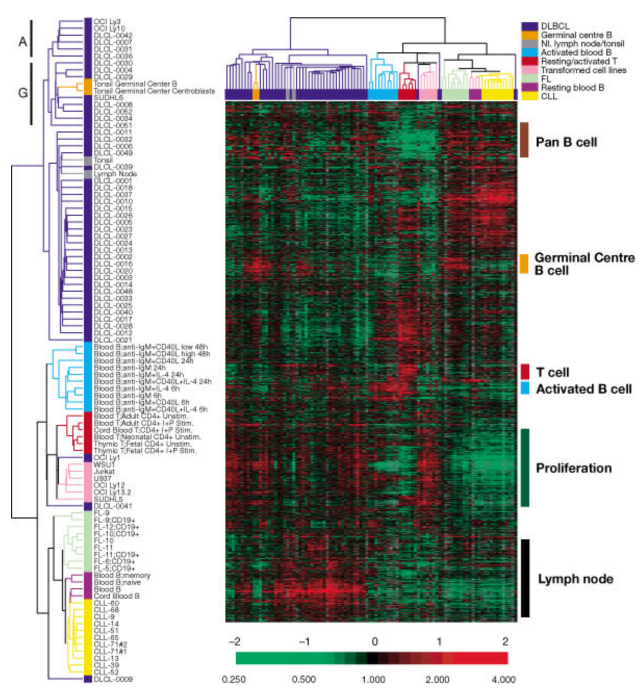


Figure from Alizadeh et al. (2000) Nature.

Dimensionality Reduction

The goal of **dimensionality reduction** is to extract low dimensional representations of high dimensional data that are useful for visualization, exploration, inference, or prediction.

The low dimensional representations should capture key sources of variation in the data.

Some Methods

- Principal component analysis
- Singular value decomposition
- Latent variable modeling
- Vector quantization
- Self-organizing maps
- Multidimensional scaling

Example: Weather Data

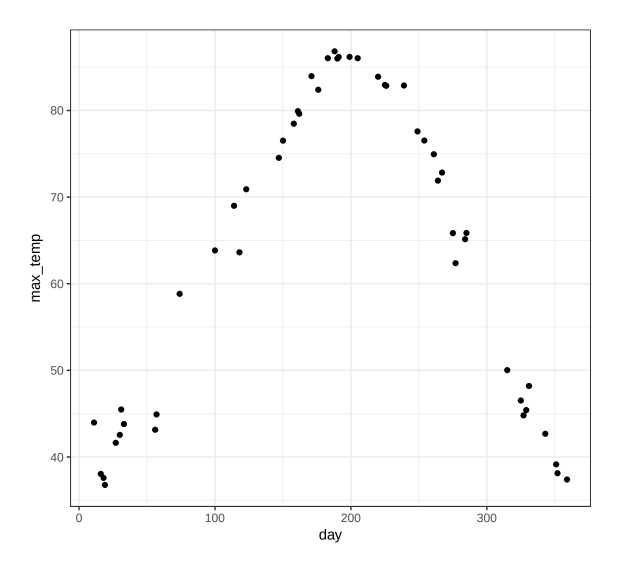
These daily temperature data (in tenths of degrees C) come from meteorogical observations for weather stations in the US for the year 2012 provided by NOAA (National Oceanic and Atmospheric Administration).:

This matrix contains temperature data on 50 days and 2811 stations that were randomly selected.

Convert temperatures to Fahrenheit:

```
> weather_data <- 0.18*weather_data + 32
> weather_data[1:5, 1:6]
                                  18
                                           19
                                                 27
                                                        30
                  11
                           16
AG000060611 56.84000 63.50000 63.14 61.52000 71.24 60.80
AGM00060369 60.44000 61.16000 59.72 60.62000 61.70 54.50
AGM00060425 81.09714 81.09714 59.36 61.34000 61.34 51.44
AGM00060444 55.04000 50.36000 50.00 51.98000 54.50 37.94
AGM00060468 50.90000 53.96000 49.46 79.44286 59.90 41.36
> apply(weather data, 1, median) %>%
    quantile(probs=seq(0,1,0.1))
        0%
                  10%
                              20%
                                         30%
                                                     40%
  8.886744
            49.010000
                       54.500000
                                   58.460000
                                              62.150000
       50%
                  60%
                              70%
                                         80%
                                                     90%
 65.930000
            69.679318
                       73.490000
                                   77.990000
                                              82.940000
      100%
140.000000
```

Here are the 2811 rows converted to a single row that captures the most variation among the rows:



Principal Component Analysis

Goal

For a given set of variables, **principal component analysis** (PCA) finds (constrained) weighted sums of the variables to produce variables (called principal components) that capture consectuive maximum levels of variation in the data.

Specifically, the first principal component is the weighted sum of the variables that results in a component with the highest variation.

This component is then "removed" from the data, and the second principal component is obtained on the resulting residuals.

This process is repeated until there is no variation left in the data.

Population PCA

Suppose we have m random variables X_1, X_2, \ldots, X_m . We wish to identify a set of weights w_1, w_2, \ldots, w_m that maximizes

$$\operatorname{Var}\left(w_{1}X_{1}+w_{2}X_{2}+\cdots+w_{m}X_{m}\right).$$

However, this is unbounded, so we need to constrain the weights. It turns out that constraining the weights so that

$$\|\boldsymbol{w}\|_2^2 = \sum_{i=1}^m w_i^2 = 1$$

is both interpretable and mathematically tractable.

Therefore we wish to maximize

$$Var(w_1X_1 + w_2X_2 + \cdots + w_mX_m)$$

subject to $\|\boldsymbol{w}\|_2^2 = 1$. Let Σ be the $m \times m$ population covariance matrix of the random variables X_1, X_2, \dots, X_m . It follows that

$$\operatorname{Var}(w_1X_1 + w_2X_2 + \dots + w_mX_m) = \boldsymbol{w}^T\boldsymbol{\Sigma}\boldsymbol{w}.$$

Using a Lagrange multiplier, we wish to maximize

$$\boldsymbol{w}^T \boldsymbol{\Sigma} \boldsymbol{w} + \lambda (\boldsymbol{w}^T \boldsymbol{w} - 1).$$

Differentiating with respect to w and setting to $\mathbf{0}$, we get $\Sigma w - \lambda w = 0$ or

$$\Sigma w = \lambda w$$
.

For any such \boldsymbol{w} and λ where this holds, note that

$$Var (w_1 X_1 + w_2 X_2 + \dots + w_m X_m) = \boldsymbol{w}^T \boldsymbol{\Sigma} \boldsymbol{w} = \lambda$$

so the variance is λ .

The eigendecompositon of a matrix identifies all such solutions to $\Sigma w = \lambda w$. Specifically, it calculates the decompositon

$$\Sigma = W \Lambda W^T$$

where \boldsymbol{W} is an $m \times m$ orthogonal matrix and $\boldsymbol{\Lambda}$ is a diagonal matrix with entries $\lambda_1 \geq \lambda_2 \geq \cdots \geq \lambda_m \geq 0$.

The fact that W is orthogonal means $WW^T = W^TW = I$.

The following therefore hold:

- For each column j_of \boldsymbol{W} , say \boldsymbol{w}_j , it follows that $\boldsymbol{\Sigma} \boldsymbol{w}_j = \lambda_j \boldsymbol{w}_j$
- $\|\boldsymbol{w}_j\|_2^2 = 1$ and $\boldsymbol{w}_j^T \boldsymbol{w}_k = \mathbf{0}$ for $\lambda_j \neq \lambda_k$
- $\operatorname{Var}(\boldsymbol{w}_{i}^{T}\boldsymbol{X}) = \lambda_{j}$

• $\operatorname{Var}(\boldsymbol{w}_{1}^{T}\boldsymbol{X}) \geq \operatorname{Var}(\boldsymbol{w}_{2}^{T}\boldsymbol{X}) \geq \cdots \geq \operatorname{Var}(\boldsymbol{w}_{m}^{T}\boldsymbol{X})$ • $\boldsymbol{\Sigma} = \sum_{j=1}^{m} \lambda_{j} \boldsymbol{w}^{j} \boldsymbol{w}_{j}^{T}$ • For $\lambda_{j} \neq \lambda_{k}$,

$$\operatorname{Cov}(\boldsymbol{w}_i^T \boldsymbol{X}, \boldsymbol{w}_k^T \boldsymbol{X}) = \boldsymbol{w}_i^T \boldsymbol{\Sigma} \boldsymbol{w}_k = \lambda_k \boldsymbol{w}_i^T \boldsymbol{w}_k = \boldsymbol{0}$$

Population PCs

The jth population principal component (PC) of X_1, X_2, \ldots, X_m is

$$\mathbf{w}_{i}^{T}\mathbf{X} = w_{1j}X_{1} + w_{2j}X_{2} + \dots + w_{mj}X_{m}$$

where $\boldsymbol{w}_j = (w_{1j}, w_{2j}, \dots, w_{mj})^T$ is column j of \boldsymbol{W} from the eigendecomposition

$$\Sigma = W \Lambda W^T$$
.

The column w_j are called the loadings of the jth principal component. The variance explained by the jth PC is λ_j , which is diagonal element j of Λ .

Sample PCA

Suppose we have m variables, each with n observations:

$$x_1 = (x_{11}, x_{12}, \dots, x_{1n})$$

 $x_2 = (x_{21}, x_{22}, \dots, x_{2n})$
 \vdots
 $x_m = (x_{m1}, x_{m2}, \dots, x_{mn})$

We can organize these variables into an $m \times n$ matrix X where row i is x_i .

PCA can be extended from the population scenario applied to rv's to the sample scenario applied to the observed data X.

Consider all possible weighted sums of these variables

$$\tilde{\boldsymbol{x}} = \sum_{i=1}^m u_i \boldsymbol{x_i}$$

where we constrain $\sum_{i=1}^{m} u_i^2 = 1$.

The first principal component of X is the results \tilde{x} with maximum sample variance

$$s_{\tilde{x}}^2 = \frac{\sum_{j=1}^n \left(\tilde{x}_j - \frac{1}{n} \sum_{k=1}^n \tilde{x}_k \right)^2}{n-1}.$$

This first sample principal component (PC) is then "removed" from the data, and the procedure is repeated until $\min(m, n-1)$ sample PCs are constructed.

The sample PCs are found in a manner analogous to the population PCs. First, we construct the $m \times m$ sample covariance matrix S with (i, j) entry

$$s_{ij} = \frac{\sum_{k=1}^{n} (x_{ik} - \bar{x}_{i.})(x_{jk} - \bar{x}_{j.})}{n-1}.$$

Identifying \boldsymbol{u} that maximizes $s_{\tilde{\boldsymbol{x}}}^2$ also maximizes

$$\boldsymbol{u}^T \boldsymbol{S} \boldsymbol{u}$$
.

Following the steps from before, we want to identify u and λ where

$$Su = \lambda u$$
.

which is accomplished with the eigendecomposition

$$S = U \Lambda U^T$$

where again $U^TU = UU^T = I$ and Λ is a diagonal matrix so that $\lambda_1 \geq \lambda_2 \geq \cdots \geq \lambda_m \geq 0$.

Sample PCs

Let $x_{ij}^* = x_{ij} - \bar{x}_i$ be the row-wise mean-centered values of X, and let X^* be the matrix composed of these values. Also, let u_j be column j of U from $S = U\Lambda U^T$.

Sample PC j is then

$$\tilde{\boldsymbol{x}}_j = \boldsymbol{u}_j^T \boldsymbol{X}^* = \sum_{i=1}^m u_{ij} \boldsymbol{x}_i^*$$

for $j = 1, 2, ..., \min(m, n - 1)$.

The loadings corresponding to PC j are u_i .

Note that the mean of PC j is zero, i.e., that

$$\frac{1}{n}\sum_{k=1}^{n}\tilde{x}_{jk}=0.$$

It can be calculated that the variance of PC j is

$$s_{\tilde{x}_j}^2 = \frac{\sum_{k=1}^n \tilde{x}_{jk}^2}{n-1} = \lambda_j.$$

Proportion of Variance Explained

The proportion of variance explained by PC j is

$$PVE_j = \frac{\lambda_j}{\sum_{k=1}^m \lambda_k}.$$

Singular Value Decomposition

One way in which PCA is performed is to carry out a singular value decomposition (SVD) of the data matrix X. Let $q = \min(m, n)$. Recalling that X^* is the row-wise mean centered X, we can take the SVD of $X^*/\sqrt{n-1}$ to obtain

$$\frac{1}{\sqrt{n-1}}\boldsymbol{X}^* = \boldsymbol{U}\boldsymbol{D}\boldsymbol{V}^T$$

where $U_{m\times q}$, $V_{n\times q}$, and diagonal $D_{q\times q}$. Also, we have the orthogonality properties $V^TV = U^TU = I_q$. Finally, D is composed of diagonal elements $d_1 \geq d_2 \geq \cdots \geq d_q \geq 0$ where $d_q = 0$ if q = n.

Note that

$$S = \frac{1}{n-1} X^* X^{*T} = UDV^T \left(UDV^T \right)^T = UD^2 U^T.$$

Therefore:

- The variance of PC j is $\lambda_j = d_j^2$
- The loadings of PC j are contained in the columns of the left-hand matrix from the decomposition of S or X^*
- PC j is row j of DV^T

My PCA Function

```
> pca <- function(x, space=c("rows", "columns"),</pre>
                    center=TRUE, scale=FALSE) {
    space <- match.arg(space)</pre>
    if(space=="columns") {x <- t(x)}</pre>
    x <- t(scale(t(x), center=center, scale=scale))</pre>
    x \leftarrow x/sqrt(nrow(x)-1)
    s \leftarrow svd(x)
    loading <- s$u
    colnames(loading) <- paste0("Loading", 1:ncol(loading))</pre>
    rownames(loading) <- rownames(x)</pre>
    pc <- diag(s$d) %*% t(s$v)
   rownames(pc) <- paste0("PC", 1:nrow(pc))</pre>
   colnames(pc) <- colnames(x)</pre>
   pve <- s$d^2 / sum(s$d^2)
    if(space=="columns") {pc <- t(pc); loading <- t(loading)}</pre>
    return(list(pc=pc, loading=loading, pve=pve))
```

How It Works

Input is as follows:

- x: a matrix of numerical values
- space: either "rows" or "columns", denoting which dimension contains the variables
- center: if TRUE then the variables are mean centered before calculating PCs
- scale: if TRUE then the variables are std dev scaled before calculating PCs

Output is a list with the following items:

- pc: a matrix of all possible PCs
- loading: the weights or "loadings" that determined each PC
- pve: the proportion of variation explained by each PC

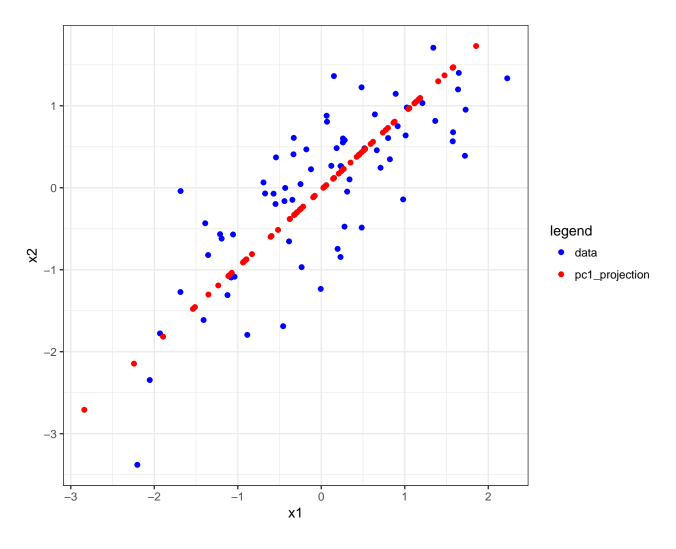
Note that the rows or columns of pc and loading have names to let you know on which dimension the values are organized.

The Ubiquitous Example

Here's an example very frequently encountered to explain PCA, but it's slightly complicated.

```
> set.seed(508)
> n <- 70
> z <- sqrt(0.8) * rnorm(n)
> x1 <- z + sqrt(0.2) * rnorm(n)
> x2 <- z + sqrt(0.2) * rnorm(n)
> X <- rbind(x1, x2)
> p <- pca(x=X, space="rows")</pre>
```

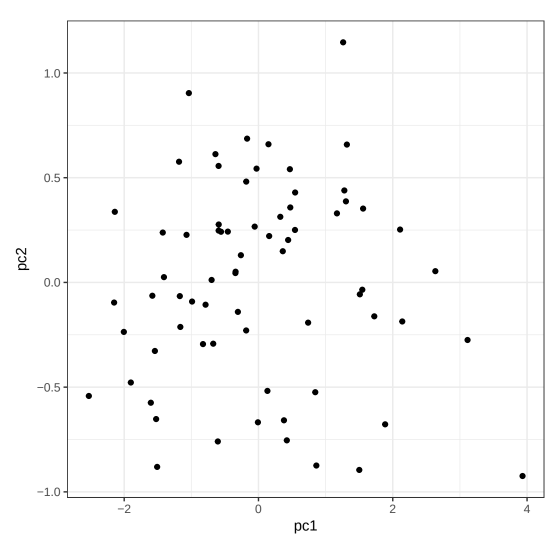
"The first PC finds the direction of maximal variance in the data. . . "



The above figure was made with the following code:

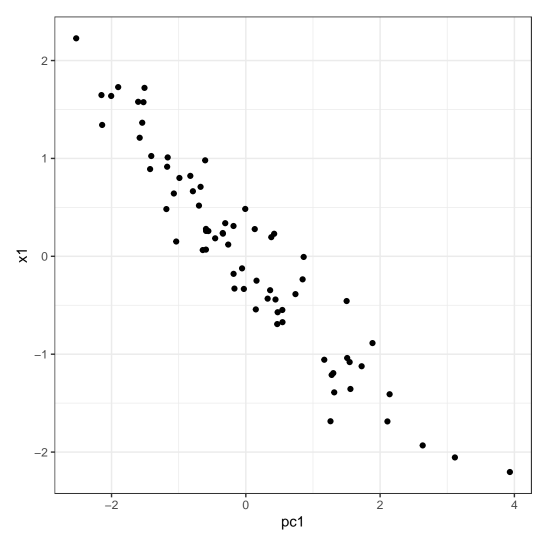
The red dots are therefore the projection of x1 and x2 onto the first PC, so they are neither the loadings nor the PC.

```
Note that
outer(p$loading[,1], p$pc[1,])[1,] + mean(x1)
# yields the same as
lm(x1 ~ p$pc[1,])$fit # and
outer(p$loading[,1], p$pc[1,])[2,] + mean(x2)
# yields the same as
lm(x2 ~ p$pc[1,])$fit
Here is PC1 vs PC2:
> data.frame(pc1=p$pc[1,], pc2=p$pc[2,]) %>%
+ ggplot() + geom_point(aes(x=pc1,y=pc2)) +
+ theme(aspect.ratio=1)
```



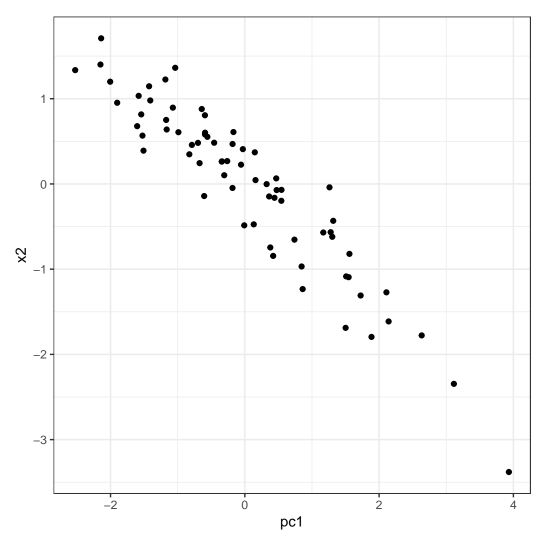
Here is PC1 vs x1:

```
> data.frame(pc1=p$pc[1,], x1=x1) %>%
+ ggplot() + geom_point(aes(x=pc1,y=x1)) +
+ theme(aspect.ratio=1)
```



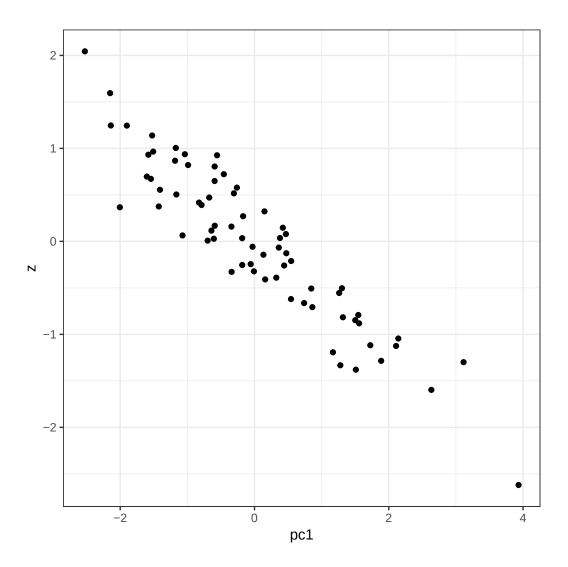
Here is PC1 vs x2:

```
> data.frame(pc1=p$pc[1,], x2=x2) %>%
+ ggplot() + geom_point(aes(x=pc1,y=x2)) +
+ theme(aspect.ratio=1)
```



Here is PC1 vs z:

```
> data.frame(pc1=p$pc[1,], z=z) %>%
+ ggplot() + geom_point(aes(x=pc1,y=z)) +
+ theme(aspect.ratio=1)
```

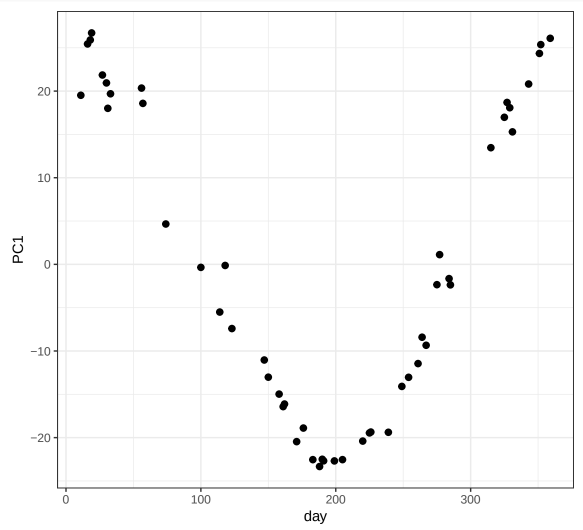


PCA Examples

Weather Data

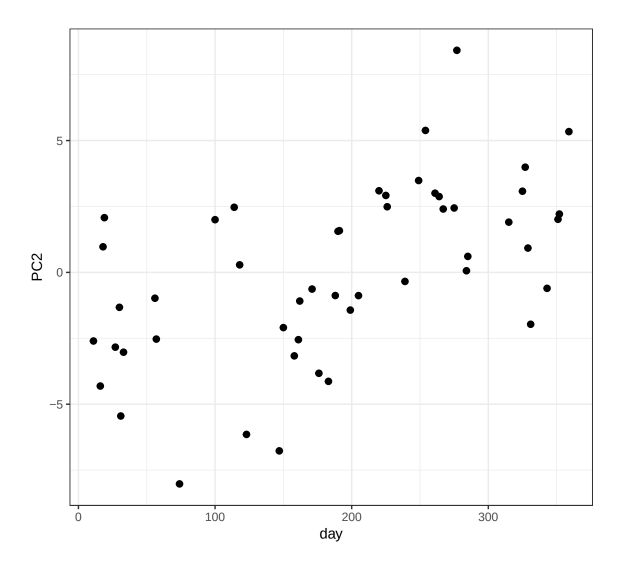
PC1 vs Time

```
> day_of_the_year <- as.numeric(colnames(weather_data))
> data.frame(day=day_of_the_year, PC1=mypca$pc[1,]) %>%
+ ggplot() + geom_point(aes(x=day, y=PC1), size=2)
```



PC2 vs Time

```
> data.frame(day=day_of_the_year, PC2=mypca$pc[2,]) %>%
+ ggplot() + geom_point(aes(x=day, y=PC2), size=2)
```



PC Biplots

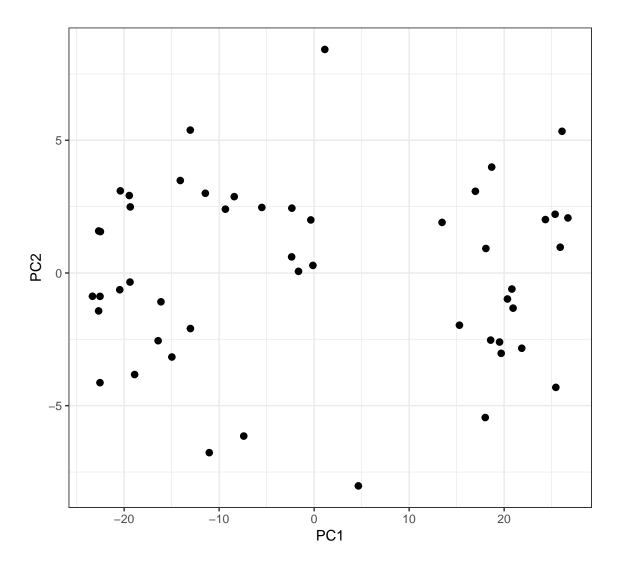
Sometimes it is informative to plot a PC versus another PC. This is called a ${\bf PC}$ biplot.

It is possible that interesting subgroups or clusters of observations will emerge.

This does not appear to be the case in the weather data set, however, due to what we observe in the next two plots.

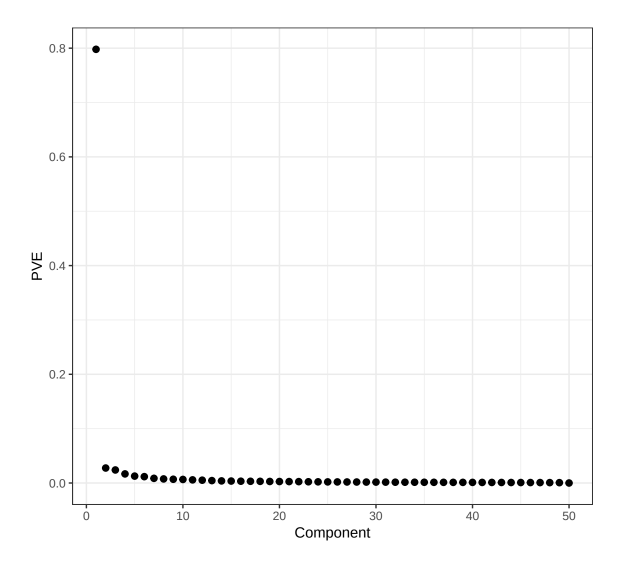
PC1 vs PC2 Biplot

```
> data.frame(PC1=mypca$pc[1,], PC2=mypca$pc[2,]) %>%
+ ggplot() + geom_point(aes(x=PC1, y=PC2), size=2)
```



Proportion of Variance Explained

```
> data.frame(Component=1:length(mypca$pve), PVE=mypca$pve) %>%
+ ggplot() + geom_point(aes(x=Component, y=PVE), size=2)
```



PCs Reproduce the Data

We can multiple the loadings matrix by the PCs matrix to reproduce the data:

Loadings

The sum of squared weights – i.e., loadings – equals one for each component:

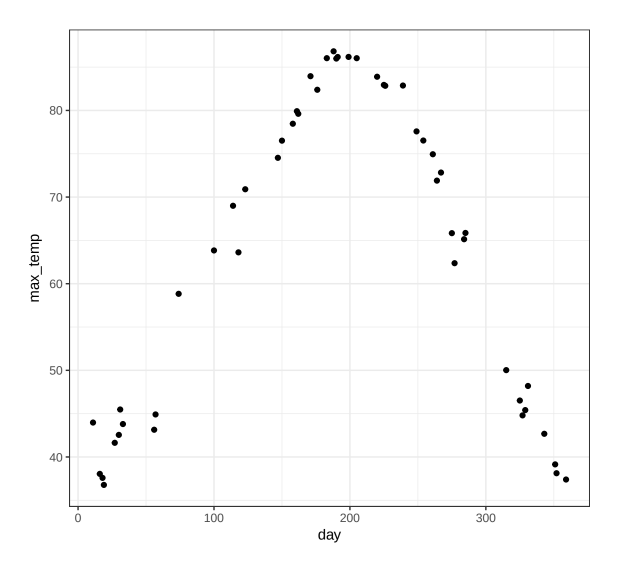
```
> sum(mypca$loading[,1]^2)
[1] 1
>
```

```
> apply(mypca$loading, 2, function(x) {sum(x^2)})
Loading1 Loading2 Loading3 Loading4 Loading5 Loading6
      1
             1
                      1
                              1
                                     1
Loading7 Loading8 Loading9 Loading10 Loading11 Loading12
      1 1 1 1 1
Loading13 Loading14 Loading15 Loading16 Loading17 Loading18
      1 1
                     1
                          1
                                  1
Loading19 Loading20 Loading21 Loading22 Loading23 Loading24
             1
                     1
                             1
Loading25 Loading26 Loading27 Loading28 Loading29 Loading30
      1 1 1 1 1
Loading31 Loading32 Loading33 Loading34 Loading35 Loading36
        1 1 1 1
Loading37 Loading38 Loading39 Loading40 Loading41 Loading42
             1
                     1
                             1
Loading43 Loading44 Loading45 Loading46 Loading47 Loading48
                      1
Loading49 Loading50
```

Pairs of PCs Have Correlaton Zero

PCs by contruction have sample correlation equal to zero:

```
> cor(mypca$pc[1,], mypca$pc[2,])
[1] 3.135149e-17
> cor(mypca$pc[1,], mypca$pc[3,])
[1] 2.273613e-16
> cor(mypca$pc[1,], mypca$pc[12,])
[1] -1.231339e-16
> cor(mypca$pc[5,], mypca$pc[27,])
[1] -2.099516e-17
> # etc...
> day_of_the_year <- as.numeric(colnames(weather_data))
> y <- -mypca$pc[1,] + mean(weather_data)
> data.frame(day=day_of_the_year, max_temp=y) %>%
+ ggplot() + geom_point(aes(x=day, y=max_temp))
```

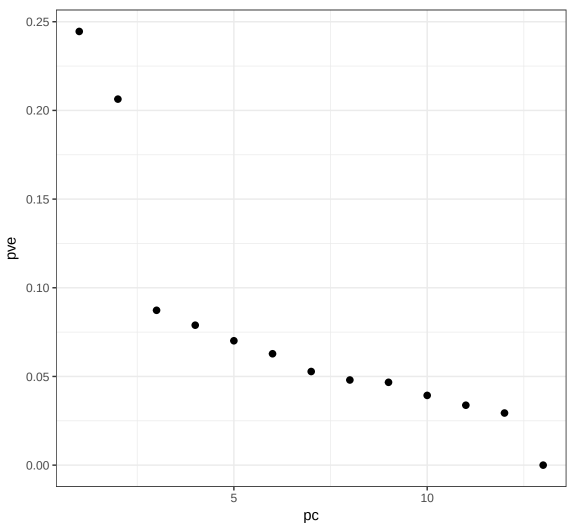


Yeast Gene Expression

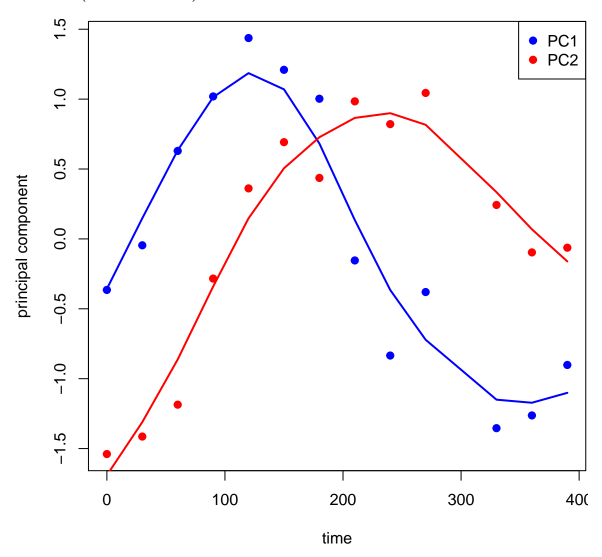
Yeast cells were synchronized so that they were on the same approximate cell cycle timing. The goal was to understand how gene expression varies over the cell cycle from a genome-wide perspective.

```
> load("./data/spellman.RData")
> time
      0 30 60 90 120 150 180 210 240 270 330 360 390
 [1]
> dim(gene_expression)
[1] 5981
> gene_expression[1:6,1:5]
                 0
                           30
                                      60
                                                90
YALOO1C 0.69542786 -0.4143538 3.2350520 1.6323737 -2.1091820
YALOO2W -0.01210662 3.0465649 1.1062193 4.0591467 -0.1166399
YALOO3W -2.78570526 -1.0156981 -2.1387564 1.9299681 0.7797033
YALOO4W 0.55165887 0.6590093 0.5857847 0.3890409 -1.0009777
YALO05C -0.53191556 0.1577985 -1.2401448 0.8170350 -1.3520947
YALOO7C -0.86693416 -1.1642322 -0.6359588 1.1179131 1.9587021
```

Proportion Variance Explained



PCs vs Time (with Smoothers)

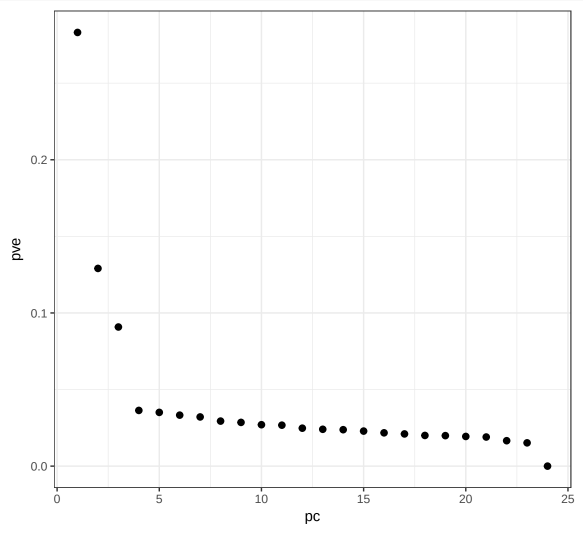


HapMap Genotypes

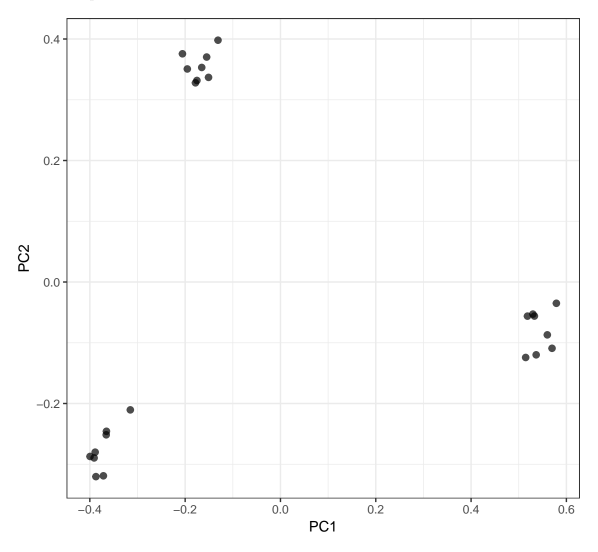
Recall the example HapMap data used to demonstrate the MCMC algorithm for estimating structure. We curated a small data set that cleanly separates human subpopulations.

```
> hapmap <- read.table("./data/hapmap_sample.txt")</pre>
> dim(hapmap)
[1] 400 24
> hapmap[1:6,1:6]
            NA18516 NA19138 NA19137 NA19223 NA19200 NA19131
rs2051075
                  0
                                    2
                                                               1
                           1
                                             1
                                                      1
                  2
                           2
                                    0
rs765546
                                             0
                                                      0
                                                               0
                  2
                           2
                                    2
                                                               2
rs10019399
                                             1
                                                      1
                  2
                           2
                                             2
                                                               2
rs7055827
                                    1
                           0
                                    2
                                                               0
rs6943479
                  0
                                             0
                                                      1
rs2095381
```

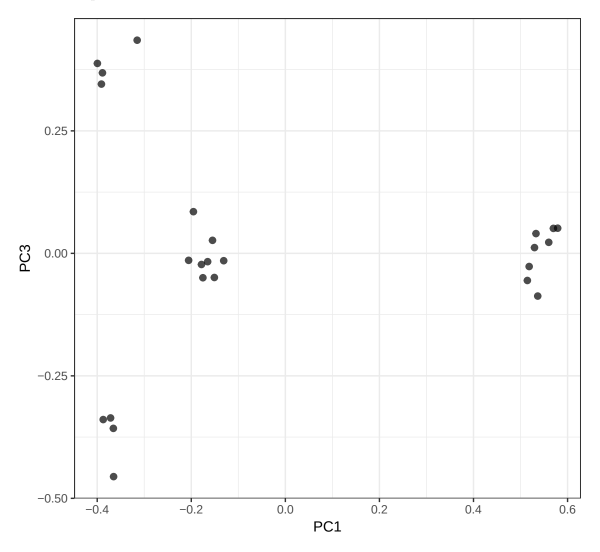
Proportion Variance Explained



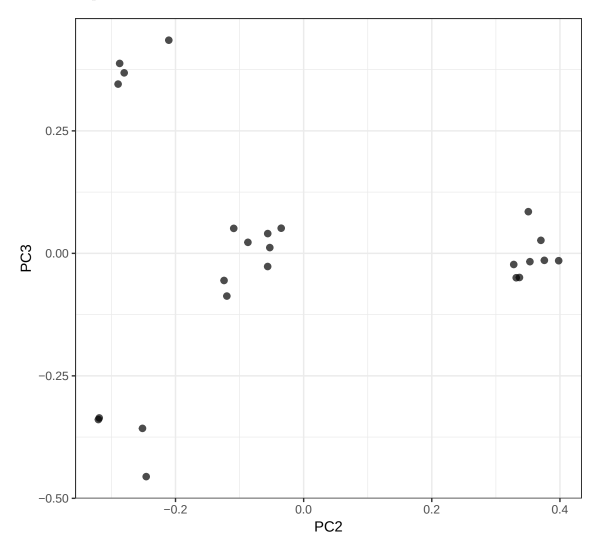
PC1 vs PC2 Biplot



PC1 vs PC3 Biplot



PC2 vs PC3 Biplot



HD Latent Variable Models

Definition

Latent variables (or hidden variables) are random variables that are present in the underlying probabilistic model of the data, but they are unobserved.

In high-dimensional data, there may be latent variables present that affect many variables simultaneously.

These are latent variables that induce **systematic variation**. A topic of much interest is how to estimate these and incorporate them into further HD inference procedures.

Model

Suppose we have observed data $Y_{m \times n}$ of m variables with n observations each. Suppose there are r latent variables contained in the r rows of $Z_{r \times n}$ where

$$E[\boldsymbol{Y}_{m\times n} \mid \boldsymbol{Z}_{r\times n}] = \boldsymbol{\Phi}_{m\times r} \boldsymbol{Z}_{r\times n}.$$

Let's also assume that $m \gg n > r$. The latent variables Z induce systematic variation in variable y_i parameterized by ϕ_i for i = 1, 2, ..., m.

Estimation

There exist methods for estimating the row space of Z with probability 1 as $m \to \infty$ for a fixed n in two scenarios.

Leek (2011) shows how to do this when $y_i|Z \sim \text{MVN}(\phi_i Z, \sigma_i^2 I)$, and the $y_i|Z$ are jointly independent.

Chen and Storey (2015) show how to do this when the $y_i|Z$ are distributed according to a single parameter exponential family distribution with mean $\phi_i Z$, and the $y_i|Z$ are jointly independent.

Jackstraw

Suppose we have a reasonable method for estimating Z in the model

$$\mathrm{E}\left[Y\mid Z\right] = \Phi Z.$$

The jackstraw method allows us to perform hypothesis tests of the form

$$H_0: \phi_i = \mathbf{0} \text{ vs } H_1: \phi_i \neq \mathbf{0}.$$

We can also perform this hypothesis test on any subset of the columns of Φ .

This is a challening problem because we have to "double dip" in the data Y, first to estimate Z, and second to perform significance tests on Φ .

Procedure

The first step is to form estimate \hat{Z} and then test statistic t_i that performs the hypothesis test for each ϕ_i from y_i and \hat{Z} (i = 1, ..., m). Assume that the larger t_i is, the more evidence there is against the null hypothesis in favor of the alternative.

Next we randomly select s rows of Y and permute them to create data set Y^0 . Let this set of s variables be indexed by S. This breaks the relationship between y_i and Z, thereby inducing a true H_0 , for each $i \in S$.

We estimate $\hat{\mathbf{Z}}^0$ from \mathbf{Y}^0 and again obtain test statistics t_i^0 . Specifically, the test statistics t_i^0 for $i \in \mathcal{S}$ are saved as draws from the null distribution.

We repeat permutation procedure B times, and then utilize all saved sB permutation null statistics to calculate empirical p-values:

$$p_i = \frac{1}{sB} \sum_{b=1}^{B} \sum_{k \in S_b} 1 \left(t_i \ge t_k^{0b} \right).$$

Example: Yeast Cell Cycle

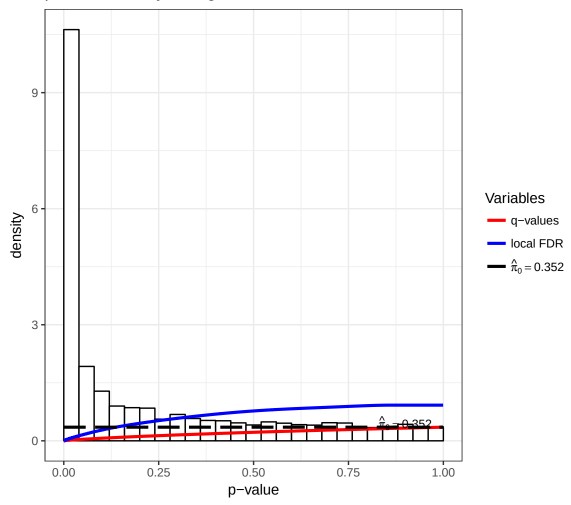
Recall the yeast cell cycle data from earlier. We will test which genes have expression significantly associated with PC1 and PC2 since these both capture cell cycle regulation.

```
> library(jackstraw)
> load("./data/spellman.RData")
> time
[1]  0  30  60  90  120  150  180  210  240  270  330  360  390
> dim(gene_expression)
[1]  5981   13
> dat <- t(scale(t(gene_expression), center=TRUE, scale=FALSE))</pre>
```

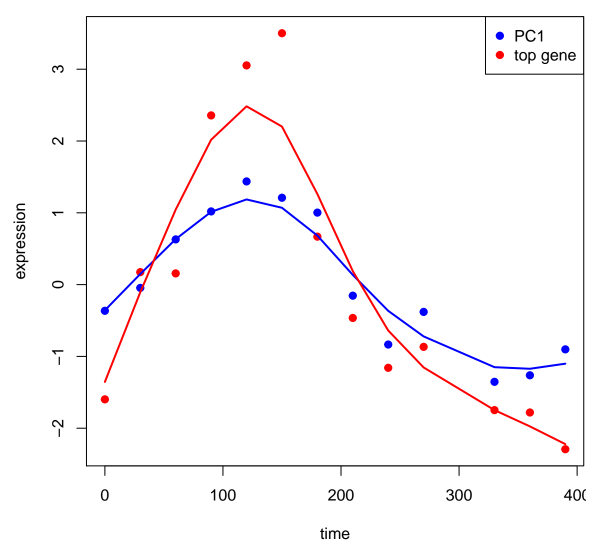
Test for associations between PC1 and each gene, conditioning on PC1 and PC2 being relevant sources of systematic variation.

```
> jsobj <- jackstraw(dat, r1=1, r=2, B=500, s=50, verbose=FALSE)
> jsobj$p.value %>% qvalue() %>% hist()
```

p-value density histogram

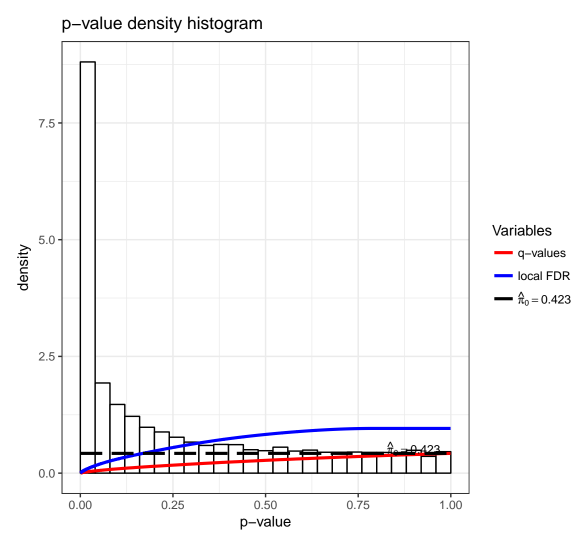


This is the most significant gene plotted with PC1.

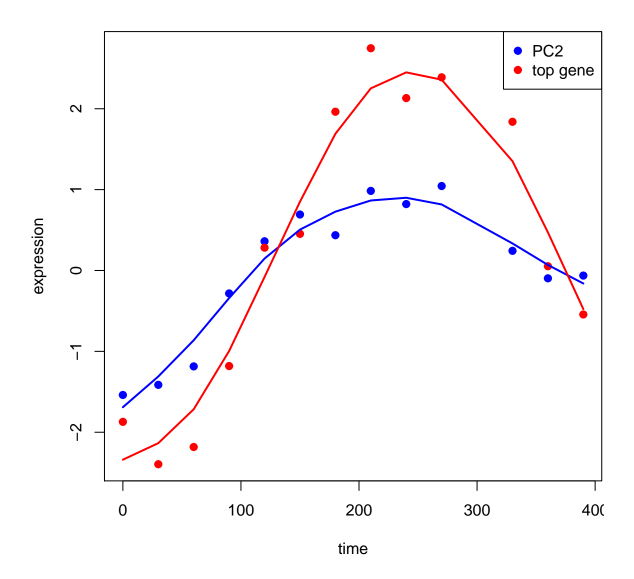


Test for associations between PC2 and each gene, conditioning on PC1 and PC2 being relevant sources of systematic variation.

```
> jsobj <- jackstraw(dat, r1=2, r=2, B=500, s=50, verbose=FALSE)
> jsobj$p.value %>% qvalue() %>% hist()
```



This is the most significant gene plotted with PC2.



Surrogate Variable Analysis

The surrogate variable analysis (SVA) model combines the many responses model with the latent variable model introduced above:

$$oldsymbol{Y}_{m imes n} = oldsymbol{B}_{m imes d} oldsymbol{X}_{d imes n} + oldsymbol{\Phi}_{m imes r} oldsymbol{Z}_{r imes n} + oldsymbol{E}_{m imes n}$$

where $m \gg n > d + r$.

Here, only Y and X are observed, so we must combine many regressors model fitting techniques with latent variable estimation.

The variables Z are called **surrogate variables** for what would be a complete model of all systematic variation.

Procedure

The main challenge is that the row spaces of X and Z may overlap. Even when X is the result of a randomized experiment, there will be a high probability that the row spaces of X and Z have some overlap.

Therefore, one cannot simply estimate Z by applying a latent variable estimation method on the residuals $Y - \hat{B}X$ or on the observed response data Y. In the former case, we will only estimate Z in the space orthogonal to $\hat{B}X$. In the latter case, the estimate of Z may modify the signal we can estimate in BX.

A recent method, takes an EM approach to esit mating \boldsymbol{Z} in the model

$$Y_{m \times n} = B_{m \times d} X_{d \times n} + \Phi_{m \times r} Z_{r \times n} + E_{m \times n}.$$

It is shown to be necessary to penalize the likelihood in the estimation of B — i.e., form shrinkage estimates of B — in order to properly balance the row spaces of X and Z.

The regularized EM algorithm, called cross-dimensional inference (CDI) iterates between

- 1. Estimate \boldsymbol{Z} from $\boldsymbol{Y} \hat{\boldsymbol{B}}^{\mathrm{Reg}} \boldsymbol{X}$
- 2. Estimate \boldsymbol{B} from $\boldsymbol{Y} \hat{\boldsymbol{\Phi}}\hat{\boldsymbol{Z}}$

where $\hat{\boldsymbol{B}}^{\mathrm{Reg}}$ is a regularized or shrunken estimate of $\boldsymbol{B}.$

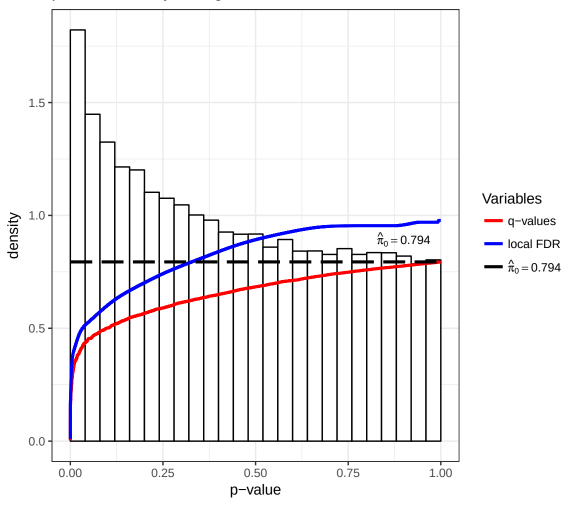
It can be shown that when the regularization can be represented by a prior distribution on B then this algorithm achieves the MAP.

Example: Kidney Expr by Age

In Storey et al. (2005), we considered a study where kidney samples were obtained on individuals across a range of ages. The goal was to identify genes with expression associated with age.

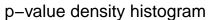
```
> library(edge)
> library(splines)
> load("./data/kidney.RData")
> age <- kidcov$age
> sex <- kidcov$sex
> dim(kidexpr)
[1] 34061
             72
> cov <- data.frame(sex = sex, age = age)
> null model <- ~sex
> full_model <- ~sex + ns(age, df = 3)</pre>
> de_obj <- build_models(data = kidexpr, cov = cov,</pre>
                          null.model = null model,
                          full.model = full model)
> de_lrt <- lrt(de_obj, nullDistn = "bootstrap", bs.its = 100, verbose=FALSE)
> qobj1 <- qvalueObj(de_lrt)</pre>
> hist(qobj1)
```

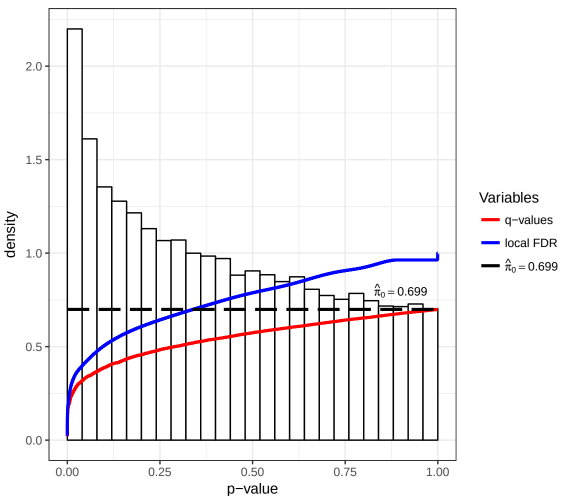




Now that we have completed a standard generalized LRT, let's estimate \boldsymbol{Z} (the surrogate variables) using the sva package as accessed via the edge package.

```
> dim(nullMatrix(de_obj))
[1] 72 2
> de_sva <- apply_sva(de_obj, n.sv=4, method="irw", B=10)
Number of significant surrogate variables is: 4
Iteration (out of 10 ):1 2 3 4 5 6 7 8 9 10
> dim(nullMatrix(de_sva))
[1] 72 6
> de_svalrt <- lrt(de_sva, nullDistn = "bootstrap", bs.its = 100, verbose=FALSE)
> qobj2 <- qvalueObj(de_svalrt)
> hist(qobj2)
```



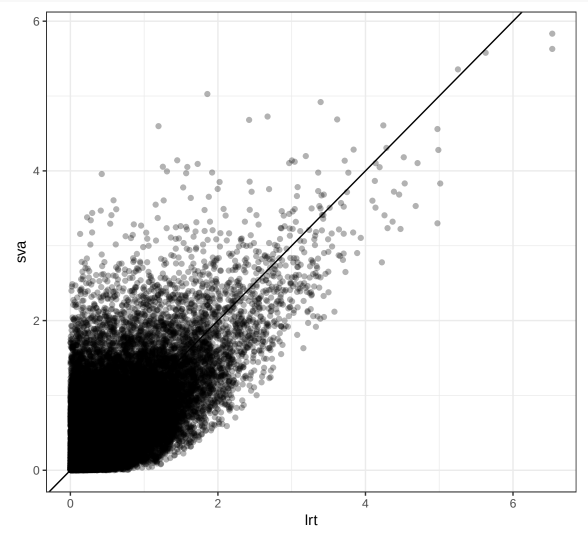


```
> summary(qobj1)
Call:
qvalue(p = pval)
       0.7939407
pi0:
Cumulative number of significant calls:
          <1e-04 <0.001 <0.01 <0.025 <0.05 <0.1
                        798
                              1709 2966 5392 34061
              26
                   145
p-value
                            2
q-value
                      0
                                   3
                                         8
                                             26 34061
local FDR
                                             12 34061
> summary(qobj2)
Call:
qvalue(p = pval)
      0.698907
pi0:
```

```
Cumulative number of significant calls:
         <1e-04 <0.001 <0.01 <0.025 <0.05 <0.1
p-value
                   158 1014 2067 3555 6141 34061
             28
q-value
              0
                                        6
                                           47 34061
                                  3
local FDR
              0
                     0
                          0
                                 1
                                      4
                                           29 34054
```

P-values from two analyses are fairly different.

```
> data.frame(lrt=-log10(qobj1$pval), sva=-log10(qobj2$pval)) %>%
+ ggplot() + geom_point(aes(x=lrt, y=sva), alpha=0.3) + geom_abline()
```



Extras

Source

License

Source Code

Session Information

```
> sessionInfo()
R version 3.3.2 (2016-10-31)
Platform: x86_64-apple-darwin13.4.0 (64-bit)
Running under: macOS Sierra 10.12.4
locale:
[1] en_US.UTF-8/en_US.UTF-8/en_US.UTF-8/en_US.UTF-8/en_US.UTF-8
attached base packages:
[1] splines
                                  graphics grDevices utils
              parallel stats
[7] datasets methods
other attached packages:
 [1] edge_2.6.0
                         Biobase_2.34.0
 [3] BiocGenerics_0.20.0 jackstraw_1.1
 [5] qvalue_2.1.1
                         MASS_7.3-47
 [7] broom 0.4.2
                         dplyr 0.5.0
                         readr_1.1.0
 [9] purrr_0.2.2
[11] tidyr_0.6.2
                         tibble 1.3.0
[13] ggplot2_2.2.1
                         tidyverse_1.1.1
[15] knitr_1.15.1
                         magrittr_1.5
[17] devtools 1.12.0
loaded via a namespace (and not attached):
 [1] httr_1.2.1
                          jsonlite_1.4
 [3] modelr_0.1.0
                          assertthat_0.2.0
 [5] highr_0.6
                          stats4_3.3.2
 [7] cellranger_1.1.0
                          yaml_2.1.14
[9] RSQLite_1.1-2
                          backports_1.0.5
[11] lattice_0.20-35
                          digest_0.6.12
[13] rvest_0.3.2
                          minqa_1.2.4
[15] colorspace_1.3-2
                          htmltools_0.3.6
[17] Matrix_1.2-10
                          plyr_1.8.4
[19] psych 1.7.5
                          XML 3.98-1.7
[21] haven_1.0.0
                          genefilter_1.56.0
[23] xtable_1.8-2
                          corpcor_1.6.9
[25] scales_0.4.1
                          lme4_1.1-13
[27] annotate_1.52.1
                          mgcv_1.8-17
[29] IRanges_2.8.2
                          withr_1.0.2
[31] lazyeval_0.2.0
                          mnormt_1.5-5
[33] survival_2.41-3
                          snm_1.22.0
[35] readxl_1.0.0
                          memoise_1.1.0
[37] evaluate_0.10
                          nlme_3.1-131
[39] forcats_0.2.0
                          xm12_1.1.1
[41] foreign_0.8-68
                          tools_3.3.2
[43] hms_0.3
                          stringr_1.2.0
[45] S4Vectors_0.12.2
                          lfa 1.4.0
[47] munsell_0.4.3
                          AnnotationDbi_1.36.2
[49] grid_3.3.2
                          RCurl_1.95-4.8
[51] nloptr_1.0.4
                          bitops_1.0-6
[53] labeling_0.3
                          rmarkdown_1.5
```

```
[55] codetools_0.2-15 gtable_0.2.0
[57] DBI_0.6-1 reshape2_1.4.2
[59] R6_2.2.0 lubridate_1.6.0
[61] rprojroot_1.2 stringi_1.1.5
[63] sva_3.22.0 Rcpp_0.12.10
```