

Principal Component Analysis

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STAT 4690—Applied Multivariate Analysis

Population PCA i

- **PCA:** Principal Component Analysis
- Dimension reduction method:
 - Let $\mathbf{Y} = (Y_1, \dots, Y_p)$ be a random vector with covariance matrix Σ . We are looking for a transformation $h : \mathbb{R}^p \rightarrow \mathbb{R}^k$, with $k \ll p$ such that $h(\mathbf{Y})$ retains “as much information as possible” about \mathbf{Y} .
- In PCA, we are looking for a **linear transformation** $h(y) = w^T y$ with **maximal variance** (where $\|w\| = 1$)

Population PCA ii

- More generally, we are looking for k linear transformations w_1, \dots, w_k such that $w_j^T \mathbf{Y}$ has maximal variance and is uncorrelated with $w_1^T \mathbf{Y}, \dots, w_{j-1}^T \mathbf{Y}$.
- First, note that $\text{Var}(w^T \mathbf{Y}) = w^T \Sigma w$. So our optimisation problem is

$$\max_w w^T \Sigma w, \quad \text{with } w^T w = 1.$$

- From the theory of Lagrange multipliers, we can look at the *unconstrained* problem

$$\max_{w, \lambda} w^T \Sigma w - \lambda(w^T w - 1).$$

Population PCA iii

- Write $\phi(w, \lambda)$ for the function we are trying to optimise.
We have

$$\begin{aligned}\frac{\partial}{\partial w} \phi(w, \lambda) &= \frac{\partial}{\partial w} w^T \Sigma w - \lambda(w^T w - 1) \\ &= 2\Sigma w - 2\lambda w; \\ \frac{\partial}{\partial \lambda} \phi(w, \lambda) &= w^T w - 1.\end{aligned}$$

- From the first partial derivative, we conclude that

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

Population PCA iv

- From the second partial derivative, we conclude that $w \neq 0$; in other words, w is an eigenvector of Σ with eigenvalue λ .
- Moreover, at this stationary point of $\phi(w, \lambda)$, we have

$$\text{Var}(w^T \mathbf{Y}) = w^T \Sigma w = w^T (\lambda w) = \lambda w^T w = \lambda.$$

- In other words, to maximise the variance $\text{Var}(w^T \mathbf{Y})$, we need to choose λ to be the *largest* eigenvalue of Σ .
- By induction, and using the extra constraints $w_i^T w_j = 0$, we can show that all other linear transformations are given by eigenvectors of Σ .

Population PCA v

PCA Theorem

Let $\lambda_1 \geq \dots \geq \lambda_p$ be the eigenvalues of Σ , with corresponding unit-norm eigenvectors w_1, \dots, w_p . To reduce the dimension of \mathbf{Y} from p to k such that every component of $W^T \mathbf{Y}$ is uncorrelated and each direction has maximal variance, we can take $W = (w_1 \ \dots \ w_k)$, whose j -th column is w_j .

Properties of PCA i

- Some vocabulary:
 - $Z_i = w_i^T \mathbf{Y}$ is called the *i-th principal component* of \mathbf{Y} .
 - w_i is the *i-th vector of loadings*.
- Note that we can take $k = p$, in which case we do not reduce the dimension of \mathbf{Y} , but we *transform* it into a random vector with uncorrelated components.
- Let $\Sigma = P\Lambda P^T$ be the eigendecomposition of Σ . We have

$$\sum_{i=1}^p \text{Var}(w_i^T \mathbf{Y}) = \sum_{i=1}^p \lambda_i = \text{tr}(\Lambda) = \text{tr}(\Sigma) = \sum_{i=1}^p \text{Var}(Y_i).$$

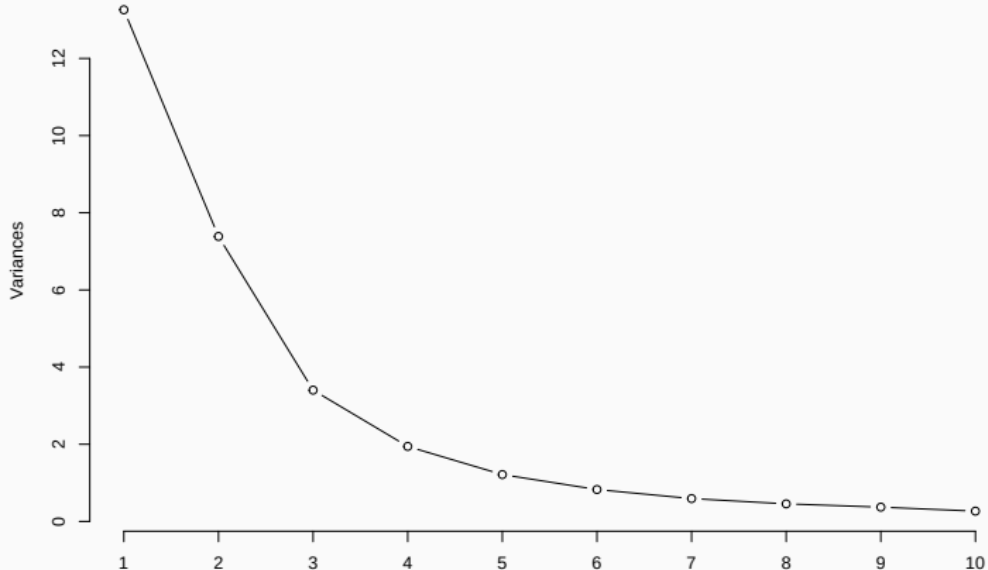
Properties of PCA ii

- Therefore, each linear transformation $w_i^T \mathbf{Y}$ contributes $\lambda_i / \sum_j \lambda_j$ as percentage of the overall variance.
- **Selecting k :** One common strategy is to select a threshold (e.g. $c = 0.9$) such that

$$\frac{\sum_{i=1}^k \lambda_i}{\sum_{i=1}^p \lambda_i} \geq c.$$

Scree plot

- A **scree plot** is a plot with the sequence $1, \dots, p$ on the x-axis, and the sequence $\lambda_1, \dots, \lambda_p$ on the y-axis.
- Another common strategy for selecting k is to choose the point where the curve starts to flatten out.
 - **Note:** This inflection point does not necessarily exist, and it may be hard to identify.



Correlation matrix

- When the observations are on the different scale, it is typically more appropriate to normalise the components of \mathbf{Y} before doing PCA.
 - The variance depends on the units, and therefore without normalising, the component with the “smallest” units (e.g. centimeters vs. meters) could be driving most of the overall variance.
- In other words, instead of using Σ , we can use the (population) correlation matrix R .
- **Note:** The loadings and components we obtain from Σ are **not** equivalent to the ones obtained from R .

Sample PCA

- In general, we do not know the population covariance matrix Σ .
- Therefore, in practice, we estimate the loadings w_i through the eigenvectors of the sample covariance matrix S_n .
- As with the population version of PCA, if the units are different, we should normalise the components or use the sample correlation matrix.

Example 1 i

```
library(mvtnorm)
Sigma <- matrix(c(1, 0.5, 0.1,
                  0.5, 1, 0.5,
                  0.1, 0.5, 1),
                  ncol = 3)

set.seed(17)
X <- rmvnorm(100, sigma = Sigma)
pca <- prcomp(X)
```

Example 1 ii

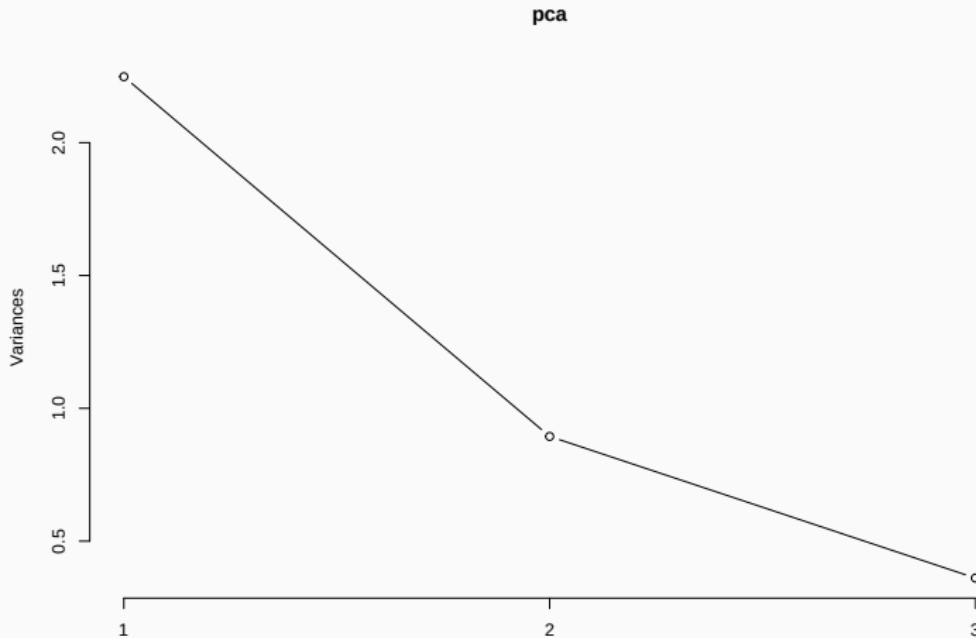
```
summary(pca)
```

```
## Importance of components:
```

	PC1	PC2	PC3
## Standard deviation	1.4994	0.9457	0.6009
## Proportion of Variance	0.6417	0.2552	0.1031
## Cumulative Proportion	0.6417	0.8969	1.0000

```
screeplot(pca, type = 'l')
```

Example 1 iii



Example 2 i

```
pca <- prcomp(USArrests, scale = TRUE)
```

```
summary(pca)
```

```
## Importance of components:
```

```
##
```

```
PC1
```

```
PC2
```

```
PC3
```

```
PC4
```

```
## Standard deviation      1.5749 0.9949 0.59713 0.41645
```

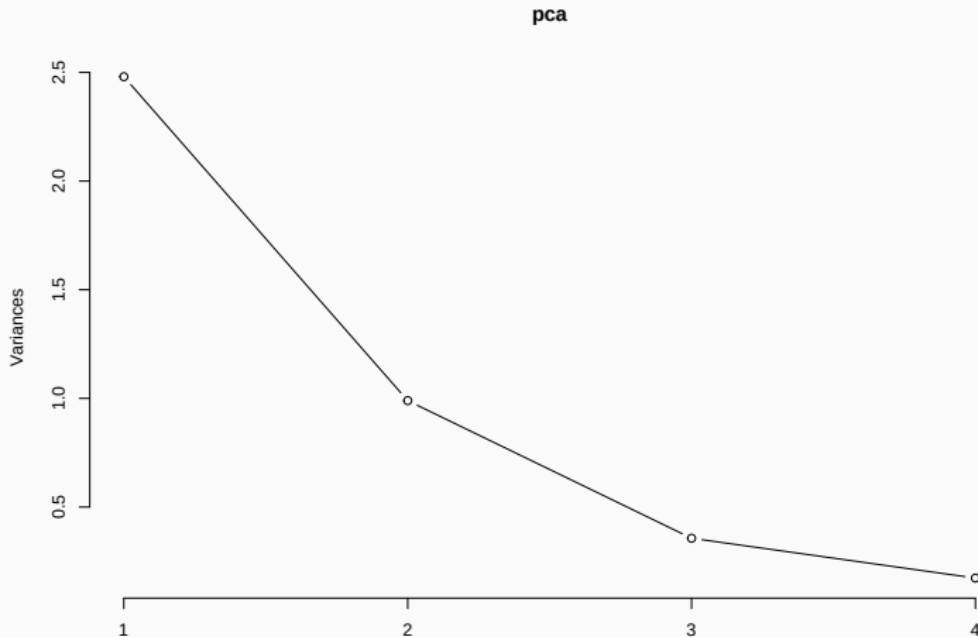
```
## Proportion of Variance 0.6201 0.2474 0.08914 0.04336
```

```
## Cumulative Proportion  0.6201 0.8675 0.95664 1.00000
```

Example 2 ii

```
screeplot(pca, type = 'l')
```

Example 2 iii



Applications of PCA

Training and testing i

- Recall: **Mean Squared Error**

$$MSE = \frac{1}{n} \sum_{i=1}^n (Y_i - \hat{Y}_i)^2,$$

where Y_i, \hat{Y}_i are the *observed* and *predicted* values.

- It is good practice to separate your dataset in two:
 - **Training** dataset, that is used to build and fit your model (e.g. choose covariates, estimate regression coefficients).
 - **Testing** dataset, that it used to compute the MSE or other performance metrics.

Training and testing ii

- PCA can be used for predictive model building in (univariate) linear regression:
 - **Feature extraction:** Perform PCA on the covariates, extract the first k PCs, and use them as predictors in your model.
 - **Feature selection:** Perform PCA on the covariates, look at the first PC, find the covariates whose loadings are the largest (in absolute value), and only use those covariates as predictors.

Feature Extraction i

```
library(ElemStatLearn)
library(tidyverse)
train <- subset(prostate, train == TRUE,
                 select = -train)
test  <- subset(prostate, train == FALSE,
                 select = -train)

# First model: Linear regression
lr_model <- lm(lpsa ~ ., data = train)
lr_pred <- predict(lr_model, newdata = test)
(lr_mse <- mean((test$lpsa - lr_pred)^2))
```

Feature Extraction ii

```
## [1] 0.521274
```

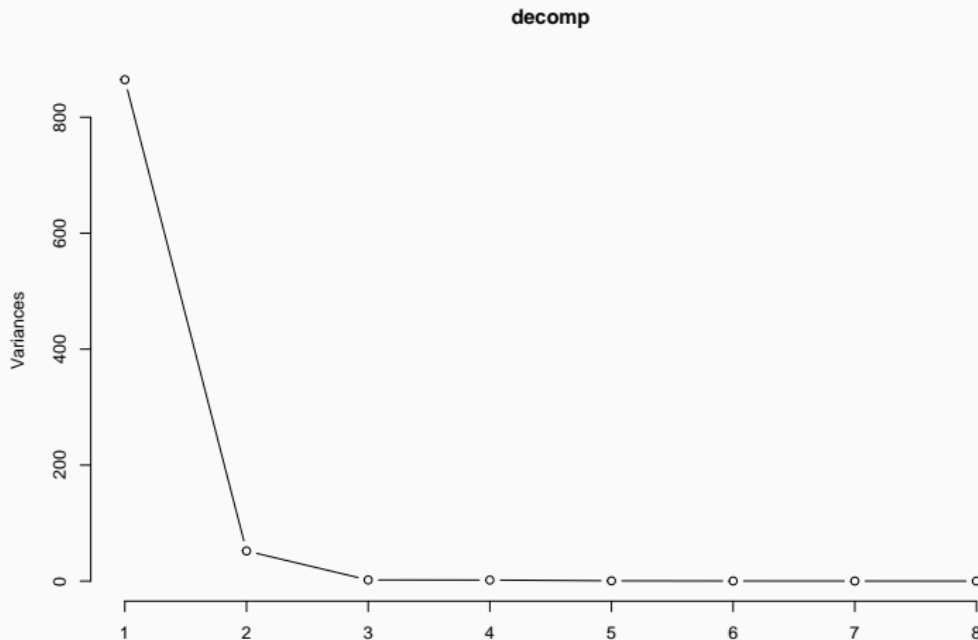
```
# PCA
decomp <- train %>%
  subset(select = -lpsa) %>%
  as.matrix() %>%
  prcomp
summary(decomp)$importance[, 1:3]
```

Feature Extraction iii

```
##                                     PC1        PC2        PC3
## Standard deviation      29.40597 7.211721 1.410789
## Proportion of Variance 0.93844 0.056440 0.002160
## Cumulative Proportion  0.93844 0.994890 0.997050
```

```
screeplot(decomp, type = 'lines')
```

Feature Extraction iv



Feature Extraction v

```
# Second model: PCs for predictors
train_pc <- train
train_pc$PC1 <- decomp$x[,1]
pc_model <- lm(lpsa ~ PC1, data = train_pc)

test_pc <- as.data.frame(predict(decomp, test))
pc_pred <- predict(pc_model,
                     newdata = test_pc)
(pc_mse <- mean((test$lpsa - pc_pred)^2))

## [1] 0.9552741
```

Feature Selection i

```
contribution <- decomp$rotation[, "PC1"]
round(contribution, 3)[1:6]
```

```
##   lcavol lweight      age     lbph      svi      lcp
##   0.021   0.001   0.075 -0.001   0.007   0.032
```

```
round(contribution, 3)[7:8]
```

```
## gleason    pgg45
##   0.018   0.996
```

Feature Selection ii

```
(keep <- names(which(abs(contribution) > 0.01)))  
  
## [1] "lcavol"   "age"       "lcp"       "gleason"  "pgg45"  
  
fs_model <- lm(lpsa ~ ., data = train[,c(keep, "lpsa")])  
fs_pred <- predict(fs_model, newdata = test)  
(fs_mse <- mean((test$lpsa - fs_pred)^2))  
  
## [1] 0.5815571
```

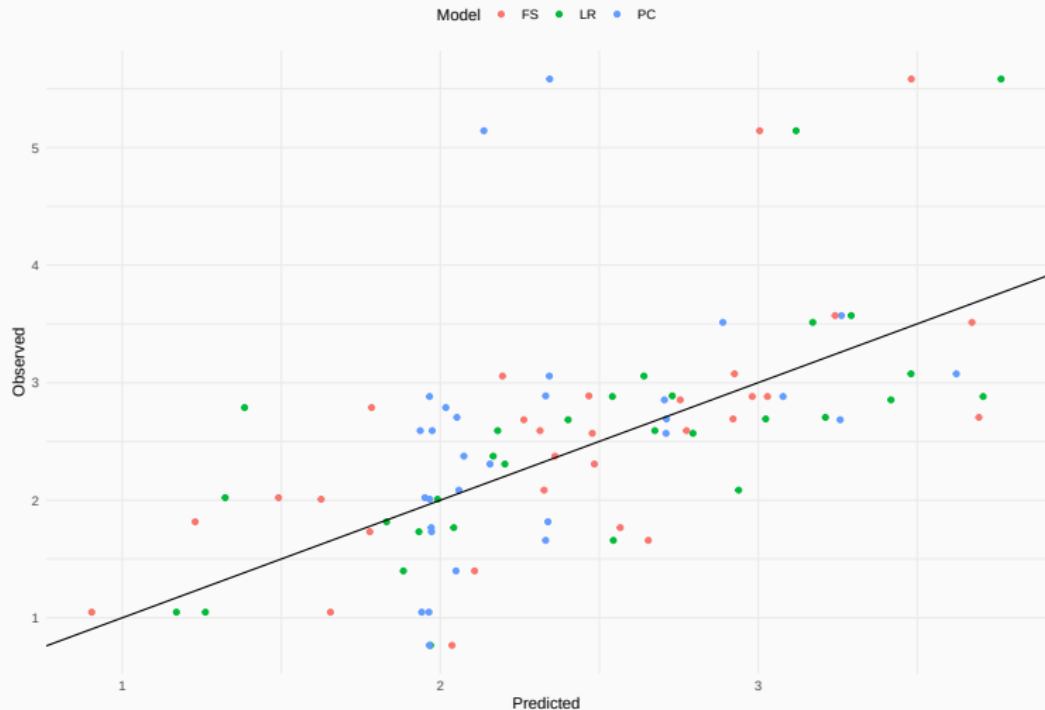
Feature Selection iii

```
model_plot <- data.frame(  
  "obs" = test$lpsa,  
  "LR" = lr_pred,  
  "PC" = pc_pred,  
  "FS" = fs_pred  
) %>%  
  gather(Model, pred, -obs)
```

Feature Selection iv

```
ggplot(model_plot,  
       aes(pred, obs, colour = Model)) +  
  geom_point() +  
  theme_minimal() +  
  geom_abline(slope = 1, intercept = 0) +  
  theme(legend.position = 'top') +  
  xlab("Predicted") + ylab("Observed")
```

Feature Selection v



Comments

- The full model performed better than the ones we created with PCA
 - It had a lower MSE
- On the other hand, if we had multicollinearity issues, or too many covariates ($p > n$), the PCA models could outperform the full model.
- However, note that PCA does not use the association between the covariates and the outcome, so it will never be the most efficient way of building a model.

Data Visualization i

```
library(dslabs)

mnist <- read_mnist()

dim(mnist$train$images)

## [1] 60000    784

dim(mnist$test$images)

## [1] 10000    784
```

Data Visualization ii

```
head(mnist$train$labels)  
  
## [1] 5 0 4 1 9 2  
  
matrix(mnist$train$images[1,], ncol = 28) %>%  
  image(col = gray.colors(12, rev = TRUE),  
        axes = FALSE)
```

Data Visualization iii

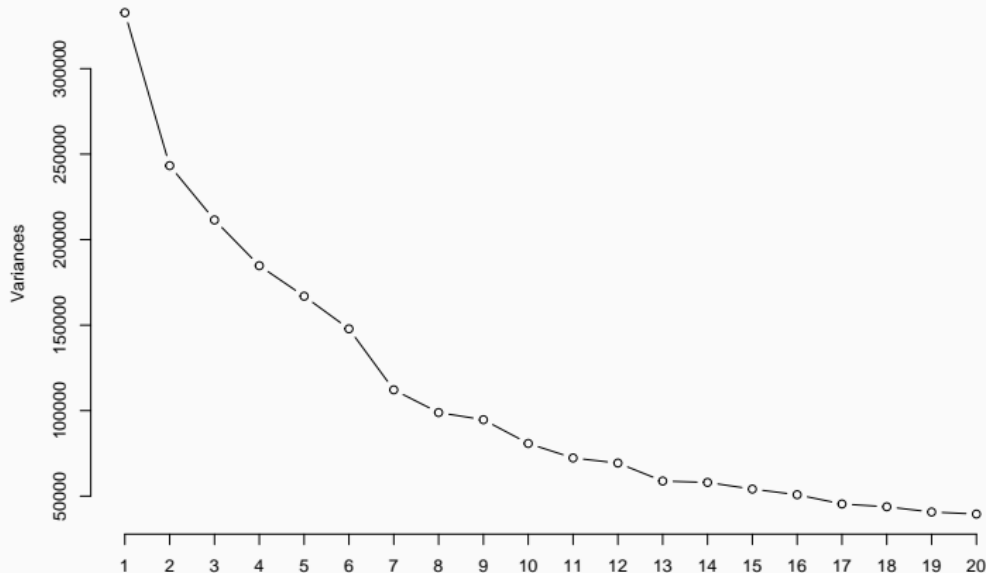


Data Visualization iv

```
decomp <- prcomp(mnist$train$images)
```

```
screeplot(decomp, type = 'lines',
          n pcs = 20, main = "")
```

Data Visualization v



Data Visualization vi

```
decomp$x[,1:2] %>%
  as.data.frame() %>%
  mutate(label = factor(mnist$train$labels)) %>%
  ggplot(aes(PC1, PC2, colour = label)) +
  geom_point(alpha = 0.5) +
  theme_minimal()
```

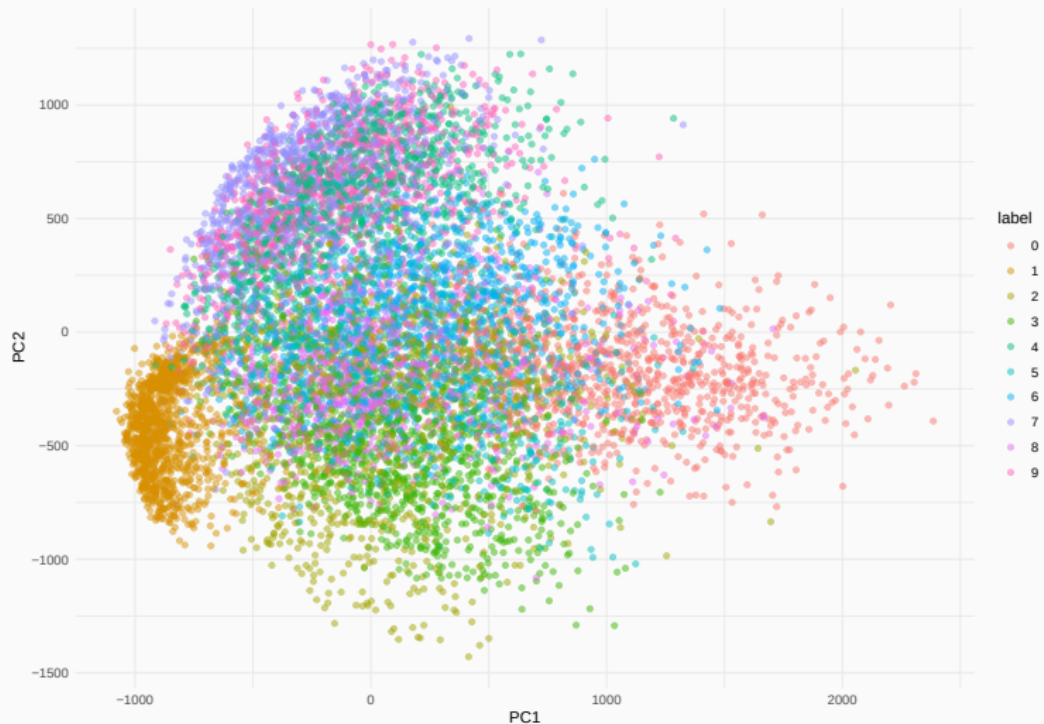
Data Visualization vii



Data Visualization viii

```
# And on the test set
decomp %>%
  predict(newdata = mnist$test$images) %>%
  as.data.frame() %>%
  mutate(label = factor(mnist$test$labels)) %>%
  ggplot(aes(PC1, PC2, colour = label)) +
  geom_point(alpha = 0.5) +
  theme_minimal()
```

Data Visualization ix

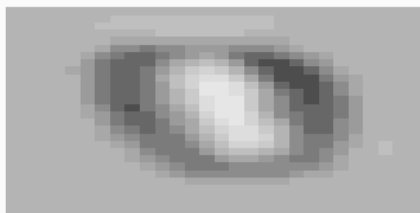


Data Visualization x

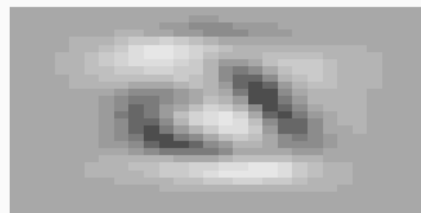
```
par(mfrow = c(2, 2))
for (i in seq_len(4)) {
  matrix(decomp$rotation[,i], ncol = 28) %>%
    image(col = gray.colors(12, rev = TRUE),
          axes = FALSE, main = paste0("PC", i))
}
```

Data Visualization xi

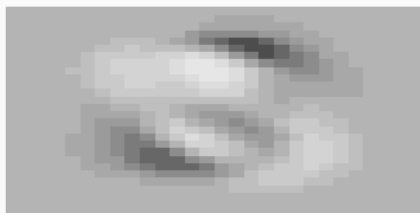
PC1



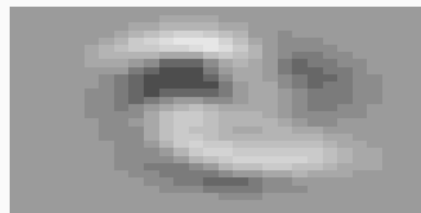
PC2



PC3



PC4



Data Visualization xii

```
# Approximation with 90 PCs
approx_mnist <- decomp$rotation[, seq_len(90)] %*%
  decomp$x[1, seq_len(90)]
par(mfrow = c(1, 2))

matrix(mnist$train$images[1,], ncol = 28) %>%
  image(col = gray.colors(12, rev = TRUE),
        axes = FALSE, main = "Original")
matrix(approx_mnist, ncol = 28) %>%
  image(col = gray.colors(12, rev = TRUE),
        axes = FALSE, main = "Approx")
```

Data Visualization xiii

Original



Approx



Additional comments about sample PCA i

- Let $\mathbf{Y}_1, \dots, \mathbf{Y}_n$ be a sample from a distribution with covariance matrix Σ . Write \mathbb{Y} for the $n \times p$ matrix whose i -th row is \mathbf{Y}_i .
- Let S_n be the sample covariance matrix, and write W_k for the matrix whose columns are the first k eigenvectors of S_n .
- You can define the matrix of k principal components as

$$\mathbb{Z} = \mathbb{Y}W_k.$$

Additional comments about sample PCA ii

- On the other hand, it is much more common to define it as

$$\mathbb{Z} = \tilde{\mathbb{Y}}W_k,$$

where $\tilde{\mathbb{Y}}$ is the centered version of \mathbb{Y} (i.e. the sample mean has been subtracted from each row).

- This leads to sample principal components with mean zero.

Example 1 (revisited) i

```
library(mvtnorm)
Sigma <- matrix(c(1, 0.5, 0.1,
                  0.5, 1, 0.5,
                  0.1, 0.5, 1),
                 ncol = 3)
mu <- c(1, 2, 2)
```

Example 1 (revisited) ii

```
set.seed(17)
X <- rmvnorm(100, mean = mu,
              sigma = Sigma)
pca <- prcomp(X)
```

```
colMeans(X)
```

```
## [1] 0.8789229 2.0517403 2.0965127
```

```
colMeans(pca$x)
```

Example 1 (revisited) iii

```
##          PC1          PC2          PC3  
## -6.169544e-17 5.433154e-17 9.228729e-18
```

On the other hand

```
pca <- prcomp(X, center = FALSE)  
colMeans(pca$x)
```

```
##          PC1          PC2          PC3  
## 3.058960918 0.142358612 0.001050088
```

Geometric interpretation of PCA i

- The definition of PCA as a linear combination that maximises variance is due to Hotelling (1933).
- But PCA was actually introduced earlier by Pearson (1901)
 - *On Lines and Planes of Closest Fit to Systems of Points in Space*
- He defined PCA as the **best approximation of the data by a linear manifold**
- Let's suppose we have a lower dimension representation of \mathbb{Y} , denoted by a $n \times k$ matrix \mathbb{Z} .

Geometric interpretation of PCA ii

- We want to *reconstruct* \mathbb{Y} using an affine transformation

$$f(z) = \mu + W_k z,$$

where W_k is a $p \times k$ matrix.

- We want to find μ, W_k, \mathbf{Z}_i that minimises the **reconstruction error**:

$$\min_{\mu, W_k, \mathbf{Z}_i} \sum_{i=1}^n \|\mathbf{Y}_i - \mu - W_k \mathbf{Z}_i\|^2.$$

Geometric interpretation of PCA iii

- First, treating W_k constant and minimising over μ, \mathbf{Z}_i , we find

$$\begin{aligned}\hat{\mu} &= \bar{\mathbf{Y}}, \\ \hat{\mathbf{Z}}_i &= W_k^T(\mathbf{Y}_i - \bar{\mathbf{Y}}).\end{aligned}$$

- Putting these quantities into the reconstruction error, we get

$$\min_{W_k} \sum_{i=1}^n \|(\mathbf{Y}_i - \bar{\mathbf{Y}}) - W_k W_k^T (\mathbf{Y}_i - \bar{\mathbf{Y}})\|^2.$$

Geometric interpretation of PCA iv

Eckart–Young theorem

The reconstruction error is minimised by taking W_k to be the matrix whose columns are the first k eigenvectors of the sampling covariance matrix S_n .

Equivalently, we can take the matrix whose columns are the first k *right singular vectors* or the centered data matrix $\tilde{\mathbb{Y}}$.

Example i

```
set.seed(1234)
# Random measurement error
sigma <- 5

# Exact relationship between
# Celsius and Fahrenheit
temp_c <- seq(-40, 40, by = 1)
temp_f <- 1.8*temp_c + 32
```

Example ii

```
# Add measurement error  
temp_c_noise <- temp_c + rnorm(n = length(temp_c),  
                                  sd = sigma)  
temp_f_noise <- temp_f + rnorm(n = length(temp_f),  
                                  sd = sigma)
```

```
# Linear model
```

```
(fit <- lm(temp_f_noise ~ temp_c_noise))
```

Example iii

```
##  
## Call:  
## lm(formula = temp_f_noise ~ temp_c_noise)  
##  
## Coefficients:  
## (Intercept)  temp_c_noise  
##           34.256          1.662  
  
confint(fit)
```

Example iv

```
##                      2.5 %    97.5 %
## (Intercept) 32.152891 36.35921
## temp_c_noise 1.577228 1.74711
```

PCA

```
pca <- prcomp(cbind(temp_c_noise, temp_f_noise))
pca$rotation
```

```
##                  PC1          PC2
## temp_c_noise 0.5012360 -0.8653106
## temp_f_noise 0.8653106  0.5012360
```

Example v

```
pca$rotation[2,"PC1"] / pca$rotation[1,"PC1"]
```

```
## [1] 1.726354
```

Large sample inference i

- If we impose distributional assumption on the data \mathbf{Y} , we can derive the sampling distributions of the sample principal components.
- Assume $\mathbf{Y} \sim N_p(\mu, \Sigma)$, with Σ positive definite. Let $\lambda_1 > \dots > \lambda_p$ be the eigenvalues of Σ ; in particular we assume they are *distinct*. Finally let w_1, \dots, w_p be the corresponding eigenvectors.
- Given a random sample of size n , let S_n be the sample covariance matrix, $\hat{\lambda}_1, \dots, \hat{\lambda}_p$ its eigenvalues, and $\hat{w}_1, \dots, \hat{w}_p$ the corresponding eigenvectors.

Large sample inference ii

- Define Λ to be the diagonal matrix whose entries are $\lambda_1, \dots, \lambda_p$, and define

$$\Omega_i = \lambda_i \sum_{k=1, k \neq i}^p \frac{\lambda_k}{(\lambda_k - \lambda_i)^2} w_k w_k^T.$$

Large sample inference iii

Asymptotic results

1. Write $\lambda = (\lambda_1, \dots, \lambda_p)$ and similarly for $\hat{\lambda}$. As $n \rightarrow \infty$, we have

$$\sqrt{n} (\hat{\lambda} - \lambda) \rightarrow N_p(0, 2\Lambda^2).$$

2. As $n \rightarrow \infty$, we have

$$\sqrt{n} (\hat{w}_i - w_i) \rightarrow N_p(0, \Omega_i).$$

3. Each $\hat{\lambda}_i$ is distributed independently of \hat{w}_i .

Comments i

- These results **only** apply to principal components derived from the covariance matrix.
 - Some asymptotic results are available for those derived from the correlation matrix, but we will not cover them in class.
- Asymptotically, all eigenvalues of S_n are independent.
- You can get a confidence interval for λ_i as follows:

$$\frac{\hat{\lambda}_i}{(1 + z_{\alpha/2}\sqrt{2/n})} \leq \lambda_i \leq \frac{\hat{\lambda}_i}{(1 - z_{\alpha/2}\sqrt{2/n})}.$$

Comments ii

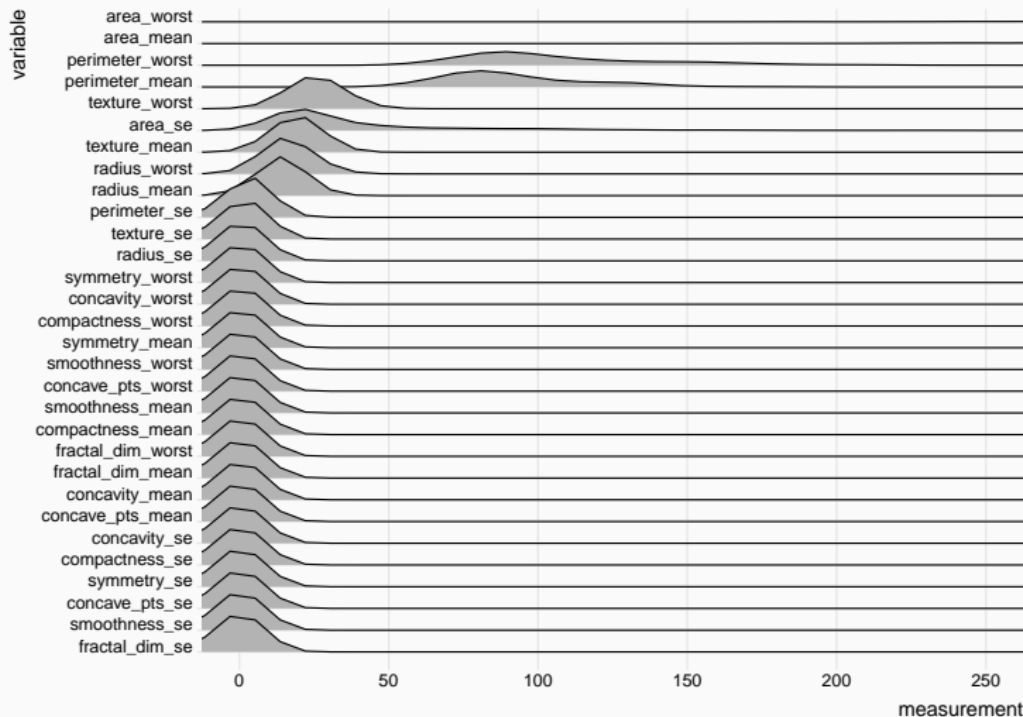
- Use Bonferroni correction if you want CIs that are simultaneously valid for all eigenvalues.
- The matrices Ω_i have rank $p - 1$, and therefore they are *singular*.
- The entries of \hat{w}_i are correlated, and this correlation depends on the *separation* between the eigenvalues.
 - Good separation \implies smaller correlation

Example i

```
library(dslabs)
library(ggridges)

# Data on Breast Cancer
as.data.frame(brca$x) %>%
  gather(variable, measurement) %>%
  mutate(variable = reorder(variable, measurement,
                            median)) %>%
  ggplot(aes(x = measurement, y = variable)) +
  geom_density_ridges() + theme_ridges() +
  coord_cartesian(xlim = c(0, 250))
```

Example ii



Example iii

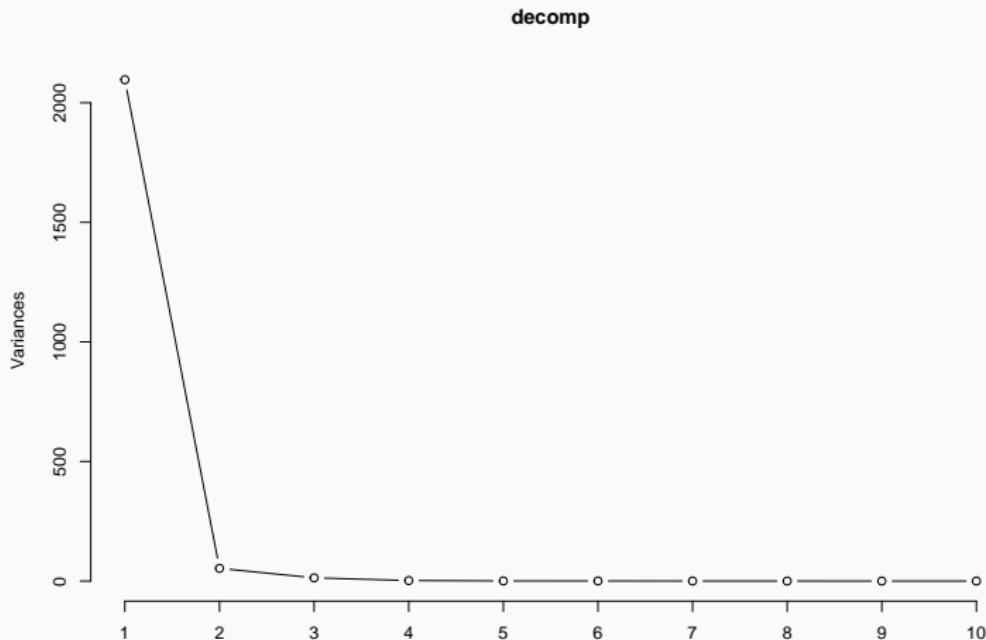
```
# Remove some variables
rem_index <- which(colnames(brca$x) %in%
                     c("area_worst", "area_mean",
                       "perimeter_worst",
                       "perimeter_mean"))
dataset <- brca$x[,-rem_index]
decomp <- prcomp(dataset)
summary(decomp)$importance[,1:3]
```

Example iv

```
##                                     PC1        PC2        PC3
## Standard deviation      45.78445 7.281664 3.677815
## Proportion of Variance 0.96776 0.024480 0.006240
## Cumulative Proportion  0.96776 0.992240 0.998490
```

```
screeplot(decomp, type = 'l')
```

Example v



Example vi

```
# Let's put a CI around the first eigenvalue
first_ev <- decomp$sdev[1]^2
n <- nrow(dataset)

# Recall that TV = 2166
c("LB" = first_ev/(1+qnorm(0.975)*sqrt(2/n)),
  "Est." = first_ev,
  "UP" = first_ev/(1-qnorm(0.975)*sqrt(2/n)))

##          LB      Est.          UP
## 1877.992 2096.216 2371.822
```

Simulations i

```
B <- 1000; n <- 100; p <- 3

results <- purrr::map_df(seq_len(B), function(b) {
  X <- matrix(rnorm(p*n, sd = sqrt(c(1, 2, 3))),  

               ncol = p, byrow = TRUE)
  tmp <- eigen(cov(X), symmetric = TRUE,  

               only.values = TRUE)
  tibble(ev1 = tmp$values[1],  

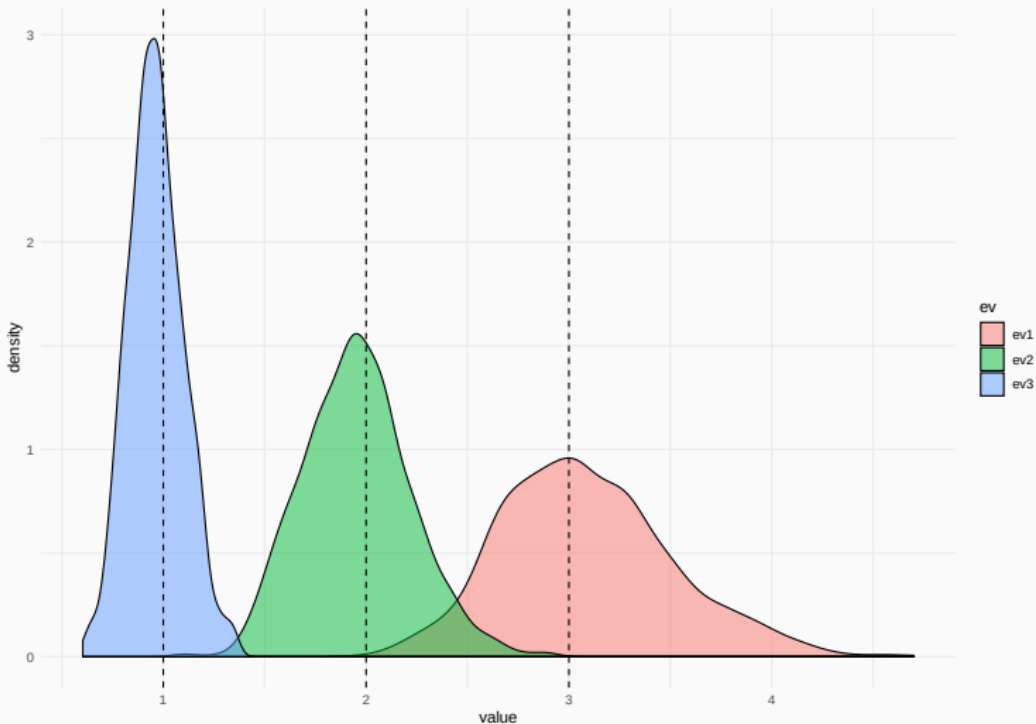
         ev2 = tmp$values[2],  

         ev3 = tmp$values[3])
})
```

Simulations ii

```
results %>%
  gather(ev, value) %>%
  ggplot(aes(value, fill = ev)) +
  geom_density(alpha = 0.5) +
  theme_minimal() +
  geom_vline(xintercept = c(1, 2, 3),
             linetype = 'dashed')
```

Simulations iii



Simulations iv

```
results %>%
  summarise_all(mean)

## # A tibble: 1 x 3
##       ev1     ev2     ev3
##   <dbl> <dbl> <dbl>
## 1  3.09  1.95  0.963
```

Simulations v

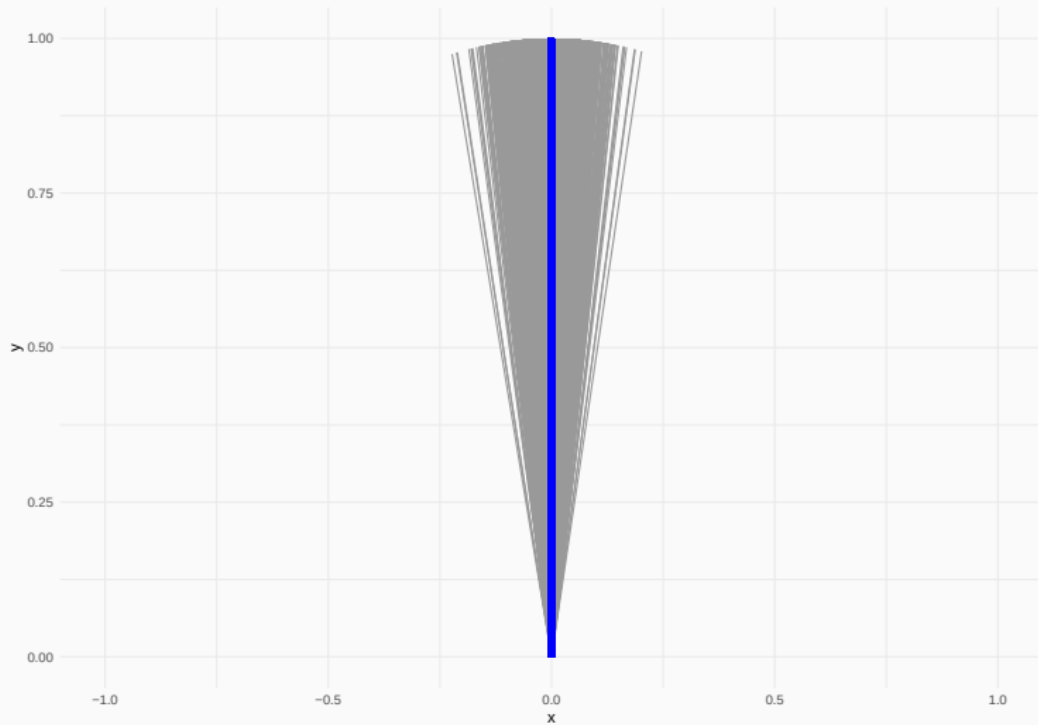
```
p <- 2
results <- purrr::map_df(seq_len(B), function(b) {
  X <- matrix(rnorm(p*n, sd = c(1, 2)), ncol = p,
              byrow = TRUE)
  tmp <- eigen(cov(X), symmetric = TRUE)

  tibble(
    xend = tmp$vectors[1,1],
    yend = tmp$vectors[2,1]
  )
})
```

Simulations vi

```
results %>%
  ggplot() +
  geom_segment(aes(xend = xend, yend = yend),
               x = 0, y = 0, colour = 'grey60') +
  geom_segment(x = 0, xend = 0,
               y = 0, yend = 1,
               colour = 'blue', size = 2) +
  expand_limits(y = 0, x = c(-1, 1)) +
  theme_minimal()
```

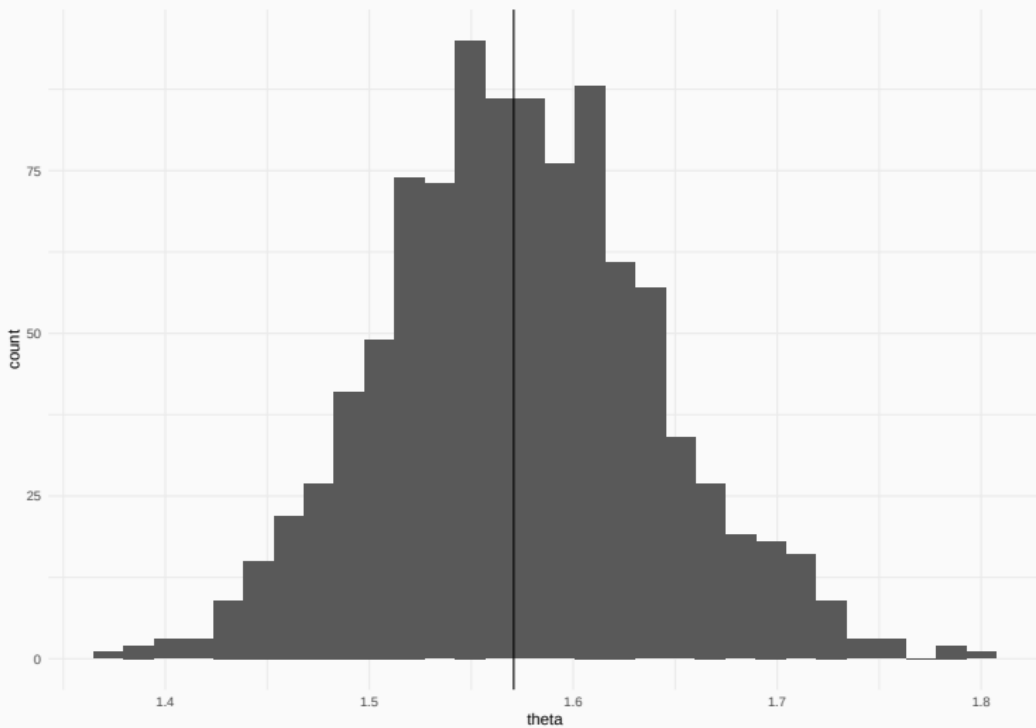
Simulations vii



Simulations viii

```
# Or looking at angles
results %>%
  transmute(theta = atan2(yend, xend)) %>%
  ggplot(aes(theta)) +
  geom_histogram() +
  theme_minimal() +
  geom_vline(xintercept = pi/2)
```

Simulations ix



Test for structured covariance i

- The asymptotic results above assumed distinct eigenvalues.
- But we may be interested in *structured* covariance matrices; for example:

$$\Sigma_0 = \sigma^2 \begin{pmatrix} 1 & \rho & \cdots & \rho \\ \rho & 1 & \cdots & \rho \\ \vdots & \vdots & \ddots & \vdots \\ \rho & \rho & \cdots & 1 \end{pmatrix}.$$

- This is called an **exchangeable** correlation structure.

Test for structured covariance ii

- Assuming $\rho > 0$, the eigenvalues of Σ_0 are

$$\lambda_1 = \sigma^2(1 + (p - 1)\rho),$$

$$\lambda_2 = \sigma^2(1 - \rho),$$

$$\vdots \quad \vdots$$

$$\lambda_p = \sigma^2(1 - \rho).$$

- Let's assume $\sigma^2 = 1$. We are interested in testing whether the correlation matrix is equal to Σ_1 .

Test for structured covariance iii

- Let $\bar{r}_k = \frac{1}{p-1} \sum_{i=1, i \neq k}^p r_{ik}$ be the average of the off-diagonal value of the k -th column of the sample correlation matrix.
- Let $\bar{r} = \frac{2}{p(p-1)} \sum_{i < j} r_{ij}$ be the average of all off-diagonal elements (we are only looking at entries below the diagonal).
- Finally, let $\hat{\gamma} = \frac{(p-1)^2[1-(1-\bar{r})^2]}{p-(p-2)(1-\bar{r})^2}$.
- We reject the null hypothesis that the correlation matrix is equal to Σ_0 if

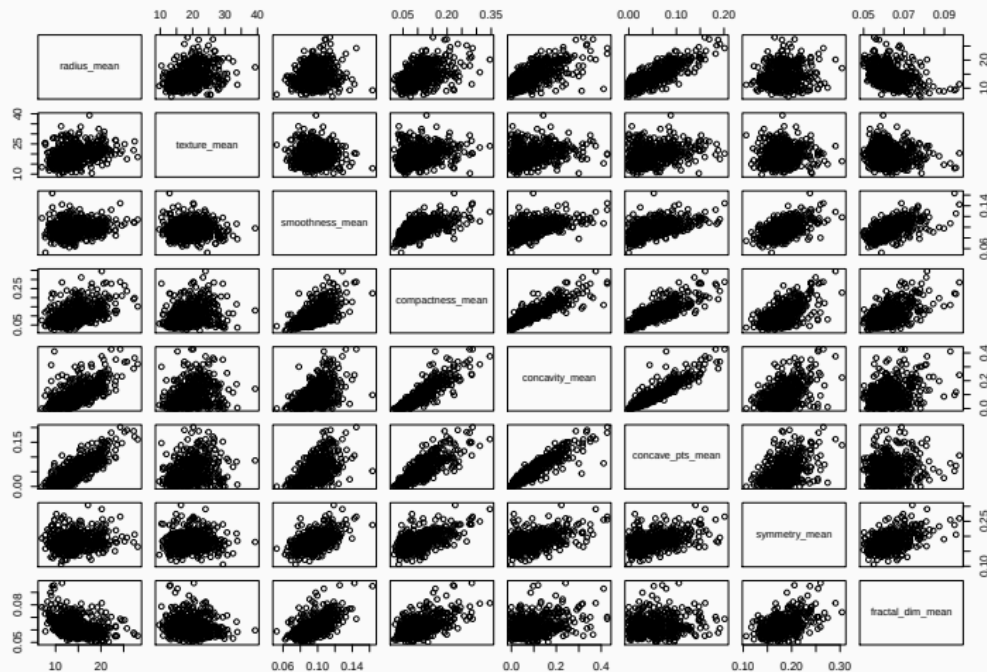
$$\frac{(n-1)}{(1-\bar{r})^2} \left[\sum_{i < j} (r_{ij} - \bar{r})^2 - \hat{\gamma} \sum_{k=1}^p (\bar{r}_k - \bar{r})^2 \right] > \chi_{\alpha}^2((p+1)(p-2)/2)$$

Example i

```
# Keep only mean measurements
rem_index2 <- grep("mean", colnames(brca$x),
                     invert = TRUE)
dataset <- brca$x[,-c(rem_index,
                      rem_index2)]
R <- cor(dataset)

pairs(dataset)
```

Example ii



Example iii

```
# Overall mean  
r_bar <- mean(R[upper.tri(R, diag = FALSE)])  
  
# Column specific means  
r_cols <- (colSums(R) - 1)/(nrow(R) - 1)  
  
# Extra quantities  
p <- ncol(dataset)  
n <- nrow(dataset)  
gamma_hat <- (p - 1)^2*(1 - (1 - r_bar)^2)/  
(p - (p - 2)*(1 - r_bar)^2)
```

Example iv

```
# Test statistic
Tstat <- sum((R[upper.tri(R,
                           diag = FALSE)] - r_bar)^2) -
  gamma_hat*sum((r_cols - r_bar)^2)
Tstat <- (n-1)*Tstat/(1-r_bar)^2

Tstat > qchisq(0.95, 0.5*(p+1)*(p-2))

## [1] TRUE
```

Selecting the number of PCs i

- We already discussed two strategies for selecting the number of principal components:
 - Look at the scree plot and find where the curve starts to be flat;
 - Retain as many PCs as required to explain the desired proportion of variance.

Selecting the number of PCs ii

- There is a **vast** literature on different strategies for selecting the number of components. Two good references:
 - Peres-Neto *et al.* (2005) *How many principal components? stopping rules for determining the number of non-trivial axes revisited*
 - Jolliffe (2012) *Principal Component Analysis* (2nd ed)
- We will discuss one more technique based on resampling.
- The idea is to try to estimate the distribution of eigenvalues if there was no correlation between the variables.

Selecting the number of PCs iii

Algorithm

1. Permute the observations of each column **independently**.
2. Perform PCA on the permuted data.
3. Repeat B times and collect the eigenvalues $\hat{\lambda}_1^{(b)}, \dots, \hat{\lambda}_p^{(b)}$.
4. Keep the components whose observed $\hat{\lambda}_i$ is greater than $(1 - \alpha)\%$ of the values $\hat{\lambda}_i^{(b)}$ obtained through permutations.

Example (cont'd) i

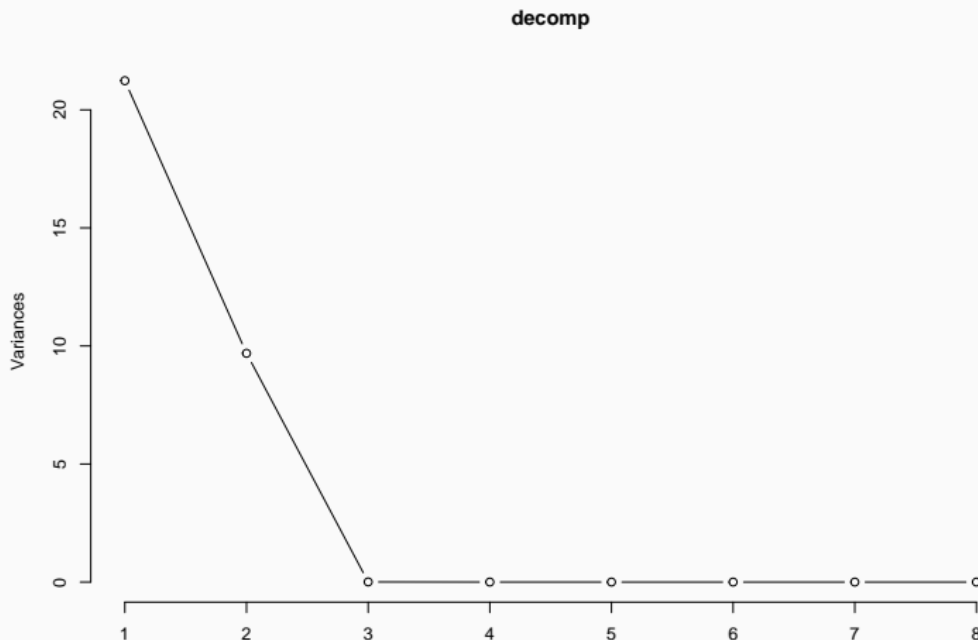
```
decomp <- prcomp(dataset)  
  
summary(decomp)$importance[, seq_len(3)]
```

```
##                                     PC1        PC2        PC3  
## Standard deviation      4.60806 3.112611 0.07664969  
## Proportion of Variance 0.68654 0.313240 0.00019000  
## Cumulative Proportion  0.68654 0.999780 0.99997000
```

Example (cont'd) ii

```
screeplot(decomp, type = 'l')
```

Example (cont'd) iii



Example (cont'd) iv

```
permute_data <- function(data) {  
  p <- ncol(data)  
  data_perm <- data  
  for (i in seq_len(p)) {  
    ind_sc <- sample(nrow(data))  
    data_perm[,i] <- data[ind_sc, i]  
  }  
  return(data_perm)  
}
```

Example (cont'd) v

```
set.seed(123)
B <- 1000
alpha <- 0.05
results <- matrix(NA, ncol = B,
                   nrow = ncol(dataset))

results[, 1] <- decomp$sdev
results[,-1] <- replicate(B - 1, {
  data_perm <- permute_data(dataset)
  prcomp(data_perm)$sdev
})
```

Example (cont'd) vi

```
cutoff <- apply(results, 1, function(row) {  
  mean(row >= row[1])  
})  
which(cutoff < alpha)  
  
## [1] 1
```

Biplots i

- In our example with the MNIST dataset, we plotted the first principal component against the second component.
 - This gave us a sense of how much discriminatory ability each PC gave us.
 - E.g. the first PC separated 1s from 0s
- What was missing from that plot was how the PCs were related to the original variables.
- A **biplot** is a graphical display of both the original observations and original variables *together* on one scatterplot.

Biplots ii

- The prefix “bi” refers to two modalities (i.e. observations and variables), not to two dimensions.
- One approach to biplots relies on the Eckart-Young theorem:
 - The “best” 2-dimensional representation of the data passes through the plane containing the first two eigenvectors of the sample covariance matrix.

Biplots iii

Construction

- Let $\tilde{\mathbb{Y}}$ be the $n \times p$ matrix of centered data, and let w_1, \dots, w_p be the p eigenvectors of $\tilde{\mathbb{Y}}^T \tilde{\mathbb{Y}}$.
- For each row \mathbf{Y}_i of \mathbb{Y} , add the point $(w_1^T \mathbf{Y}_i, w_2^T \mathbf{Y}_i)$ to the plot.
- The j -th column of \mathbb{Y} is represented by an arrow from the origin to the point (w_{1j}, w_{2j}) .
- It may be necessary to rescale the PCs and/or the loadings in order to see the relationship better.

Example (cont'd) i

```
# Continuing with our example on breast cancer
decomp <- prcomp(dataset)

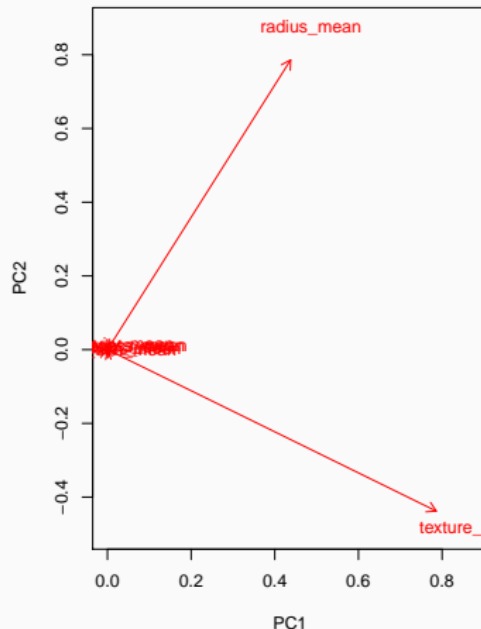
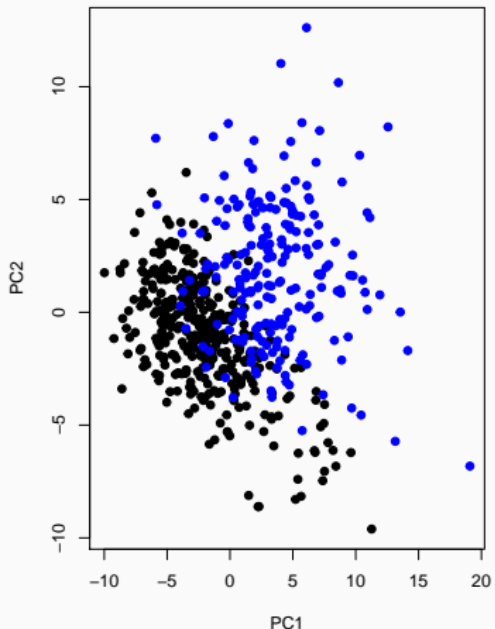
# Extract PCs and loadings
PCs <- decomp$x[, 1:2]
loadings <- decomp$rotation[, 1:2]

# Extract data on tumour type
colour <- ifelse(brca$y == "B", "black", 'blue')
```

Example (cont'd) ii

```
par(mfrow = c(1,2))
plot(PCs, pch = 19, col = colour)
plot(loadings, type = 'n')
text(loadings,
      labels = colnames(dataset),
      col = 'red')
arrows(0, 0, 0.9 * loadings[, 1],
       0.9 * loadings[, 2],
       col = 'red',
       length = 0.1)
```

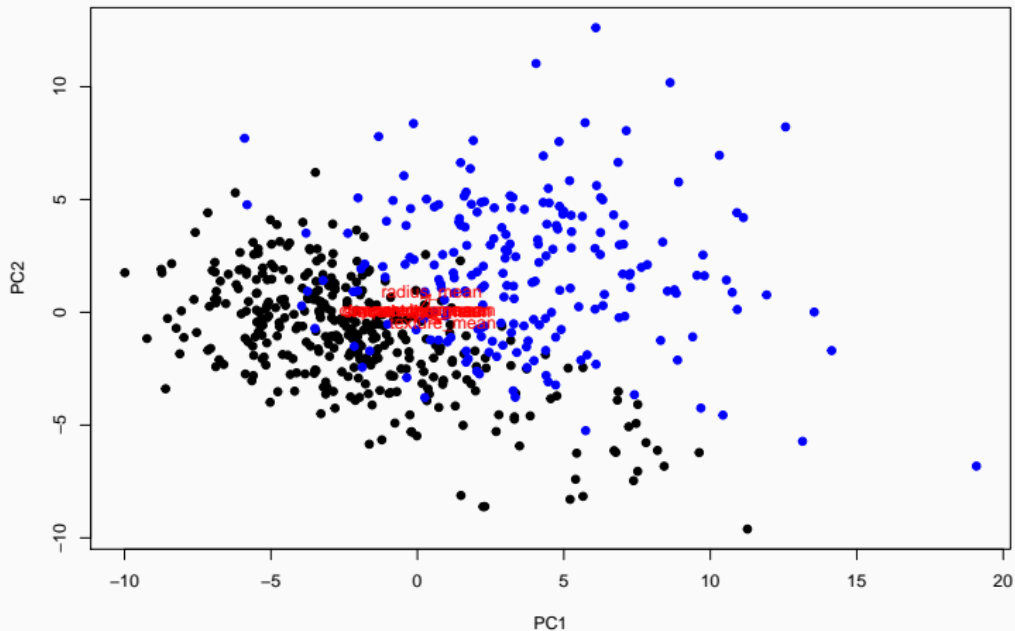
Example (cont'd) iii



Example (cont'd) iv

```
# Or both on the same plot
plot(PCs, pch = 19, col = colour)
text(loadings,
      labels = colnames(dataset),
      col = 'red')
arrows(0, 0, 0.9 * loadings[, 1],
       0.9 * loadings[, 2],
       col = 'red',
       length = 0.1)
```

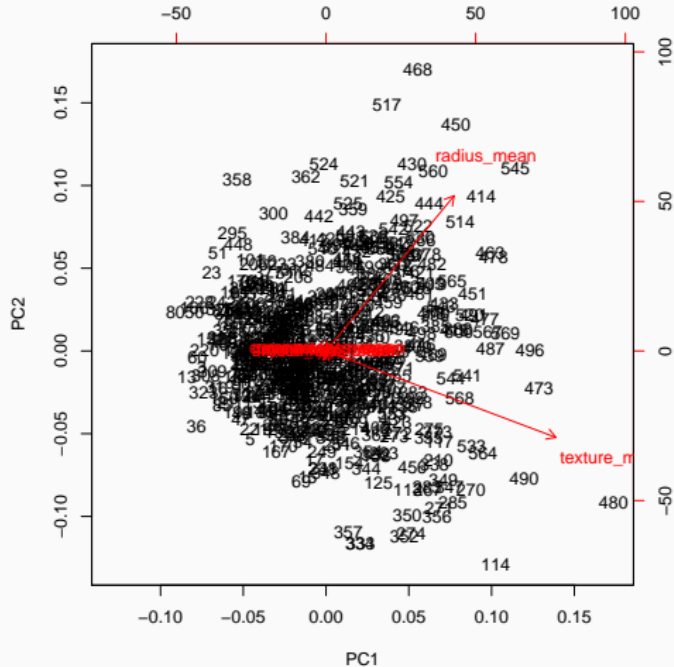
Example (cont'd) v



Example (cont'd) vi

```
# The biplot function rescales for us  
biplot(decomp)
```

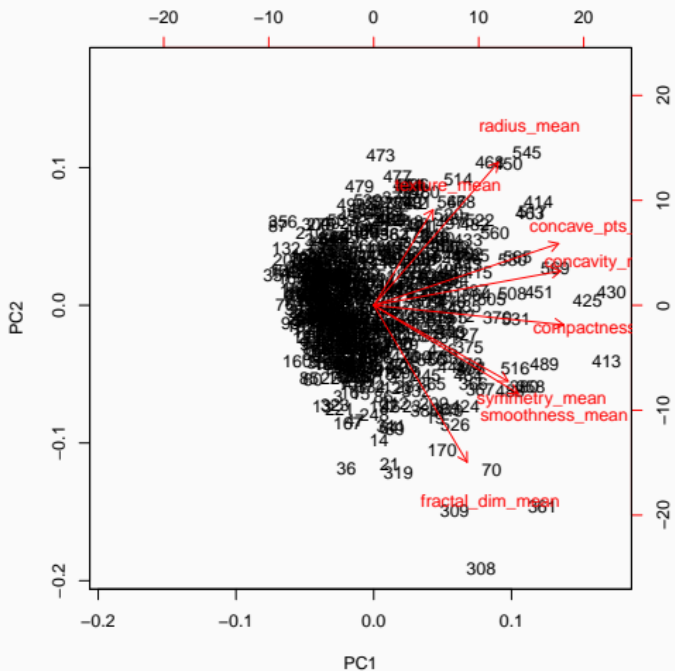
Example (cont'd) vii



Example (cont'd) viii

```
# With scaled data  
biplot(prcomp(dataset, scale = TRUE))
```

Example (cont'd) ix



Summary of graphical displays

- When we plot the first PC against the second PC, we are looking for similarity between *observations*.
- When we plot the first loading against the second loading, we are looking for similarity between *variables*.
 - Orthogonal loadings \implies Uncorrelated variables
 - Obtuse angle between loadings \implies Negative correlation
- A **biplot** combines both pieces of information.
 - You can think of it as a projection of the p -dimensional scatter plot (points and axes) onto a 2-dimensional plane.
- A **scree plot** displays the amount of variation in each principal component.