# Principal components analysis

Reference: Introduction to Statistical Learning Chapter 10.1-10.2

### Outline

- I. Introduction to PCA
- 2. Brief reminder of some linear algebra notation
- 3. PCA as a projection
- 4. PCA as a regression on latent variables

#### Introduction to PCA

The goal of PCA is to find low-dimensional summaries of high-dimensional data sets.

This is useful for compression, for denoising, for plotting, and for making sense of data sets that initially seem too complicated to understand.

#### It differs from clustering:

- Clustering assumes that each data point is a member of one, and only one, cluster. (Clusters are mutually exclusive.)
- PCA assumes that each data point is like a combination of multiple basic "ingredients." (Ingredients are not mutually exclusive.)

## Think about recipes:

Nestle Toll House Chocolate-chip cookies: 280 grams flour, 150 grams white sugar, 165 grams brown sugar, 225 grams butter, 2 eggs, 0 grams water...

Mary Berry's Victoria sponge cake: 225 grams flour, 225 grams white sugar, 0 grams brown sugar, 225 grams butter, 4 eggs, 0 grams water...

seriouseats.com old fashioned flaky pie dough: 225 grams flour, 15 grams white sugar, 0 grams brown sugar, 225 grams butter, 0 eggs, 115 grams water...

## Think about recipes:

Each baked good is constructed by following a recipe: a combination of the same basic ingredients.

- Each data point  $x_i$  is like a baked good.
- In PCA, the **principal components** are like the ingredients.

The amounts of each ingredient differ from one baked good to the next:

- E.g. 225g sugar for sponge cake versus 15 grams sugar for pie dough.
- In PCA, the **scores** are like the amounts of each ingredient in a given baked good.

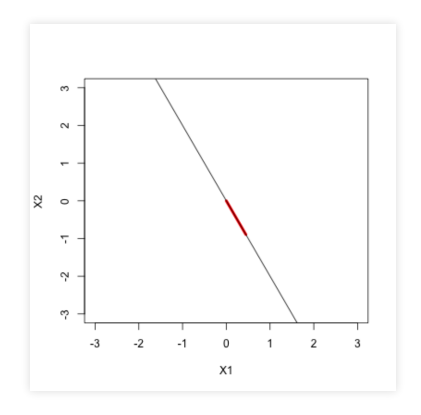
## Think about recipes:

Our goal is to reverse-engineer both the **ingredients** and the **amounts/recipes** from an observed set of "baked goods" (i.e. original data points).

# Some linear algebra reminders

Alas, PCA is less delicious than baking, and it uses more linear algebra. Say that  $v \in \mathbb{R}^P$  is some vector. This defines a *subspace* of  $\mathbb{R}^P$ :

$$\mathcal{V} = \{z : z = \alpha_i v, \alpha_i \in \mathbb{R}\}\$$



# Some linear algebra reminders

Now let X be our usual  $N \times P$  data matrix with rows  $x_i^T$ .

Suppose we project each  $x_i^T$  in our data matrix onto the subspace  $\mathcal{V}$ .

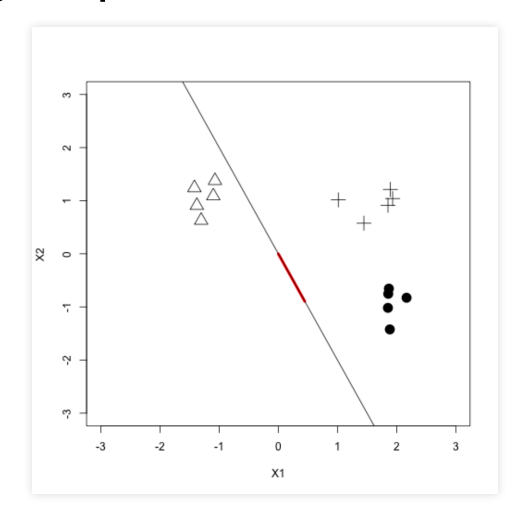
• The scalar location along the vector v is

$$\alpha_i = x_i \cdot v = x_i^T v$$

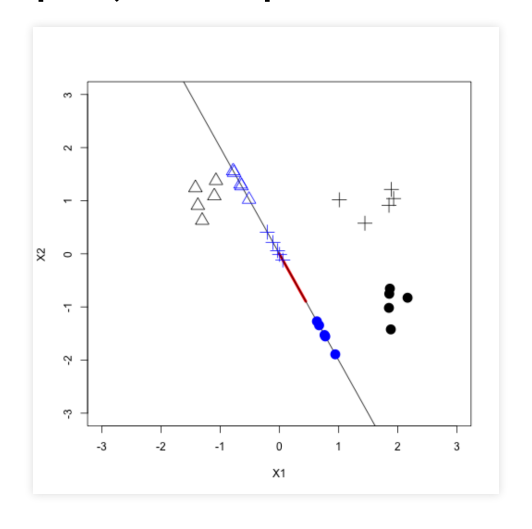
• The location of the projected point in  $\mathcal{R}^P$  is

$$\hat{x_i} = \alpha_i v = (x_i \cdot v)v$$

# The original points



# With the projected points



# Key ideas

Key idea 1: projection = summary

• Each point's location along the subspace is a **one-number linear summary** of a *P*-dimensional data vector:

$$\alpha_i = x_i \cdot v = x_i^T v$$

• The goal of principal components analysis (PCA) is to find the "best" projection, i.e. the best linear summary of the data points.

Key idea 2: the "best summary" is the one that preserves as much of the **variance** in the original data points as possible.

Let's walk through pca intro.R

# PCA as an optimization problem

Given data points  $x_1, \ldots, x_N$ , with each  $x_i \in \mathbb{R}^P$ , and a candidate vector  $v_1$ , the variance of the projected points is

variance = 
$$\frac{1}{n} \sum_{i=1}^{n} [\alpha_i - \bar{\alpha}]^2$$

where  $\alpha_i x_i \cdot v_1$ .

So we solve:

Note: we constrain  $v_1$  to have length 1; otherwise we could blow up the variance of the projected points as large as we wanted to.

# PCA as an optimization problem

The solution  $v_1$  to this optimization problem:

- is called the first principal component (synonyms: loading, rotation.)
- is the one-dimensional subspace capturing as much of the information in the original data matrix as possible.

The projected points  $\alpha_i = x_i \cdot v$  are called the *scores* on the first principal component.

# PCA as regression on latent variables

We can think of the projected location of  $x_i$  as an approximation to the original data point:  $x_i \approx \hat{x}_i = \alpha_i v$ .

Or to make the approximation error explicit:

$$x_{ij} = \hat{x}_{ij} + e_i$$
$$= \alpha_i v_i + e_i$$

This is like a regression problem for the jth feature variable.

- The  $\alpha_i$ 's are like hidden (latent) features.
- $v_j$  is like a regression coefficient for observed variable j.

Thus PCA is like estimating P regression coefficients  $v_1 = (v_{11}, \dots, v_{1P})$  all at once, where the feature variable is hidden.

# PCA as regression on latent variables

We can write the approximation for the whole matrix as follows:

$$X \approx \begin{pmatrix} \alpha_1 v_{11} & \alpha_1 v_{12} & \cdots & \alpha_1 v_{1P} \\ \alpha_2 v_{11} & \alpha_2 v_{12} & \cdots & \alpha_2 v_{1P} \\ \vdots & \vdots & & \vdots \\ \alpha_N v_{11} & \alpha_N v_{12} & \cdots & \alpha_N v_{1P} \end{pmatrix}$$
$$= \alpha v_1^T \quad \text{(outer product of } \alpha \text{ and } v_1 \text{)}$$

# PCA as regression on latent variables

And if we explicitly include the error term:

$$X = \alpha v_1^T + E$$

where E is a residual matrix with entries

$$E_{ij} = x_{ij} - \hat{x}_{ij}$$
$$= x_{ij} - \alpha_i v_{1j}$$

# Higher-order principal components

With this in place, we can now define principal components 2 and up!

- PC 2: run PCA on the residual matrix from PC 1.
- PC 3: run PCA on the residual matrix from PCs I-2.
- ...
- PC P: run PCA on the residual matrix from PCs I-(P-I).

Thus principal component M is defined recursively in terms of the fit from principal components I through M-1.

# Higher-order principal components

Note: in practice we often stop with far fewer than P principal components.

Let's see the examples in congress109.R and NCI60.R.