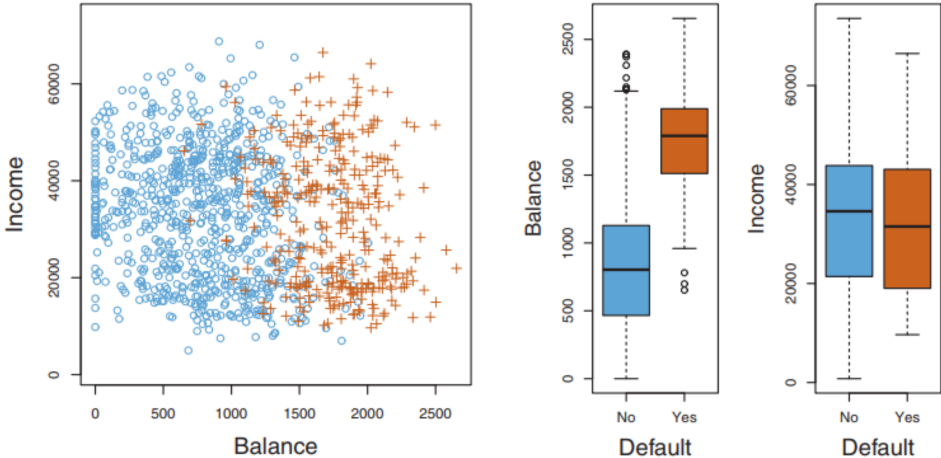
# Dimensionality Reduction Feature Selection & Extraction

Jing Sun



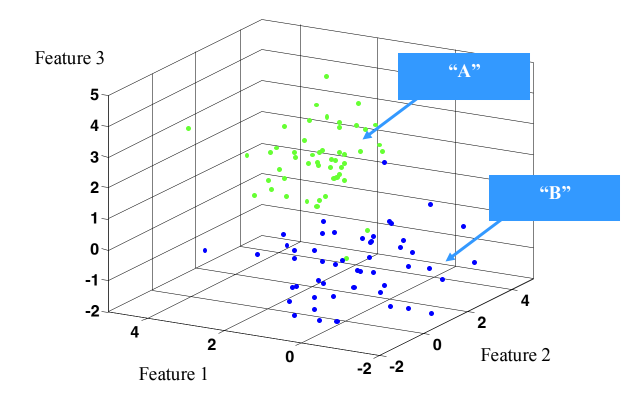
### Is "Income" an informative feature?





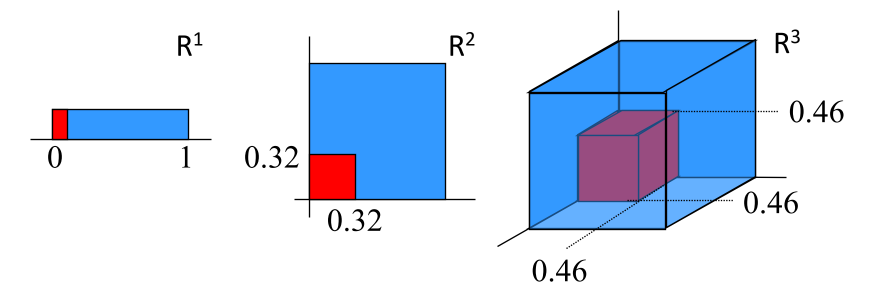
### Feature Space

- A p-dimensional space,
   in which each dimension is a feature
   containing n [labeled] samples [objects]
- What will happen if p is very large?
- [the curse of dimensionality]



In high-dimensional spaces, our 2D/3D intuition does not work anymore...





- Example:
- Neighborhood capturing 10% of uniformly distributed data in hypercube
- E.g. in  $\mathbb{R}^{20}$  side length of  $\sqrt[20]{.1} \approx 0.89$

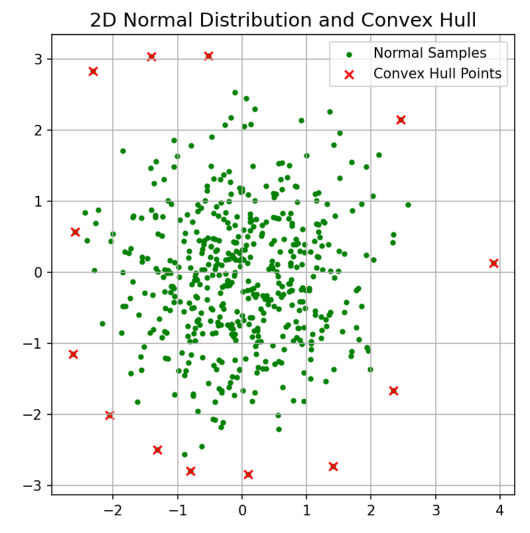
So, not a small block anymore...



Example: Boundary points ?

500 samples from normal distribution

In a 2-D space, only 2% are on the convex hull





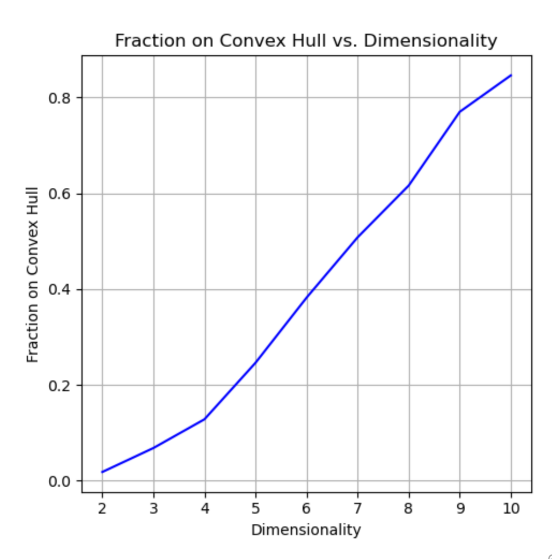
Example: Boundary points ?

500 samples from normal distribution

In a 2-D space, only 2% are on the convex hull

In a 20-D space, 95% are on the convex hull!



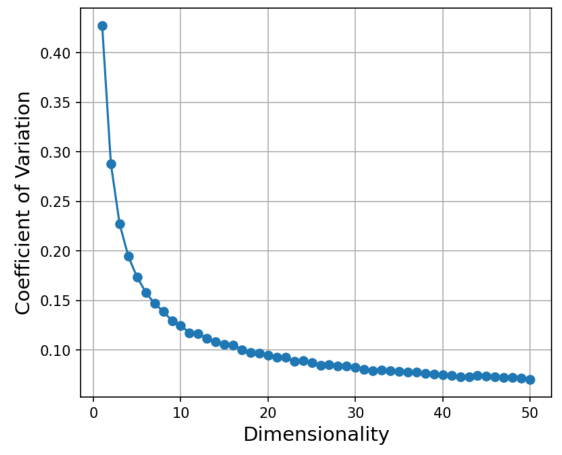


Example: Points tend to have equal distances

200 samples from normal distribution N(2000, 8000)

In a  $\mathbb{R}^1$  to  $\mathbb{R}^{1000}$  space

Consider  $\frac{\operatorname{std}(d^2)}{\operatorname{mean}(d^2)}$  for squared distance  $d^2$ 



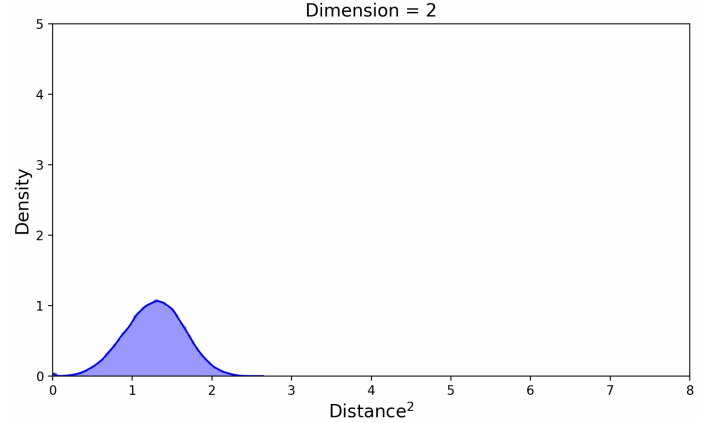


Example: Points tend to have equal distances

200 samples from normal distribution

N(2000,8000)

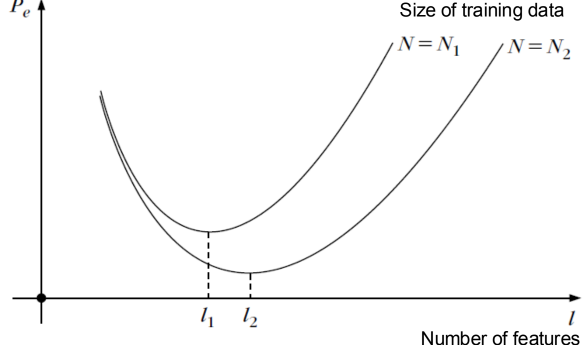
In a  $\mathbb{R}^1$  to  $\mathbb{R}^{1000}$  space





### **Dimensionality Reduction**

- Problem: too few samples in too many dimensions [the curse of dimensionality]
- Solution: drop dimensions / features
  - Feature selection
  - Feature extraction





### **Dimensionality Reduction**

- Uses/Benefits:
  - Fewer parameters give faster algorithms and parameters are easier to estimate

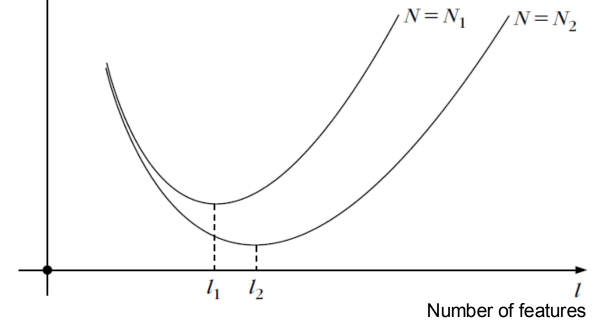
- Explaining which measurements are useful and which are not [reducing redundancy]
- Visualization of data can be a powerful tool when designing pattern recognition systems



### **Dimensionality Reduction**

Problem: too few samples in too many dimensions
 [the curse of dimensionality]

- Solution: drop dimensions / features
  - Feature selection
  - Feature extraction
- Questions:
  - Which dimensions to drop?
  - What feature subset to keep?





Size of training data

### Dimensionality Reduction by Selection or Extraction

Overview – Feature Selection vs Feature Extraction

- Criteria
  - Mahalanobis distance (vs Euclidean distance)
  - Scatter matrices (what are  $S_W$ ,  $S_B$ ,  $S_T$ ?)
- Approaches
  - Sequential **feature selection** (individual, forward, backward, etc.)
  - Principal Component Analysis & Recall LDA (∈ linear **feature extraction**)



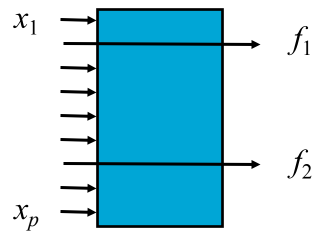
#### Feature Selection vs Extraction

Feature selection :

**Select** *d* **out of** *p* measurements

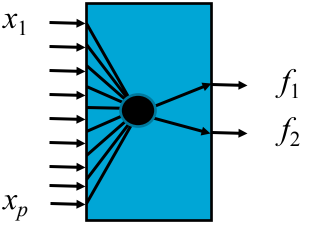
Only a subset of the original features are selected.

There are 
$$\binom{p}{d} = \frac{p!}{d!(p-d)!}$$
 subsets.



Feature extraction :

**Map** p measurements **to** d measurements All original features are used (they are transferred)





### Dimensionality Reduction by Selection or Extraction

Overview – Feature Selection vs Feature Extraction

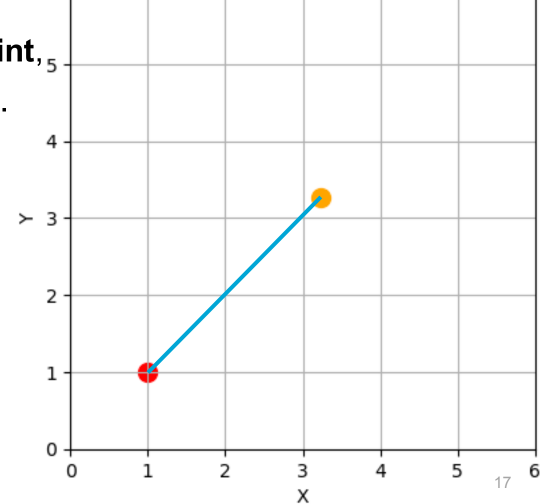
- Criteria
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### Why Mahalanobis distance?

• When measuring the distance from a single point to another single point, susing (squared) Euclidean distance is fine.

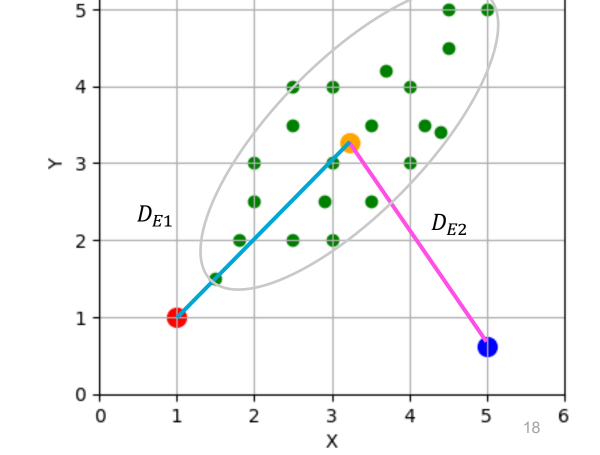
$$D_E = (x_{red} - x_{yellow})^2 + (y_{red} - y_{yellow})^2$$





### Why Mahalanobis distance?

- However,
- when there is a group of data points:
- Centroid (mean vector) =  $\begin{pmatrix} \bar{x} \\ \bar{y} \end{pmatrix}$
- Euclidean distances  $D_{E1} = D_{E2}$





#### Mahalanobis distance

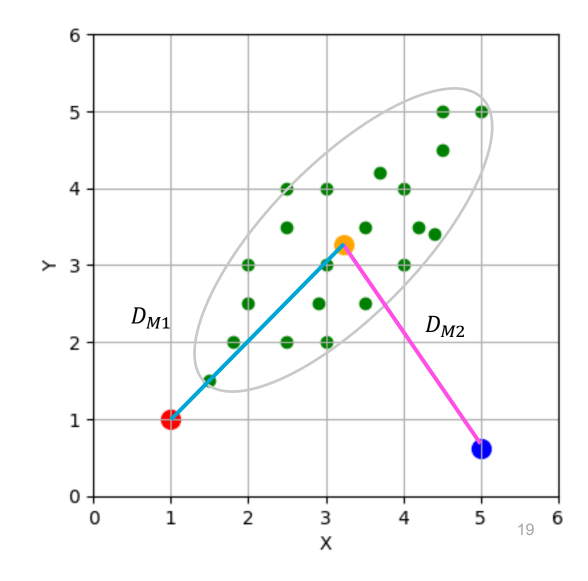
Takes the variance into account.

- It is a distance measure
   between a point and a distribution.
- For red and blue points,

$$D_{M} = \begin{pmatrix} x - \bar{x} \\ y - \bar{y} \end{pmatrix}^{T} \Sigma^{-1} \begin{pmatrix} x - \bar{x} \\ y - \bar{y} \end{pmatrix}$$

• You will see  $D_{M2} > D_{M1}$ 



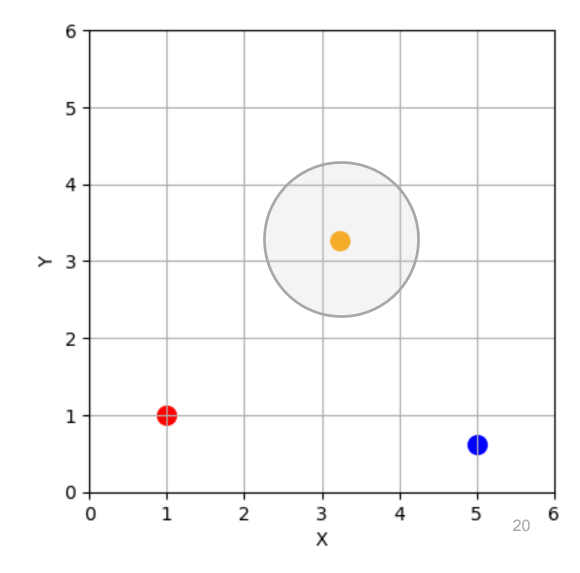


### Mahalanobis distance

- Think about:
- What if Σ is an identity matrix?

$$D_{M} = \begin{pmatrix} x - \bar{x} \\ y - \bar{y} \end{pmatrix}^{T} I \begin{pmatrix} x - \bar{x} \\ y - \bar{y} \end{pmatrix} = D_{E}$$





#### Mahalanobis distance

- Mahalanobis distance between two classes:
  - Assumes Gaussian distributions with equal covariance matrix

$$D_M = (\mu_1 - \mu_2)^T S_W^{-1} (\mu_1 - \mu_2)$$

- E.g., Exercise 6.21
- What is this  $S_W$ ?



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### **Scatter Matrices**

Within-class scatter matrix:

$$S_W = \sum_{i=1}^M \frac{n_i}{N} \Sigma_i$$
,  $\Sigma_i$  is the covariance matrix of class  $w_i$ ;  $M$  is the number of classes;  $n_i$  is the number of samples in class  $w_i$ , out of a total of  $N$  samples.

Between-class scatter matrix:

$$S_B = \sum_{i=1}^M \frac{n_i}{N} (\mu_i - \mu) (\mu_i - \mu)^T, \quad \mu_i \text{ is the mean of class } w_i, \\ \mu \text{ is the global mean.} \qquad \mu = \sum_{i=1}^M \frac{n_i}{N} \mu_i$$

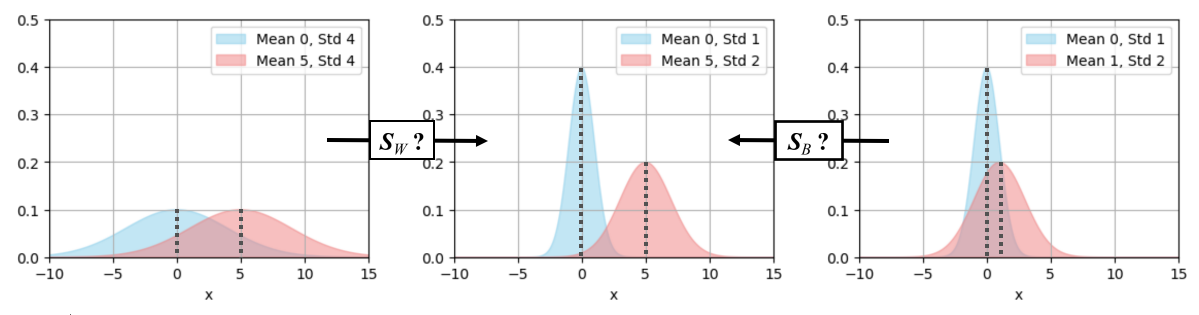
• Total scatter matrix:  $S_T = S_W + S_B$ 



### **Scatter Matrices**

#### For a classification task

- $S_W$  = "average class width"; **the smaller, the better**
- $S_B$  = "average distance between class means"; the larger, the better
- $S_T$  = "overall width"

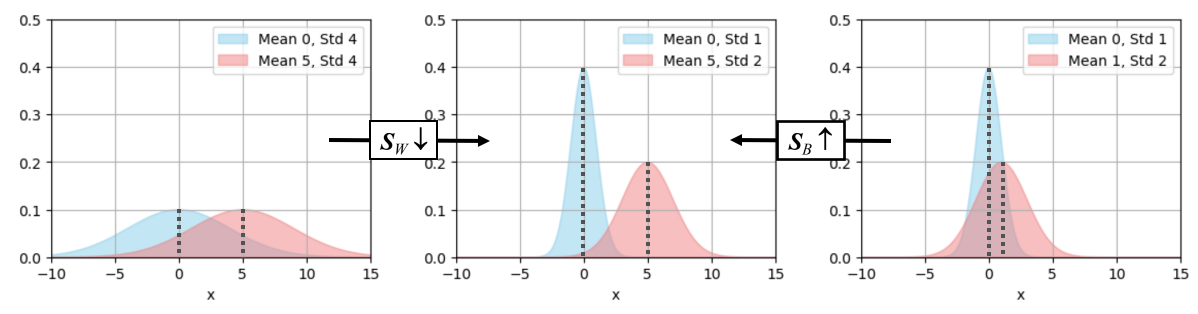




### **Scatter Matrices**

#### For a classification task

- $S_W$  = "average class width"; **the smaller, the better**
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- $S_T$  = "overall width"





#### Scatter-based Criteria

$$J_1 = \frac{trace\{S_T\}}{trace\{S_W\}}$$

$$J_2 = \frac{|S_T|}{|S_W|}$$

- etc.
- by using various combinations of  $S_W$ ,  $S_B$ ,  $S_T$  in a "trace" or "determinant" formulation...

 PS: The "trace" is equal to the sum of the eigenvalues; the "determinant" is equal to their product.



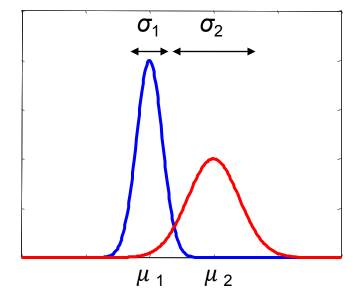
#### FDR: Fisher Discriminant Ratio

1-D, two-class problem

• 
$$S_W \propto (\sigma_1^2 + \sigma_2^2), S_B \propto (\mu_1 - \mu_2)^2,$$

• Combining  $S_W$  and  $S_B$ , you get Fisher's criterion

$$J_F = \frac{(\mu_1 - \mu_2)^2}{\sigma_1^2 + \sigma_2^2}$$



It is often used to quantify the separability capabilities of individual features.



### Dimensionality Reduction by Selection or Extraction

Overview – Feature Selection vs Feature Extraction

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- Approaches
  - Sequential feature selection (individual, forward, backward, etc.)
  - Principal Component Analysis & Recall LDA (∈ linear **feature extraction**)



### Which method would guarantee optimal performance?

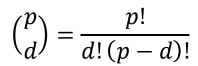
- I have p features (let's say p = 40).
- I think this is too many to handle...
- I want to select d features from p.
- But what should d = ? Well, I'm not sure...

• What can I do?



## Which method would guarantee optimal performance?

Trying all possible feature combinations



Exhaustive feature selection

$$\sum_{i=1}^{p} {p \choose i}$$
 combinations

If originally there are 4 features, we will end up with 15 combinations.

• 
$$\binom{4}{1} + \binom{4}{2} + \binom{4}{3} + \binom{4}{4} = 15$$

But, what if there are 40 features...?





### **Sub-optimal Strategies**

- Trying all possible feature combinations
- Exhaustive feature selection

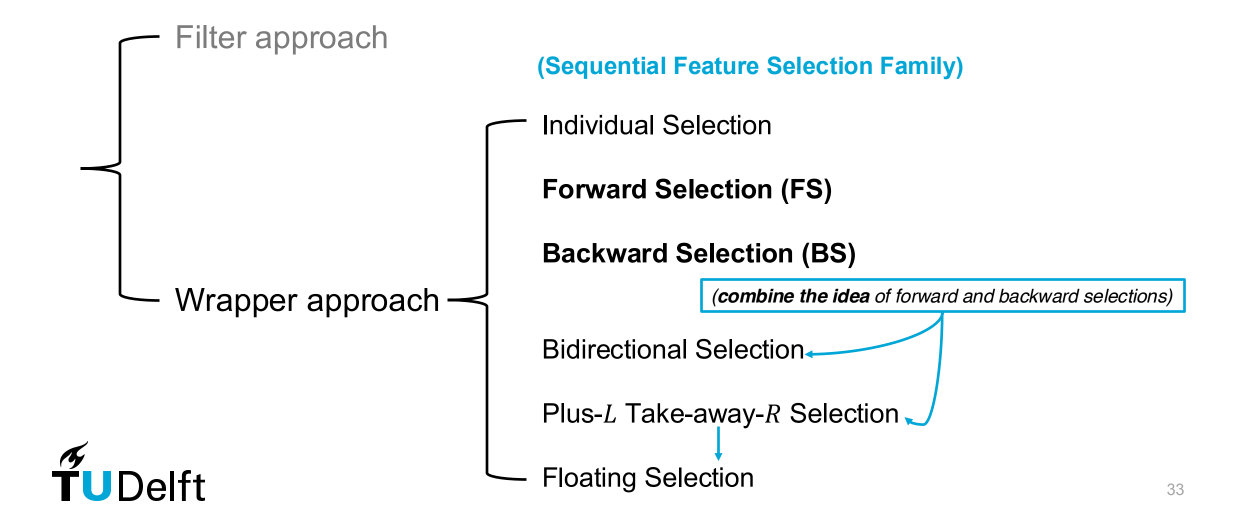
- It can be super Expensive! And Exhaustive!! :-(
- Let's use Sequential Feature Selection!

$$\binom{p}{d} = \frac{p!}{d! (p-d)!}$$

$$\sum_{i=1}^{p} {p \choose i}$$
 combinations



#### **Feature Selection Methods**



Start with empty feature set





Start with empty feature set



Compute the criterion value for each feature individually and select the best one,



Start with empty feature set



Compute the criterion value for each feature individually and select the best one,

$$x_2 > x_4 > x_2 > x_3 \implies x_2$$



Start with empty feature set



Compute the criterion value for each feature individually and select the best one,

$$x_2 > x_4 > x_2 > x_3 \implies x_2$$

 Keep the winner and compute the criterion value for all two-feature combinations that include it.



Start with empty feature set



Compute the criterion value for each feature individually and select the best one,

$$x_2 > x_4 > x_2 > x_3 \implies x_2$$

 Keep the winner and compute the criterion value for all two-feature combinations that include it.

$$[x_2, x_1] > [x_2, x_4] > [x_2, x_3]$$



Start with empty feature set



Compute the criterion value for each feature individually and select the best one,

$$x_2 > x_4 > x_2 > x_3 \implies x_2$$

Keep the winner and compute the criterion value for all two-feature combinations that include it.

$$[x_2, x_1] > [x_2, x_4] > [x_2, x_3]$$

... until a predefined number of features are left.



Start with all originally available features





Start with all originally available features



Compute the criterion value for all possible combinations after eliminating one feature,



Start with all originally available features



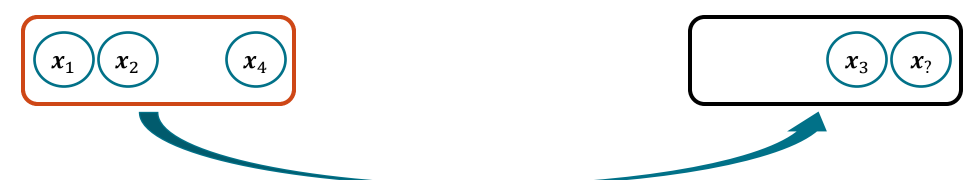
Compute the criterion value for all possible combinations after eliminating one feature,

$$[x_1, x_2, x_4] > [x_1, x_2, x_3] > [x_2, x_3, x_4] > [x_1, x_3, x_4]$$

Keep the winner combination (i.e., remove one feature);



Start with all originally available features



Compute the criterion value for all possible combinations after eliminating one feature,

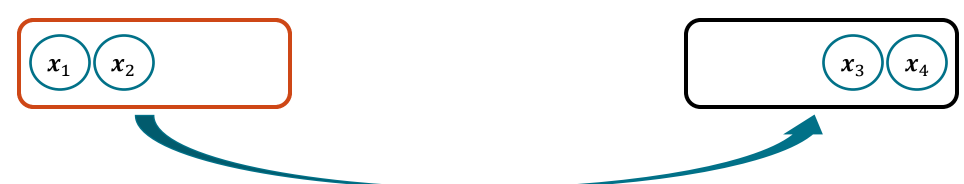
$$[x_1, x_2, x_4] > [x_1, x_2, x_3] > [x_2, x_3, x_4] > [x_1, x_3, x_4]$$

Keep the winner combination (i.e., remove one feature);

Repeat step above: from the winner vector, eliminate one feature, and for each of the resulting combinations, compute the criterion value...



Start with all originally available features



Compute the criterion value for all possible combinations after eliminating one feature,

$$[x_1, x_2, x_4] > [x_1, x_2, x_3] > [x_2, x_3, x_4] > [x_1, x_3, x_4]$$

Keep the winner combination (i.e., remove one feature);

Repeat step above: from the winner vector, eliminate one feature, and for each of the resulting combinations, compute the criterion value...

$$[x_1, x_2] > [x_2, x_4] > [x_1, x_4]$$

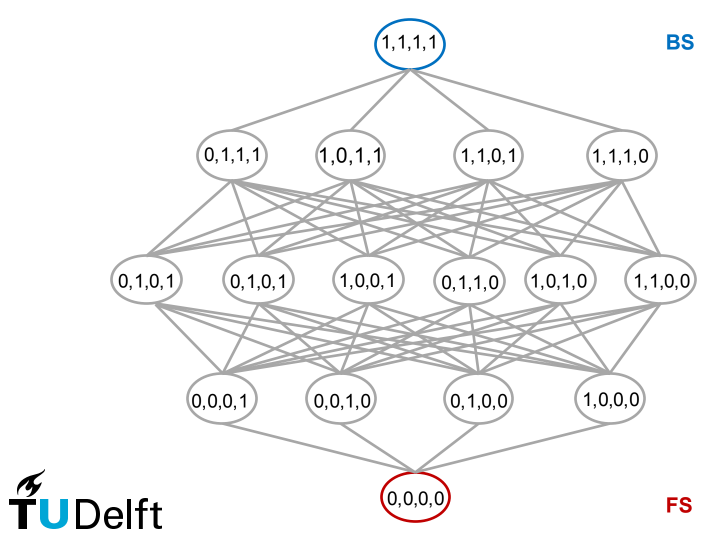


... until a **predefined number** of features are left.

- It applies FS and BS simultaneously:
  - FS starts from the empty feature set.
  - BS starts from the full set of all originally available features.
- To make sure they converge to the same solution:
  - Features already selected by FS are not removed by BS.
  - Features already removed by BS are not selected by FS.



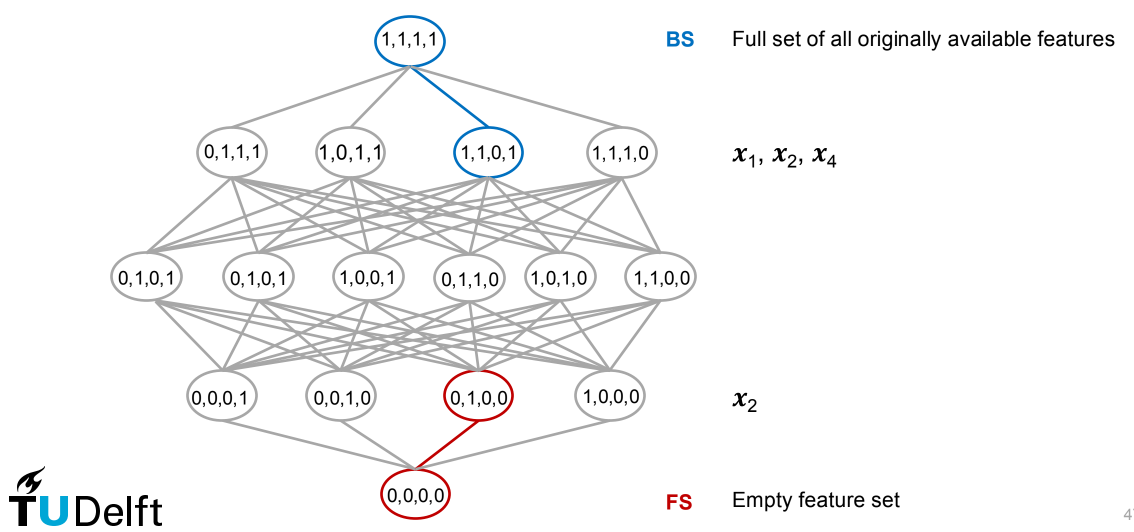
Four features in order of  $x_1$ ,  $x_2$ ,  $x_3$ ,  $x_{4,1}$ , 1 means selected, 0 means not selected, e.g., (0,0,0,1) means only  $x_4$  is selected.



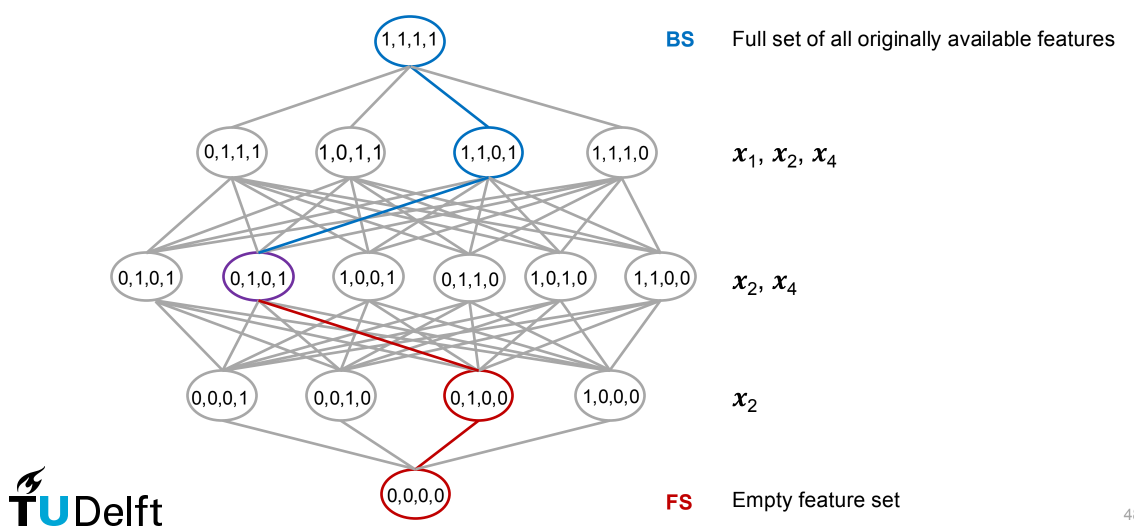
Full set of all originally available features

Empty feature set

Four features in order of  $x_1$ ,  $x_2$ ,  $x_3$ ,  $x_4$ 1 means selected, 0 means not selected, e.g., (0,0,0,1) means only  $x_4$  is selected.



Four features in order of  $x_1$ ,  $x_2$ ,  $x_3$ ,  $x_4$ 1 means selected, 0 means not selected, e.g., (0,0,0,1) means only  $x_4$  is selected.



## Plus-L Take-away-R Selection

- Also based on the ideas of FS and BS. It has two forms.
- If L > R, it starts from the **empty set** and
  - -- repeatedly add *L* features
  - -- repeatedly remove *R* features
- If L < R, it starts from the **full set** of all available features and
  - -- repeatedly remove *R* features
  - -- repeatedly add *L* features
- There is no way of foreseeing the best values of L and R. :-(



## Floating Selection

- FS and BS suffer from the so-called nesting effect. That is,
  - For FS, once a feature is chosen, there is no way for it to be discarded later on.
  - For BS, once a feature is discarded, there is no way for it to be reconsidered again.
- Plus-L Take-away-R Selection doesn't have a flexible backtracking capability.
  - Every round, we have to plus L and have to take away R.
- Floating Selection allows flexible backtracking:
  - The dimensionality of the subset during the search can be "floating" up and down.



## Floating Selection

- There are two floating methods:
  - Floating forward selection & Floating backward selection

- Floating forward selection starts from the empty set,
  - after each forward step, it performs backward steps as long as the criterion function increases.
- Floating backward selection starts from the full set,
  - after each backward step, it performs forward steps as long as the criterion function increases.



## Dimensionality Reduction by Selection or Extraction

Overview – Feature Selection vs Feature Extraction

- Criteria
  - Mahalanobis distance (vs Euclidean distance)
  - Scatter matrices (what are  $S_W$ ,  $S_B$ ,  $S_T$ ?)
- Approaches
  - Sequential feature selection (individual, forward, backward, etc.)
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## Interesting facts about PCA

 PCA is widely recognized as the most classical method for dimensionality reduction, having been invented in 1901.

- However, it doesn't automatically reduce the dimensionality!
- Rather, it transforms the data into a new coordinate system where the choice to retain fewer principal components effectively reduces dimensionality.
  - Retain the variance as much as possible
    - i.e., Minimize the reconstruction error



# PCA: offers different view of your data

Data:

$$m{X} = egin{bmatrix} m{x}_1 \ m{x}_2 \ dots \ m{x}_p \end{bmatrix}$$

 $X = \begin{bmatrix} x_1 \\ x_2 \\ \vdots \\ x_n \end{bmatrix}$  mean-centered data (the mean of each feature is 0); p is number of features

(Variance-) Covariance matrix:

$$\sum_{(p \times p)} \mathbf{X} \mathbf{X}^T = \begin{bmatrix} \sigma_1^2 & \sigma_{12} & \dots & \sigma_{1p} \\ \sigma_{21} & \sigma_2^2 & \dots & \sigma_{2p} \\ \vdots & \vdots & \ddots & \vdots \\ \sigma_{p1} & \sigma_{p2} & \sigma_p^2 \end{bmatrix}$$



## PCA: offers different view of your data

**Eigen-decomposition** of the covariance matrix:

$$m{\Sigma}m{v}=m{v}\lambda,\,\|m{v}\|^2=1$$
  $m{v}_i=egin{bmatrix}v_{1i}\v_{2i}\dots\v_{pi}\end{bmatrix}$  ,  $\lambda_i$  ,  $i=1,2,...p$ 

Transform the data to a new space, in which the coordinate system is defined by the principal components.

$$\mathbf{V} = \begin{bmatrix} \mathbf{v}_1 & \mathbf{v}_2 \dots \mathbf{v}_k \dots \mathbf{v}_p \end{bmatrix}$$

Each column of *V* is a principal component

**ORDERED** by the value of  $\lambda$ ,

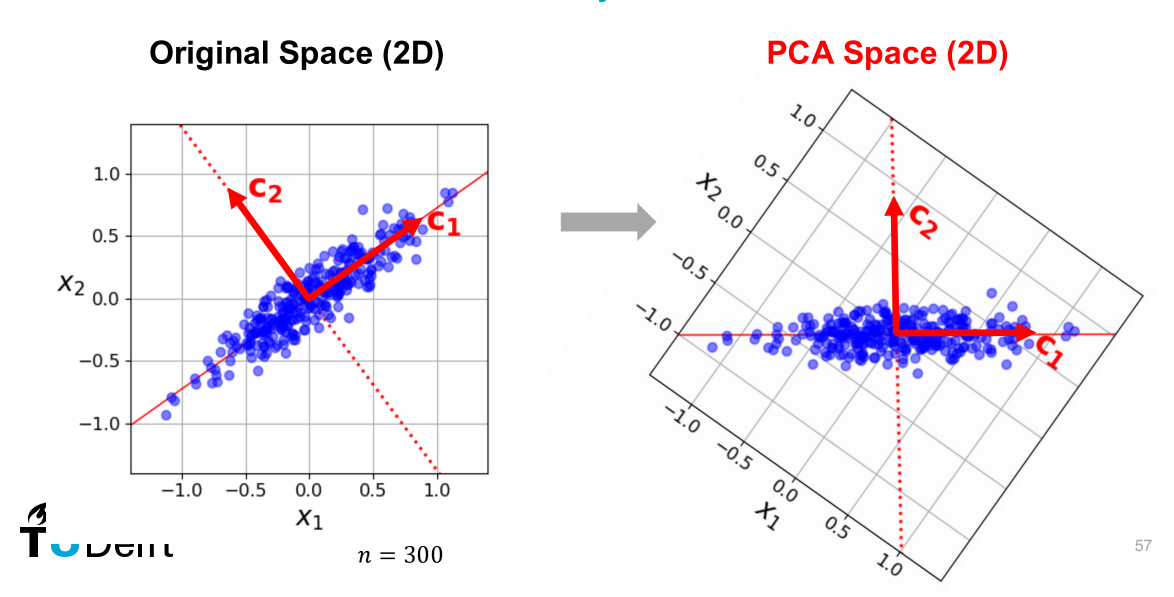
 $\lambda_1$  is the largest eigenvalue



 $egin{align*} oldsymbol{v}_1^T X \ oldsymbol{v}_1^T X \ oldsymbol{v}_2^T X \ oldsymbol{arphi}_1^T X \ oldsymbol{arphi}_2^T X \ oldsymbol{arphi}_1^T X \ oldsymbol{arphi}_1$ 



## PCA: offers different view of your data



## PCA: choose to reduce dimensionality

Again, PCA doesn't automatically reduce the dimensionality.

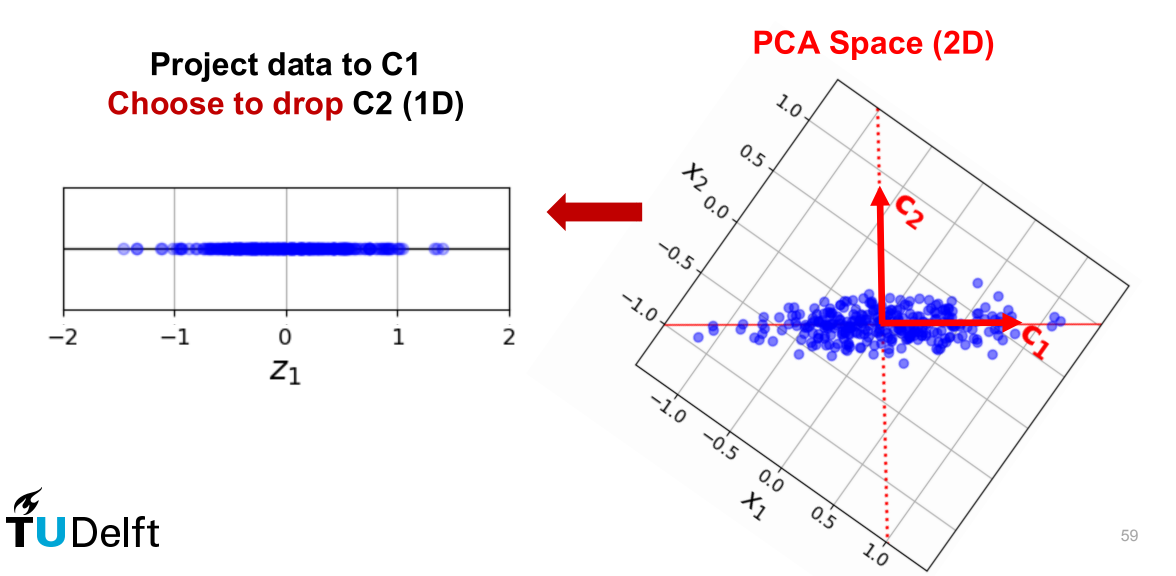
$$T = V^T X$$

• Choose to retain the first k principal components because e.g., 95% variance is captured

What is the dimensionality of 
$$T_k$$
? 
$$T_k = V_k^T X \qquad V_k^T = \begin{bmatrix} \boldsymbol{v}_1^T \\ \boldsymbol{v}_2^T \\ (k \times p) \end{bmatrix} \begin{bmatrix} \boldsymbol{v}_1^T \\ \boldsymbol{v}_2^T \\ \vdots \\ \boldsymbol{v}_k^T \end{bmatrix}$$
 keep drop



# PCA: choose to reduce dimensionality



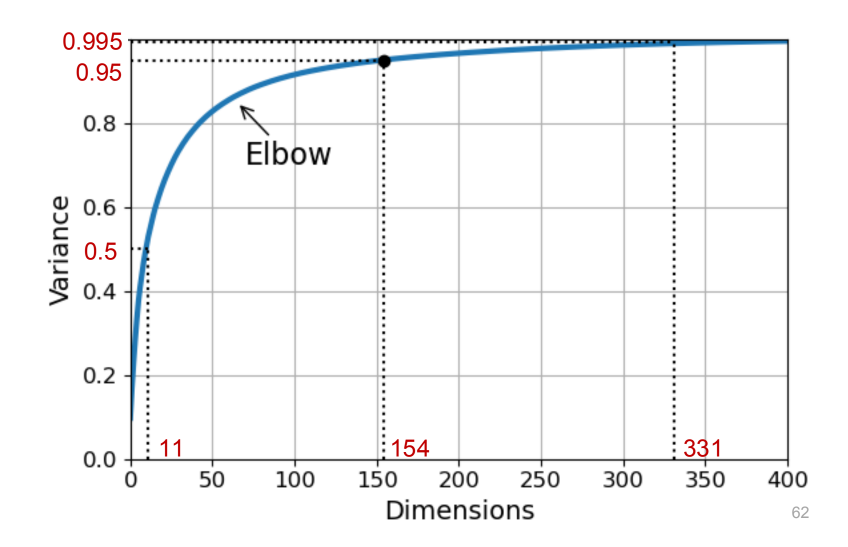
## Quiz

$$egin{aligned} oldsymbol{V} &= egin{bmatrix} oldsymbol{v}_1 & oldsymbol{v}_2 & \dots & oldsymbol{v}_k & \dots & oldsymbol{v}_p \end{bmatrix} & oldsymbol{T}_k &= oldsymbol{V}_k^T oldsymbol{X} & oldsymbol{v}_1^T oldsymbol{X} \ \vdots \ oldsymbol{v}_k^T oldsymbol{X} \end{bmatrix} \ & oldsymbol{v}_k &= oldsymbol{v}_1^T oldsymbol{X} & oldsymbol{v}_1^T oldsymbol{X} \ \vdots \ oldsymbol{v}_k^T oldsymbol{X} \end{bmatrix}$$

- When k = p, T<sub>k</sub> contain exactly the same amount of information as the original data X.
  True or False?
- What does  $v_1^T X$  in  $T_k$  represent?
- What does  $v_1^T \Sigma v_1$  represent?

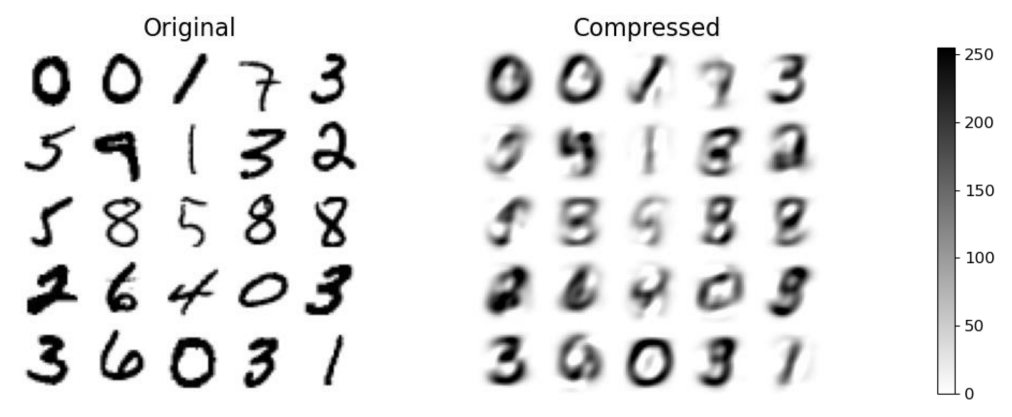


- PCA reconstructions
- Original space (784 D)





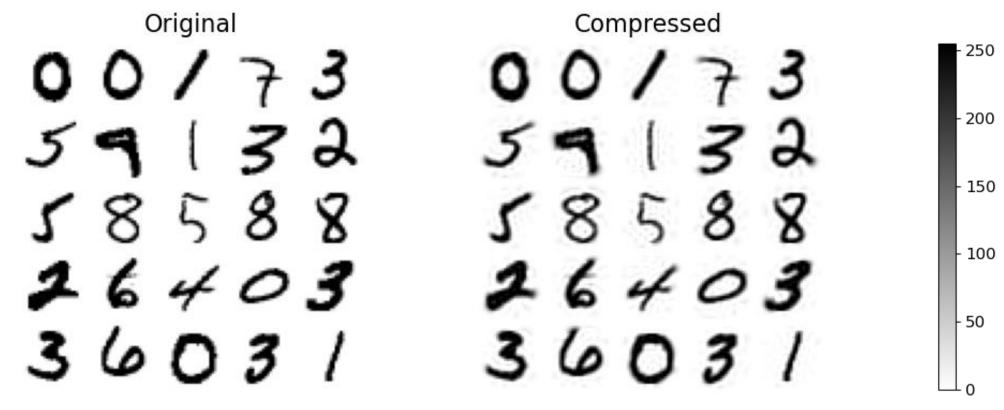
50% Variance: Dim = 11





The more PCs we retain, the smaller the reconstruction error becomes.

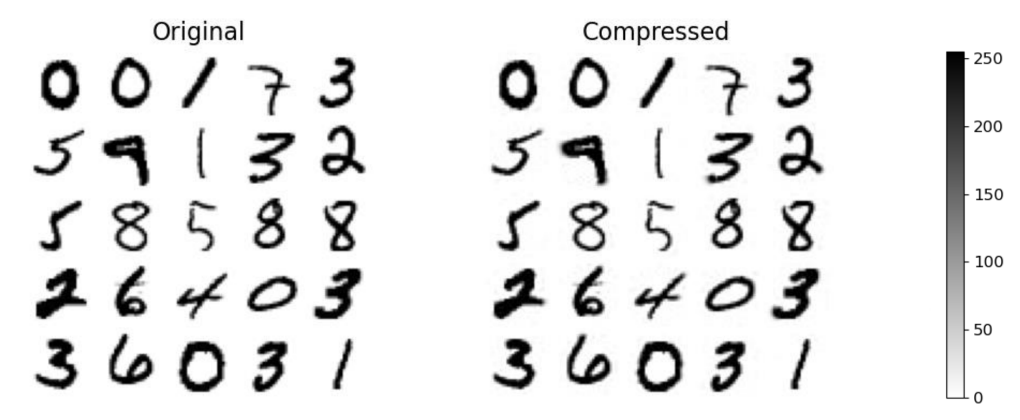
95% Variance: Dim = 154





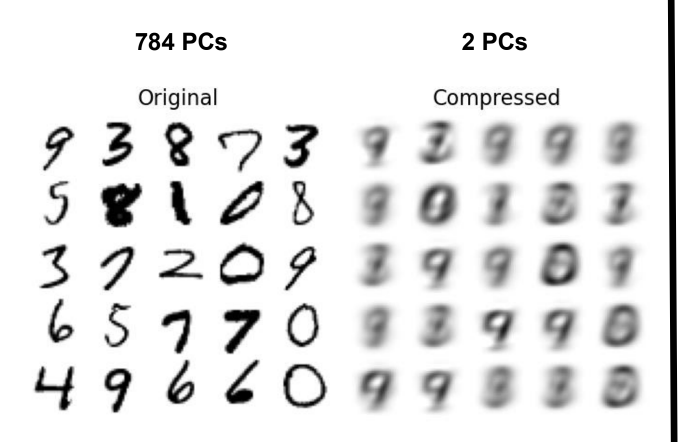
The more PCs we retain, the smaller the reconstruction error becomes.

99.5% Variance: Dim = 331



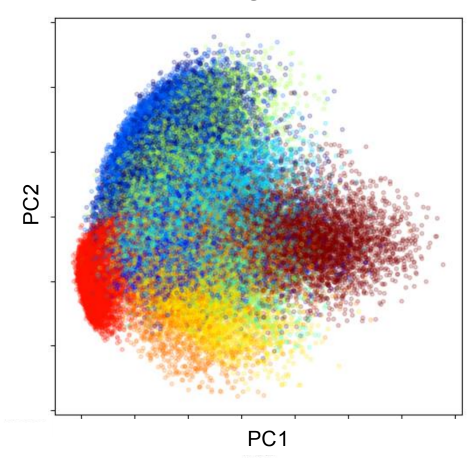


The more PCs we retain, the smaller the reconstruction error becomes.





#### **PCA SPACE**



Colours indicate the class of the object

### Two classical linear feature extractors

LDA (or Fisher mapping):

#### Supervised

-- Find projection vector a that captures the greatest separability between

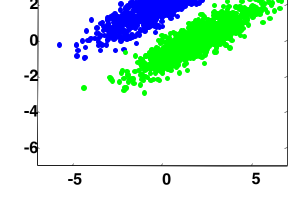
the classes, i.e., choose a to maximize Fisher criterion:

$$J_F(\boldsymbol{a}) = \frac{\boldsymbol{a}^T \boldsymbol{S}_B \boldsymbol{a}}{\boldsymbol{a}^T \boldsymbol{S}_W \boldsymbol{a}}$$

PCA:

#### Unsupervised

-- Capture the greatest variance in the total data



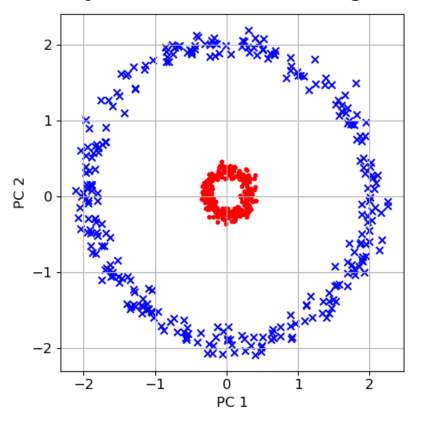


## Kernel PCA - Non-linear Dimensionality Reduction.

- Linear subspaces may be inefficient for some cases.
- The data are not linearly separable in the original dimension.

# Data in 2D space $\mathbf{x}_2$

#### Projection of the data using PCA



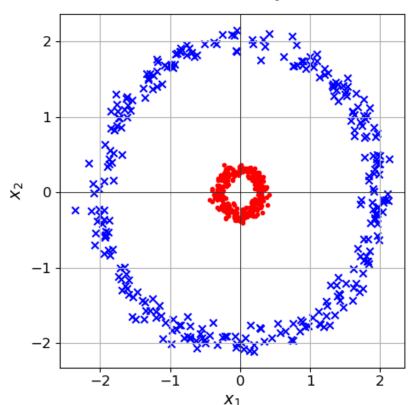


## Kernel PCA – What it does

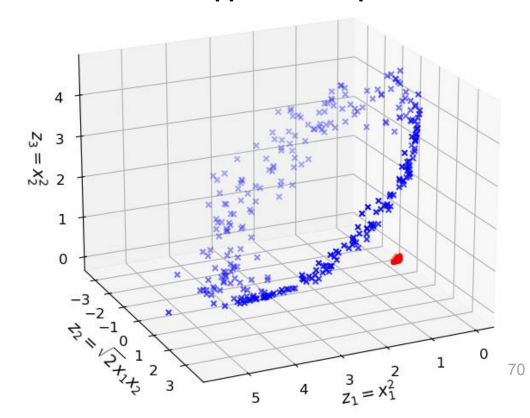
Use a kernel function to project data into a higher-dim. space where they are linearly separable.

$$(x_1, x_2)$$
  $\longrightarrow$   $(z_1, z_2, z_3) = (x_1^2, \sqrt{2}x_1x_2, x_2^2)$ 

#### Data in 2D space

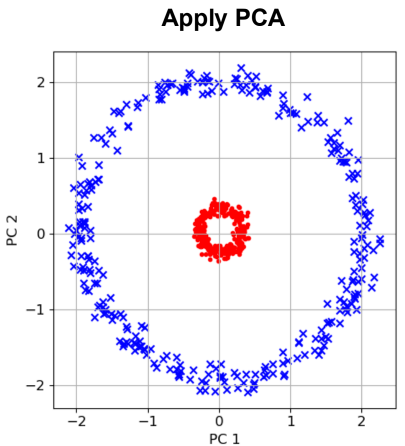


#### Data mapped to 3D space

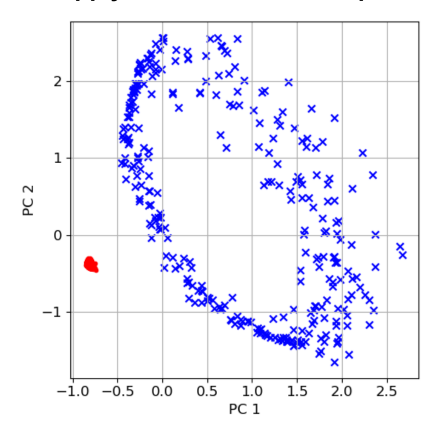




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#### **Apply Kernel PCA and drop PC3**





#### Practice

Given mean-centered data in 3D for which the covariance matrix is given by

$$\Sigma = \begin{pmatrix} 1 & 0 & 0 \\ 0 & 2 & 0 \\ 0 & 0 & 4 \end{pmatrix}.$$

Also given is a data transformation matrix

$$R = \begin{pmatrix} 1 & 0 & 0 \\ 0 & \frac{1}{2} & -\frac{\sqrt{3}}{2} \\ 0 & \frac{\sqrt{3}}{2} & \frac{1}{2} \end{pmatrix},$$

by which we can linearly transform every data vector x (taken as a column vector) to a new 3D column vector z through z = Rx.

We note that *R* is actually a rotation matrix that rotates in the second and third coordinate.

Also note that for its inverse, we have

$$R^{-1} = \begin{pmatrix} 1 & 0 & 0 \\ 0 & \frac{1}{2} & \frac{\sqrt{3}}{2} \\ 0 & -\frac{\sqrt{3}}{2} & \frac{1}{2} \end{pmatrix}.$$

Q1: What is the first principal component of the original data for which we have the covariance matrix  $\Sigma$ ?

Q2: Assume we transform all the data by the transformation matrix R, what does the covariance of the transformed data become?

Q3: What is the first principal component for the transformed data?



#### **Dataset Collection Participants Wanted**

We are collecting a dataset regarding multiple people having conversations in the multiview capture system. **Participants are wanted!** 



What you are going to do: having conversations with other participants (3-6 in total), including having a discussion on a particular movie, playing a game, and having a debate

What we are going to collect: a video and an audio recording of your performance throughout the conversation

The entire process will take approximately one and a half hours. After your participation, we will provide you with a **35€ gift card** as compensation for your time.

Contact: If you are interested, please contact Xiangwei Shi (X.Shi-3@tudelft.nl) or scan the QR code below.

