Lecture 2: Data analysis



Departament de Ciències Matemàtiques i Informàtica 11752 Aprendizaje Automático
11752 Machine Learning
Máster Universitario
en Sistemas Inteligentes

Alberto ORTIZ RODRÍGUEZ

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- Introduction
- Data exploration
- Data preprocessing
- Goodness measures
- Feature selection
- Dimensionality reduction
- Pipelines

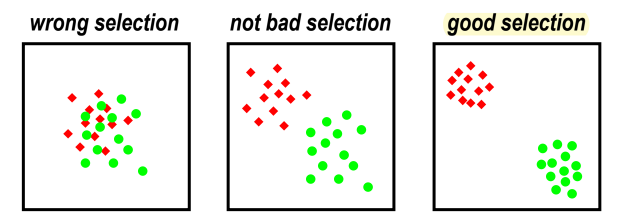
Introduction

Feature engineering

- Given a machine learning problem and a collection of features, select the
 minimum set (and/or maybe a transformation) that improves/retains as much as possible the ability to discriminate among samples
 - Generally we find a large number of possible features
 - the more features, the more complex the classifier is
 more parameters, harder to tune ⇒ feature selection / dimensionality reduction
 - the computational complexity related to their calculation can be taken into account
 - Consider whether a transformation of the features could enhance performance
 - Given the number of samples N, the number of features L should be:
 - big enough to learn
 what makes classes different
 what makes individuals of the same class similar
 - small enough not to make individuals of the same class different
 - it is well known that the **classification error** gets lower as the ratio N/L gets higher
 - as a rule of thumb, N/L > 3, though, in some cases, N/L > 10 or 20
 - in other words, the more features, the more samples you need

Introduction

- Feature engineering
 - General criterion:
 - select those features that result in a large between-class distance and a reduced variance between class elements (within-class variance)



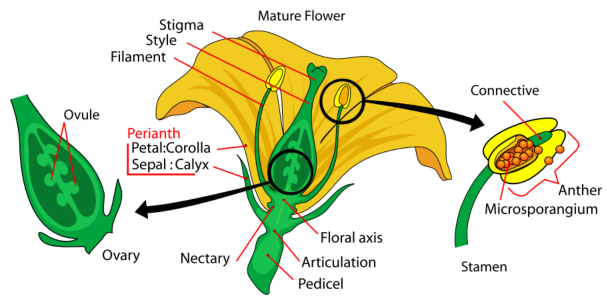
- Actions to do to "engineer" the feature set:
 - understand your data, i.e. explore your data (maybe to know which is your case above)
 - if needed, transform the data to compensate well known problems during training
 - examine features in isolation
 - examine features in combination
 - combine your features

→ feature selection / dimensionality reduction

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- Data exploration is the first, basic step to understand your data
- This includes data visualization for qualitative assessment, detection of anomalies, trends and relationships, as well as to detect the necessity for data cleaning
- Let us consider the **Iris** flower dataset (Fisher's Iris data set)
 - multivariate dataset by the British statistician and biologist Ronald Fisher (1936)
 - 150 samples under four attributes:
 - sepal length
 - sepal width
 - petal length
 - petal width
 - 3 species:
 - setosa
 - versicolor
 - virginica



• Basic descriptive data:

```
import numpy as np
from sklearn.datasets import load iris
iris = load iris()
X = iris.data
y = iris.target
nos, nod = X.shape
print('no. samples = %d, no. dimensions = %d' % (nos, nod))
noc = len(np.unique(y))
print('no. classes = %d' % (noc))
mu = np.mean(X, axis=0)
std = np.std(X, axis=0)
std2 = np.var(X, axis=0)
          mean std var')
print('
for i in range (nod):
    print('x%d: %.2f %.2f %.2f' % (i+1,mu[i],std[i],std2[i]))
```

```
no. samples = 150, no. dimensions = 4
no. classes = 3
    mean std var
x1: 5.84 0.83 0.68
x2: 3.06 0.43 0.19
x3: 3.76 1.76 3.10
x4: 1.20 0.76 0.58
```

• Basic **descriptive data**:

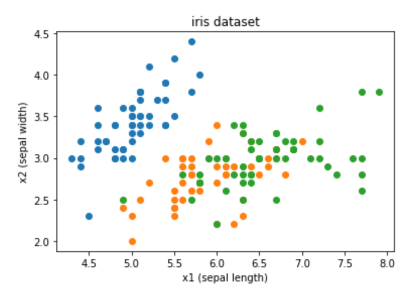
```
import numpy as np
from sklearn.datasets import load_iris
iris = load_iris()
print(iris.DESCR)
```

=========	====	====	======	=====	========	
	Min	Max	Mean	SD	Class Cor	relation
=========	====	====	======	=====	========	
sepal length:	4.3	7.9	5.84	0.83	0.7826	
sepal width:	2.0	4.4	3.05	0.43	-0.4194	
petal length:	1.0	6.9	3.76	1.76	0.9490	(high!)
petal width:	0.1	2.5	1.20	0.76	0.9565	(high!)
==========	====	====	======	=====	========	=======

[:]Missing Attribute Values: None :Class Distribution: 33.3% for each of 3 classes.

• Basic visualization:

```
import matplotlib.pyplot as plt
plt.figure()
for c in range(noc):
    i = np.where(y == c)[0]
    plt.scatter(X[i,0],X[i,1])
plt.xlabel('x1 (sepal length)')
plt.ylabel('x2 (sepal width)')
plt.title('iris dataset')
plt.show()
```



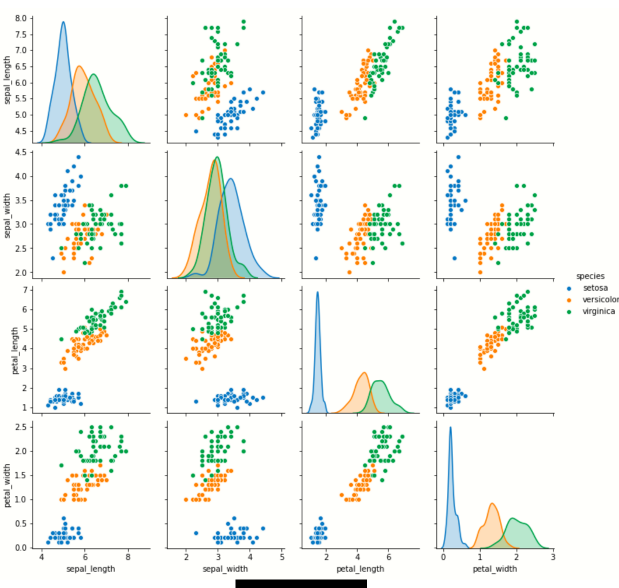
- For multidimensional datasets, i.e. more than 2 / 3 dimensions, the standard methods of visualization are not an option
 - Among many others:
 - the Scatter Plot Matrix (SPLOM) and
 - the parallel coordinates plot

are alternative visualization tools, though of limited capability

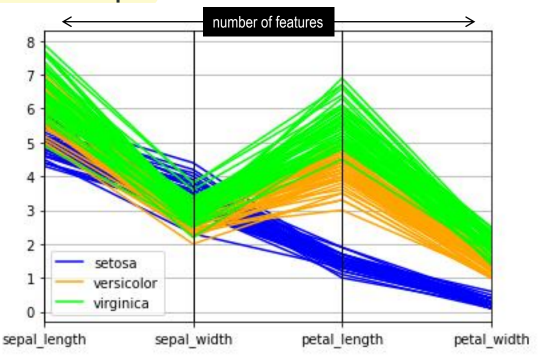
Scatter PLOt Matrix

Correlation plots & Histograms

```
import seaborn as sb
df = sb.load_dataset('iris')
sb.pairplot(df, hue='species')
```



Parallel coordinates plot



```
import matplotlib.pyplot as plt
import pandas as pd
import seaborn as sb
df = sb.load_dataset('iris')
pd.plotting.parallel_coordinates(df, 'species', color=('#0000FF', '#FFA500', '#00FF00'))
plt.legend(loc='lower left')
plt.show()
```

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- Pandas is a library for data manipulation and analysis which can be useful for ML
 - The pandas dataframe class may be particularly useful for data manipulation and indexing, as well as for file input / output
- To create a dataframe we need an array of values.
 Moreover, we can add labels for the columns and for the samples:

```
import pandas as pd
df = pd.DataFrame([[1, 2], [4, 5], [7, 8]],
    index=['cobra', 'viper', 'sidewinder'],
    columns=['max_speed', 'shield'])
print(df.head())

max_speed shield
cobra 1 2
viper 4 5
sidewinder 7 8
```

• In dataframes, indexing can be very flexible with the df.loc() method:

```
print(df.loc[['viper', 'sidewinder']])
print(df.loc['cobra':'viper', 'max_speed'])
print(df.loc[df['shield'] > 4, ['max_speed']])

cobra 1
viper 4
Name: max_speed, dtype: int64

max_speed
viper 4
Sidewinder 7
```

• Let us use the *Titanic* dataset to illustrate other functionalities of *dataframes*:

```
<class 'pandas.core.frame.DataFrame'>
         import seaborn as sb
                                                              RangeIndex: 891 entries, 0 to 890
         titanic = sb.load dataset('titanic')
                                                              Data columns (total 15 columns):
         df = titanic
                                                                                Non-Null Count
                                                                   Column
                                                                                                Dtype
         print(df.info()) -
         print(df.head(3))
                                                               ()
                                                                   survived
                                                                                891 non-null
                                                                                                int.64
                                                                   pclass
                                                                                891 non-null
                                                                                                int64
                                                                                891 non-null
                                                                                                object
                                                                   sex
survived pclass sex age ... deck embark town alive alone
                                                                                714 non-null
                                                                                                float64
                                                                   age
                 male 22.0 ... NaN Southampton
                                                    no False
                                                                                891 non-null
                                                                   sibsp
                                                                                                int.64
              1 female 38.0 ...
                                   С
                                       Cherbourg
                                                   yes False
                                                                   parch
                                                                                891 non-null
                                                                                                int64
              3 female 26.0 ... NaN Southampton
                                                   yes True
                                                                   fare
                                                                                891 non-null
                                                                                                float.64
                                                                   embarked
                                                                                889 non-null
                                                                                                object
                                                                   class
                                                                               891 non-null
                                                                                                category

    df.tail(n) displays the last n samples

                                                                               891 non-null
                                                                                                object
                                                                   who
                                                                   adult male 891 non-null
                                                                                                bool
                                                                   deck
                                                                                203 non-null
                                                               11
                                                                                                category
                                                               12
                                                                   embark town 889 non-null
                                                                                                object

    We can also load the dataset from disk.

                                                               13
                                                                   alive
                                                                                                object
                                                                                891 non-null
          Let us assume the dataset is
                                                               14
                                                                   alone
                                                                                891 non-null
                                                                                                bool
                                                              dtypes: bool(2), category(2), float64(2),
          in file titanic.csv:
                                                              int64(4), object(5)
                                                              memory usage: 80.7+ KB
            df = pd.read csv('titanic.csv')
            print(df.info()) —
```

Other formats also available for input/output, e.g. JSON, excel, etc.

The df.describe() method provides a summary of the dataset statistics:

```
print(df.describe())
                        pclass
         survived
                                                  sibsp
                                                               parch
                                                                            fare
                                        age
       891.000000
                    891.000000
                                714.000000
                                             891.000000
                                                          891.000000
                                                                      891.000000
count
         0.383838
                      2.308642
                                 29.699118
                                               0.523008
                                                            0.381594
                                                                       32.204208
mean
         0.486592
                      0.836071
                                 14.526497
                                               1.102743
                                                           0.806057
                                                                       49.693429
std
                      1.000000
                                 0.420000
                                               0.000000
                                                           0.000000
min
         0.000000
                                                                        0.000000
25%
                      2.000000
                                 20.125000
                                               0.000000
                                                           0.000000
         0.000000
                                                                        7.910400
50%
                      3.000000
                                 28.000000
                                               0.000000
                                                           0.000000
         0.000000
                                                                       14.454200
75%
         1.000000
                      3.000000
                                 38.000000
                                               1.000000
                                                           0.000000
                                                                       31.000000
         1.000000
                      3.000000
                                 80.000000
                                               8.000000
                                                            6.000000
                                                                      512.329200
max
```

• For selecting elements of the dataset, one can additionally use column labels and the **df.iloc()** method:

```
X1 = df.iloc[:, [1,2,3,4,5,6]].to numpy()
                                                       [[3 'male'
                                                                    22.0 1 0 7.25]
X2 =
                                                         [1 'female' 38.0 1 0 71.2833]
df[['pclass','sex','age','sibsp','parch','fare']]
                                                         [3 'female' 26.0 0 0 7.925]]
v = df['survived']
print(X1[0:3,:]) -
                                                         pclass
                                                                  sex age sibsp parch
                                                                                            fare
print(X2.head(3)) —
                                                                  male 22.0
                                                                                          7.2500
                                                              1 female 38.0
                                                                                       0 71.2833
                                                               3 female 26.0
                                                                                          7.9250
                                                                                 0
```

Conditions can also be used for selecting samples:

```
print(df[df['deck'] == 'C'].head(3))
                                      urvived pclass
                                                         sex age ... deck embark town alive alone
                                                    1 female 38.0 ...
                                                                             Cherbourg
                                                                                          yes False
                                                    1 female 35.0 ...
                                                                         C Southampton
                                                                                          ves False
                                                    1 female 58.0 ...
                                                                         C Southampton
                                    11
                                                                                          yes
                                                                                               True
print(df[(df['age'] > 50) & (df['pclass'] < 2)].head(3))
                                     survived pclass
                                                              age ...
                                                                      deck embark town alive alone
                                                         sex
                                                        male 54.0 ...
                                                                         E Southampton
                                                                                               True
                                                                                           no
                                                    1 female 58.0 ...
                                                                         C Southampton
                                    11
                                                                                          yes
                                                                                               True
                                    54
                                                        male 65.0 ...
                                                                             Cherbourg
                                                                                           no False
```

- The dataframe object provides a number of ways to get more details of the dataset:
 - The **df.columns** attribute is a list with the labels of the dataset columns
 - df.values() or df.to_numpy() provide the dataset values as a numpy array
 - df.count_values() returns the number of times the different values occur in a column
 - With df.nunique() we can see the counts of unique values in each column

```
print(df[['age','deck']].nunique())
print(df['sex'].value_counts())

male 577
female 314
Name: sex, dtype: int64
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```

• We can remove some features (columns) which are useless:

```
udf = df
udf.drop('embarked',axis=1,inplace=True)
udf.drop('class',axis=1,inplace=True)
udf.drop('who',axis=1,inplace=True)
udf.drop('adult_male',axis=1,inplace=True)
udf.drop('deck',axis=1,inplace=True)
udf.drop('embark_town',axis=1,inplace=True)
udf.drop('alive',axis=1,inplace=True)
udf.drop('alone',axis=1,inplace=True)
print(udf.info())
```

```
RangeIndex: 891 entries, 0 to 890
Data columns (total 7 columns):
    Column
              Non-Null Count
                             Dtvpe
    survived 891 non-null
                              int.64
    pclass
              891 non-null
                             int.64
              891 non-null
                             object
    sex
                             float64
    age
              714 non-null
    sibsp
              891 non-null
                             int64
    parch
              891 non-null
                             int.64
    fare
              891 non-null
                             float64
```

We can as well drop duplicates, if any:

```
import pandas as pd
df = pd.DataFrame({
    'brand': ['Yum','Yum','Indo','Indo','Indo'],
    'style': ['cup','cup','pack','pack'],
    'rating': [4, 4, 3.5, 15, 5]
})
print(df)
print(df.drop_duplicates())
print(df.drop_duplicates(subset='brand'))
```

```
brand style
                rating
    Yum
                   4.0
0
          cup
    Yum
          cup
                   4.0
  Indo
                   3.5
          cup
   Indo
         pack
                  15.0
   Indo
         pack
                   5.0
  brand style
                rating
    Yum
          cup
                   4.0
                   3.5
  Indo
          cup
  Indo
         pack
                  15.0
   Indo
                   5.0
         pack
  brand style
                rating
                   4.0
0
    Yum
          cup
   Tndo
                   3.5
          cup
```

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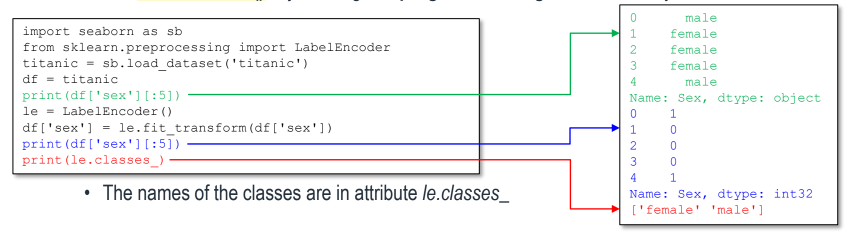
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Data preprocessing

- Preparation of data samples before proceeding to their use
 - Handling categorical data
 - Outlier detection (and removal)
 - Data normalization / standardization
 - Filling in missing data

Data preprocessing: Categorical data

- Handling categorical data
 - Categorical data must be converted to numeric values before learning
 - The LabelEncoder() object assigns a progressive integer label to every class label



 Unfortunately, on some occasions, this is not a good encoding for training, and one-hot encoding must be used instead:

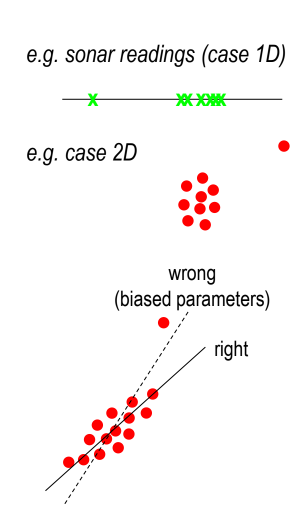
```
from sklearn.preprocessing import OneHotEncoder
df = titanic
ohe = OneHotEncoder()
data = np.expand_dims(df['sex'], axis=-1)
ohe.fit(data)
data_ = ohe.transform(data).toarray()
print(data_[:5])

[[0. 1.]
[1. 0.]
[1. 0.]
[1. 0.]
[0. 1.]]
```

- Outlier detection (and removal)
 - outlier = sample that does not agree with the rest of the population



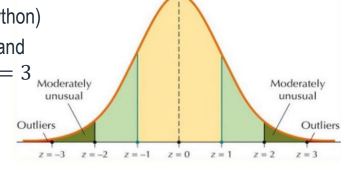
- normally, distance to the mean is $k\sigma$, $k\uparrow\uparrow$
- an outlier can distort training
 - the resulting classifier / regressor may not classify / predict for new samples in the right way



Sources of outliers:

- measurement error (instrument error) or experimental error (wrong data extraction)
- data entry error (data collection/typing) or data processing error
- extreme noisy sample (natural outlier)
- If you need to counteract the outliers, these are some of the **possible actions**:
 - discard the outliers (= full sample) if the dataset permits to do so, i.e. it is big enough
 - alter the data:
 - trimming: extreme values are set to "missing", i.e. NaN (Python)
 - **winsorization**: replace given parts of a sample at the high and low ends with the most extreme values, i.e. use $k\sigma$, e.g. k=3

Winsorized mean. After sorting the data, we replace x_1 and x_{10} by resp. x_2 and x_9 $\underbrace{x_2 + x_2}_{10} + x_3 + x_4 + x_5 + x_6 + x_7 + x_8 + x_9 + x_9}_{10}$ (20% winsorized mean)



Not unusual

1, 5, 7, 8, 9, 10, 10, 12, 12, 34 $\rightarrow \mu$ = 10.8 5, 5, 7, 8, 9, 10, 10, 12, 12, $\frac{12}{12} \rightarrow \mu$ = 9.0

tolerate the outliers by reducing their influence

scipy.stats.mstats.winsorize()

use optimization methods from robust statistics (large values are attenuated "on-line")

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• **z-score method** (Gausssian data): $\mu \pm 3\sigma$ accumulates 99.7% of the probability

```
import numpy as np
from sklearn.datasets import load wine
wine = load wine()
X = wine.data
y = wine.target
print(wine.DESCR)
                                        from scipy import stats
                                        z = np.abs(stats.zscore(X))
outl = np.zeros((3,13))
                                        \# discard samples with z > 3
for c in range(3):
    for f in range (13):
        cc = X[y == c, f]
        mu, sg = np.mean(cc), np.std(cc)
        cut off = sq * 3
        lower, upper = mu - cut off, mu + cut off
        # identify outliers
        outliers = [x for x in cc if x < lower or x > upper]
        nout = len(outliers)
        # non-outliers
        non outliers = [x \text{ for } x \text{ in } cc \text{ if } x \ge lower \text{ and } x \le lower]
        nok = len(non outliers)
        outl[c,f] = nout
print (outl)
```

$$z = \frac{x - \mu}{\sigma}$$

	Min	Max	Mean	SD
			======	
Alcohol:	11.0	14.8	13.0	0.8
Malic Acid:	0.74	5.80	2.34	1.12
Ash:	1.36	3.23	2.36	0.27
Alcalinity of Ash:	10.6	30.0	19.5	3.3
Magnesium:	70.0	162.0	99.7	14.3
Total Phenols:	0.98	3.88	2.29	0.63
Flavanoids:	0.34	5.08	2.03	1.00
Nonflavanoid Phenols:	0.13	0.66	0.36	0.12
Proanthocyanins:	0.41	3.58	1.59	0.57
Colour Intensity:	1.3	13.0	5.1	2.3
Hue:	0.48	1.71	0.96	0.23
OD280/OD315 of diluted wines:	1.27	4.00	2.61	0.71
Proline:	278	1680	746	315
	====	=====		=====

:Missing Attribute Values: None :Class Distribution: class_0 (59), class_1 (71), class_2 (48)

f12 f13 **f7** f10 f11 f1 f3 f4 f6 f8 f9 class 1 0 1 0 0 0 0

With this code, we know that there are outliers in all classes, but we should discover which samples are affected!!

and next do trimming or winsorizing

Inter-quartile range method (non-Gausssian data)

```
import numpy as np
from sklearn.datasets import load wine
wine = load wine()
X = wine.data
y = wine.target
from numpy import percentile
outl = np.zeros((3,13))
for c in range(3):
    for f in range (13):
        cc = X[y == c, f]
        q25, q75 = percentile(cc, 25), percentile(cc, 75)
        iqr = q75 - q25
        cut off = iqr * 1.5
        lower, upper = q25 - cut off, q75 + cut off
        # identify outliers
        outliers = [x for x in cc if x < lower or x > upper]
        nout = len(outliers)
        # non-outliers
        non outliers = [x \text{ for } x \text{ in } cc \text{ if } x \ge lower \text{ and } x \le lower]
        nok = len(non outliers)
        outl[c,f] = nout
print(outl)
```

- IQR = difference between the 75th and the 25th percentiles of the data (Q3, Q1)
- Situates outliers out of the ±k × IQR interval

class	f1	f2	f3	f4	f5	f6	f7	f8	f9	f10	f11	f12	f13
1	0	9	1	3	0	2	0	4	4	1	0	0	0
2	3	7	2	4	5	0	1	0	8	4	1	0	1
3	0	0	0	0	0	2	1	1	2	0	0	2	0

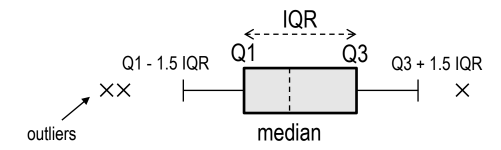
With this code, we know that there are outliers in all classes, but we should discover which samples are affected!!

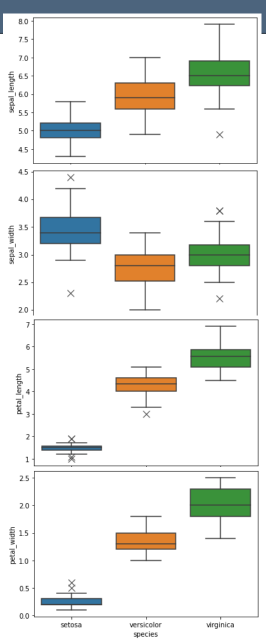
and next do trimming or winsorizing

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Box-plots

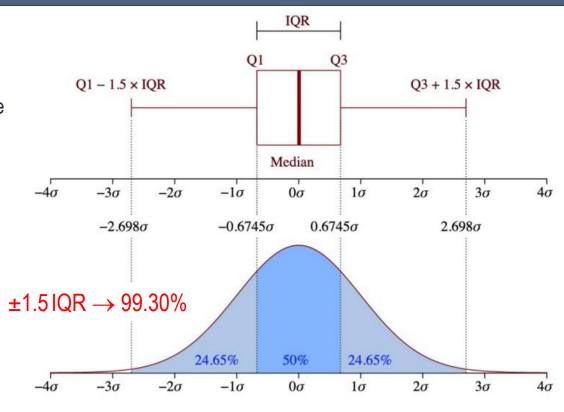
```
import matplotlib.pyplot as plt
import seaborn as sb
                            # also in matplotlib
df = sb.load dataset('iris')
plt.figure()
sb.boxplot(y=df['species'], x=df['sepal length'])
plt.show()
plt.figure()
sb.boxplot(y=df['species'], x=df['sepal width'])
plt.show()
plt.figure()
sb.boxplot(y=df['species'], x=df['petal length'])
plt.show()
plt.figure()
sb.boxplot(y=df['species'], x=df['petal width'])
plt.show()
```

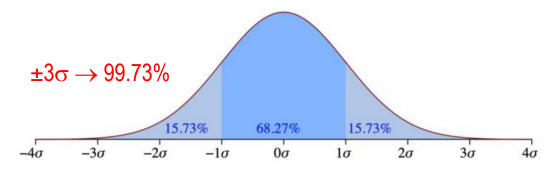




Box-plots

- Why 1.5IQR?
 - Related with the 68–95–99 rule from the Gaussian distribution
 - In the Gaussian distribution, ±1.5IQR covers approx.
 the same probability as ±3σ



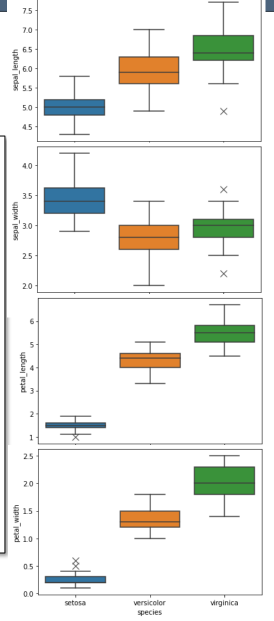


Automatic detection of outliers

 Local Outlier Factor (LOF): measures the local deviation of the density of a sample with respect to its neighbors

```
import numpy as np
import pandas as pd
from sklearn.neighbors import LocalOutlierFactor
import matplotlib.pyplot as plt
import seaborn as sb
df = sb.load dataset('iris')
data = df.values
X = data[:,:-1]
y = data[:,-1]
# identify outliers in the training dataset
lof = LocalOutlierFactor(n neighbors=20)
yhat = lof.fit predict(X)
# select all rows that are not outliers
mask = yhat != -1
X, y = X[mask, :], y[mask]
y = np.expand dims(y,axis=1)
df2 = pd.DataFrame(np.hstack((X,y)))
df2.columns = df.columns
```

Others: IsolationForest, etc.



Data preprocessing

- Data normalization / standardization
 - Often, different features do not have the same dynamic range (range of values)
 - characteristics with wider ranges will have more influence on the classification regardless of whether they are more relevant to the design of the classifier or not



- Solution:
 - normalize/scale features so that their dynamic ranges are similar
 - linear scaling
 - mu-sigma normalization (standardization)
 - max-min normalization
 - others
 - non-linear scaling
 - softmax normalization
 - others

• μ-σ (mu-sigma) normalization

− Given L − feature descriptors:

$$\forall k = 1, \dots, L, \quad \overline{x}_k = \frac{\sum_i x_{ik}}{N}$$

$$\sigma_k^2 = \frac{\sum_i (x_{ik} - \overline{x}_k)^2}{N - 1}$$

$$\widehat{x}_{ik} = \frac{x_{ik} - \overline{x}_k}{\sigma_k}$$

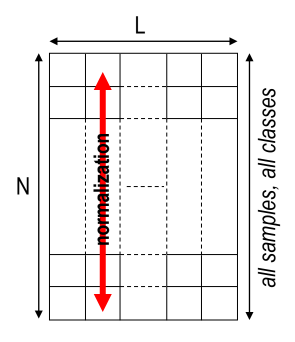
– After the transformation:

$$E[\widehat{x}_{ik}] = 0, \quad Var[\widehat{x}_{ik}] = 1$$

$$x_{ik} - \overline{x}_k = 0 \qquad \Rightarrow \quad \widehat{x}_{ik} = 0$$

$$x_{ik} - \overline{x}_k = +k\sigma \quad \Rightarrow \quad \widehat{x}_{ik} = +k$$

$$x_{ik} - \overline{x}_k = -k\sigma \quad \Rightarrow \quad \widehat{x}_{ik} = -k$$

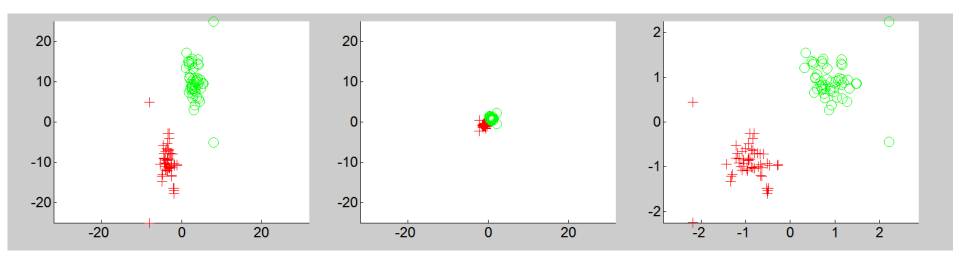


VERY IMPORTANT

Apply the transformation to the full dataset once.

- **Typical mistake**. Standardize the training set separately from the test set.
- ***** Keep the transformation parameters (\bar{x}, σ) to standardize new samples

μ-σ (mu-sigma) normalization



```
normalization: mu-sigma k=1 org: -8.00 - 8.00: 16.00 \leftarrow \text{dynamic range of } x_1 k=2 org: -25.00 - 25.00: 50.00 \leftarrow \text{dynamic range of } x_2 ratio : 3.13 k=1 nor: -2.18 - 2.21: 4.38 \leftarrow \text{dynamic range of } \hat{x}_1 k=2 nor: -2.25 - 2.25: 4.50 \leftarrow \text{dynamic range of } \hat{x}_2 ratio : 1.03
```

Max-min normalization

− Given L − feature descriptors:

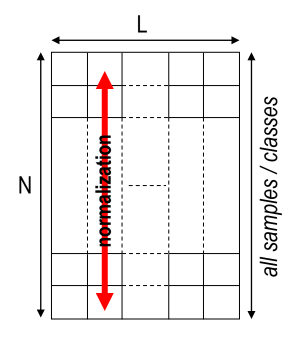
$$\forall k = 1, \dots, L, \quad X_k = \max_i \{x_{ik}\}$$
$$x_k = \min_i \{x_{ik}\}$$
$$\hat{x}_{ik} = \frac{x_{ik} - x_k}{X_k - x_k}$$

– After the transformation: $\widehat{x}_{ik} \in [0, 1]$

$$x_{ik} = \max_{i} \{x_{ik}\} \Rightarrow \widehat{x}_{ik} = 1$$
$$x_{ik} = \min_{i} \{x_{ik}\} \Rightarrow \widehat{x}_{ik} = 0$$

... distributes the data within the range [0,1]

- the original ends correspond to 0 and 1
- e.g. if originally the range of values was [-30, 100], after normalization, value -30 will become 0 for that feature, while a value of 100 will become 1

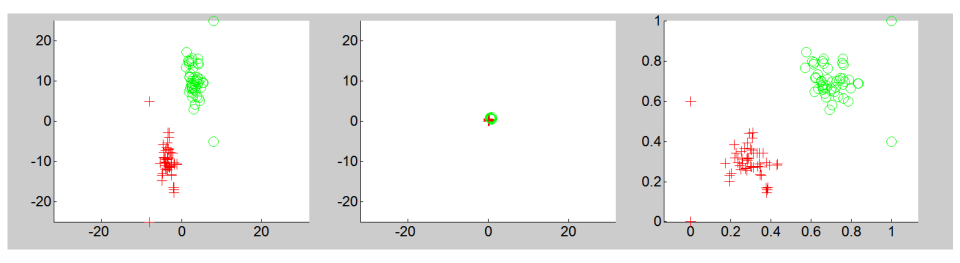


VERY IMPORTANT:

Apply the transformation to the full dataset once.

❖ Keep the transformation parameters (x, X) to normalize new samples

Max-min normalization



```
normalization: min-max  
k=1 \text{ org: } -8.00 - 8.00 : 16.00 \leftarrow \text{dynamic range of } x_1  
k=2 \text{ org: } -25.00 - 25.00 : 50.00 \leftarrow \text{dynamic range of } x_2  
ratio : 3.13  
k=1 \text{ nor: } 0.00 - 1.00 : 1.00 \leftarrow \text{dynamic range of } \hat{x}_1  
k=2 \text{ nor: } 0.00 - 1.00 : 1.00 \leftarrow \text{dynamic range of } \hat{x}_2  
ratio : 1.00
```

- **Softmax normalization** (non-linear transformation)
 - Given L feature descriptors:

$$\forall k = 1, \dots, L, \quad \overline{x}_k = \frac{\sum_i x_{ik}}{N}$$

$$\sigma_k^2 = \frac{\sum_i (x_{ik} - \overline{x}_k)^2}{N - 1}$$

$$z_{ik} = \frac{x_{ik} - \overline{x}_k}{r\sigma_k}$$

$$\widehat{x}_{ik} = \frac{1}{1 + e^{-z_{ik}}}$$

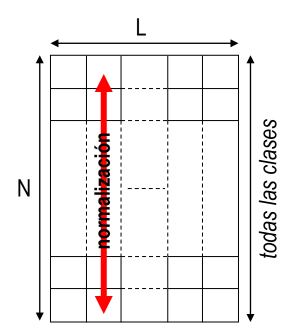
– After the transformation:

$$\widehat{x}_{ik} \in [0, 1]$$

$$\begin{vmatrix} x_{ik} = \overline{x}_k \Rightarrow \widehat{x}_{ik} = \frac{1}{2} \\ x_{ik} = +\infty \Rightarrow \widehat{x}_{ik} = 1 \\ x_{ik} = -\infty \Rightarrow \widehat{x}_{ik} = 0 \end{vmatrix}$$

... but it does not distribute evenly the data within [0,1]

- exponentially "concentrates" values far from the mean as a function of σ and r:
 - the higher r, the closer get the farthest samples



VERY IMPORTANT:

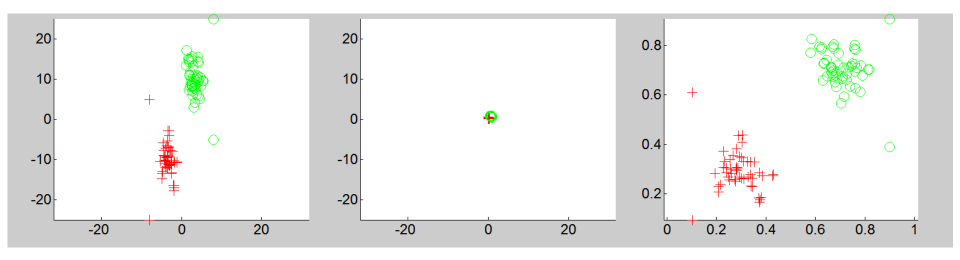
Apply the transformation to the full dataset once.

* Keep the transformation parameters (\bar{x}, σ, r) to standardize new samples

Alberto Ortiz (última revisión 16/10/2023)

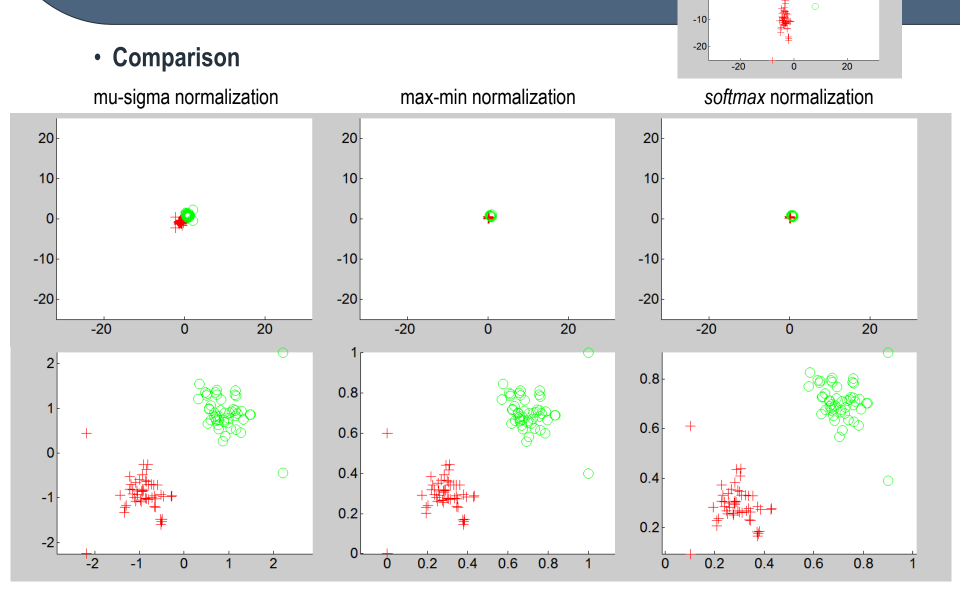
Data preprocessing

Softmax normalization



```
normalization: softmax (r=1)  
k=1 org: -8.00 - 8.00: 16.00 \leftarrow \text{dynamic range of } x_1
k=2 org: -25.00 - 25.00: 50.00 \leftarrow \text{dynamic range of } x_2
ratio : 3.13
k=1 nor: 0.10 - 0.90: 0.80 \leftarrow \text{dynamic range of } \hat{x}_1
k=2 nor: 0.10 - 0.90: 0.81 \leftarrow \text{dynamic range of } \hat{x}_2
ratio : 1.01
```

Data preprocessing



20

10

Alberto Ortiz (última revisión 16/10/2023)

• Support in **Python**:

```
from sklearn import preprocessing
                                               from sklearn import preprocessing
                                               from sklearn.datasets import load iris
from sklearn.datasets import load iris
iris = load iris()
                                               iris = load iris()
X = iris.data
                                               X = iris.data
scaler = preprocessing.StandardScaler()
                                               scaler = preprocessing.MinMaxScaler()
scaler.fit(X)
                                               scaler.fit(X)
Xhat = scaler.transform(X)
                                               Xhat = scaler.transform(X)
                                               print(scaler.min )
print(scaler.mean )
                                               print(scaler.scale )
print(scaler.scale )
                                               print(Xhat.max(axis=0))
print(Xhat.mean(axis=0))
print(Xhat.std(axis=0))
                                               print(Xhat.min(axis=0))
                                                                       \equiv - min / (max - min)
[5.8433 \ 3.0573 \ 3.7580 \ 1.1993] \equiv \mu
                                               [-1.1944 - 0.8333 - 0.1695 - 0.0417] \leftarrow
[0.8253 \ 0.4344 \ 1.7594 \ 0.7597] \equiv \sigma
                                               [ 0.2778  0.4167  0.1695
                                                                            0.41671
[-1.7e-15 -1.8e-15 -1.7e-15 -1.4e-15]
                                               [1. 1. 1. 1.]
                                                                       \equiv 1 / (max - min)
[1. 1. 1. 1.]
                                               [0. 0. 0. 0.]
```

Data preprocessing

Filling in missing data

- Sometimes, a dataset is incomplete:

\mathbf{X}_{1}	X_2	x_3	X_4
1	2	-4	3
3	2	?	2
2	5	3	?
2.5	?	2	3
1	1	0	2
6	2	4	1

- In Python, ? typically appear as Nan,
 Null or None values
- Machine learning models cannot handle these kind of values
- Filling with 0s is not an option
- If the training set is large enough, one can **discard** incomplete samples
- With some datasets, discarding samples is not an option → heuristic prediction
 - e.g. fill missing data using the average value from complete samples
 - e.g. fill missing data according to the inherent distribution

• We will illustrate the process with the *Titanic* dataset:

```
import seaborn as sb
titanic = sb.load_dataset('titanic')
df = titanic.iloc[:,0:13]
print(df.info())
```

- We can see that column Age contains missing (null) values
 - Also other columns, sometimes they are not useful from the ML point of view
- This can also be obtained by means of the isnull() and the isna() methods:

```
print(df.isnull().sum())
print(df.isna().sum())
```

```
RangeIndex: 891 entries, 0 to 890
Data columns (total 12 columns):
    Column
                 Non-Null Count
                                 Dtype
    survived
                891 non-null
                                 int64
    pclass
                891 non-null
                                int64
    sex
                891 non-null
                                object
                714 non-null
                                float64
 3
    age
    sibsp
                891 non-null
                                int64
    parch
                891 non-null
                                int64
                891 non-null
                                float64
    fare
    embarked
                889 non-null
                                object
    class
                891 non-null
                                category
                891 non-null
                                object
    who
    adult male 891 non-null
                                bool
                                category
 11
    deck
                 203 non-null
```

```
survived 0
pclass 0
sex 0
age 177
sibsp 0
parch 0
fare 0
embarked 2
class 0
who 0
adult_male 0
deck 688
```

- We can proceed in several ways:
 - Delete the columns with missing data:

```
udf = df.dropna(axis=1)
print(udf.info())
```

Delete the rows with missing data:

```
udf = df.dropna(axis=0)
print(udf.info())
```

 In this way, we remove too many entries because of the deck column:

```
-> 183 entries, 1 to 889
```

• If we remove first the *deck* column and next the rows with missing data:

```
udf = df.drop('deck', axis=1)
udf.dropna(axis=0,inplace=True)
print(udf.info())
```

```
RangeIndex: 891 entries, 0 to 890
Data columns (total 9 columns):
                  Non-Null Count
    Column
                                  Dtype
                 891 non-null
     survived
                                 int64
                 891 non-null
 1
    pclass
                                 int64
                 891 non-null
                                 object
     sex
                891 non-null
                                 int64
    sibsp
    parch
                891 non-null
                                 int64
    fare
                 891 non-null
                                 float64
    class
                 891 non-null
                                 category
                 891 non-null
                                 object
     who
    adult male 891 non-null
                                 bool
```

```
Int64Index: 712 entries, 0 to 890
Data columns (total 11 columns):
    Column
                  Non-Null Count
                                   Dtype
     survived
                 712 non-null
                                  int64
0
    pclass
                 712 non-null
                                  int64
     sex
                 712 non-null
                                  object
 3
                                  float64
                 712 non-null
     age
     sibsp
                 712 non-null
                                  int64
    parch
                 712 non-null
                                  int64
    fare
                 712 non-null
                                  float64
     embarked
                 712 non-null
                                  object
    class
                 712 non-null
                                  category
                                  object
                 712 non-null
     who
                 712 non-null
     adult male
                                  bool
```

- We can proceed in several ways:
 - Fill the missing values by means of feature imputation:

Numerical data ...

- Fill with the mean
- Fill with the median
- Fill with extreme values that do not occur in the data

Categorical data ...

- Fill with the mode of the distribution
- Fill with a new label

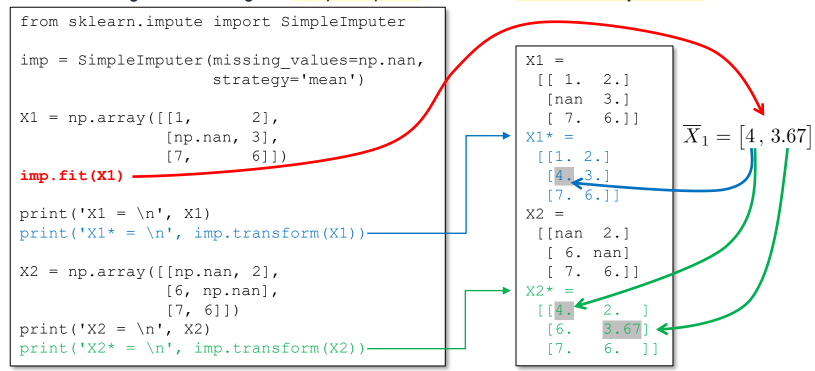
```
udf = df
udf['age'].fillna(udf['age'].mean(),
inplace=True)
```

 It is good practice to register which data values have been *filled in* before the imputation:

```
udf['missing_age'] = df['age'].isnull()
print(udf.info())
```

```
RangeIndex: 891 entries, 0 to 890
Data columns (total 13 columns):
     Column
                  Non-Null Count
                                   Dtype
 0
     survived
                  891 non-null
                                   int64
     pclass
                  891 non-null
                                   int64
                  891 non-null
                                   object
     sex
 3
                                   float64
                  891 non-null
     age
     sibsp
                  891 non-null
                                   int64
     parch
                  891 non-null
                                   int64
     fare
                  891 non-null
                                   float64
     embarked
                  889 non-null
                                   object
 8
     class
                  891 non-null
                                   category
 9
     who
                  891 non-null
                                   object
                 891 non-null
                                   bool
 10
     adult male
11
     deck
                  203 non-null
                                   category
 12
     missing age
                  891 non-null
                                   bool
```

- We can proceed in several ways:
 - Fill the missing values using the <u>SimpleImputer</u> class for univariate imputation



- **Multivariate imputation** is available experimentally in the *IterativeImputer* class
 - Take into account all columns, instead of only the column with missing values
- You can also use a **multivariate regression model** to interpolate missing values

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- Goodness measures
- Feature selection
- Dimensionality reduction
- Pipelines

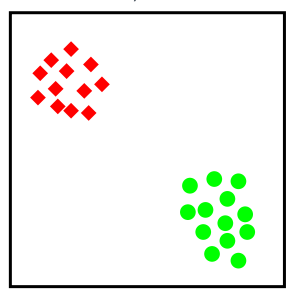
Goodness measures

General criterion:

- Features should result in a large distance between classes (between-class distance) and a reduced variance between class elements (within-class variance)

– Options:

- examine features in isolation
 - not optimal, but it serves to discard bad selections easily
- examine features in combination
- We will consider
 - measures based on scatter matrices (examination in combination)
- but we are not going to consider:
 - measures based on statistical inference tests (isolated examination)
 - Each test allows you to work with one feature and two classes only, and requires assumptions about the distribution of classes



- Measures based on scatter matrices allow multiple classes and several
 characteristics to be treated simultaneously and do not require the assumption of
 normality in the data
- Let us suppose the following dataset:
 - 10 samples/class, 3 classes, 2 features

	ω_1		ω_2		ω_3		_	caso 1			
$\overline{x_{k,1}}$	7.00	9.03	8.02	7.01	8.99	9.00	12				
$x_{k,2}$	7.05	8.90	7.90	7.13	8.97	9.10	11-				
$x_{k,3}$	7.01	8.99	8.12	6.74	9.06	9.29	10-				
$x_{k,4}$	7.07	9.19	7.79	7.04	8.88	9.02	9-	+4	± of		
$x_{k,5}$	7.03	8.99	7.91	6.88	9.08	8.94	8 -			~	
$x_{k,6}$	7.00	8.94	8.26	6.91	9.05	9.17			Ü		
$x_{k,7}$	6.83	9.05	7.88	7.03	9.08	9.09	7-		* *×		
$x_{k,8}$	6.96	9.04	7.93	6.89	8.98	9.03	6				
$x_{k,9}$	7.01	8.95	7.86	6.94	8.95	8.98	5-				
$x_{k,10}$	7.02	9.07	8.05	6.85	8.99	9.10	44	6	8	10	12

We first calculate the within-class scatter matrix (vectors are always column vectors):

$$S_w = \sum_{k=1}^M P_k S_k$$
, where $P_k \approx \frac{n_k}{N}$, $S_k = \frac{1}{n_k - 1} \sum_{j=1}^{n_k} (x_{k,j} - \mu_k) (x_{k,j} - \mu_k)^T$

where:

- P_k is the probability a priori of class ω_k
- S_k is the covariance matrix of class ω_k

$$\mu_k = \frac{1}{n_k} \sum_{j=1}^{n_k} x_{k,j}$$

• Following with the example: M = 3 classes, $n_k = 10$ samples/class, N = 30 samples

We next calculate the between-class scatter matrix (vectors are always column vectors):

$$S_b = \sum_{k=1}^{M} P_k (\mu_k - \mu_0) (\mu_k - \mu_0)^T$$
, where $P_k \approx \frac{n_k}{N}$, $\mu_0 = \sum_{k=1}^{M} P_k \mu_k$

Following with the example:

$$\mu_0 = \frac{1}{3}\mu_1 + \frac{1}{3}\mu_2 + \frac{1}{3}\mu_3 = (7.99, 8.34)^T$$
 $S_b = \begin{pmatrix} 0.6702 & 0.0321 \\ 0.0321 & 0.9817 \end{pmatrix}$

Finally, we obtain the mixture scatter matrix:

$$S_m = S_w + S_b$$
 $S_m = \begin{pmatrix} 0.6799 & 0.0301 \\ 0.0301 & 0.9916 \end{pmatrix}$

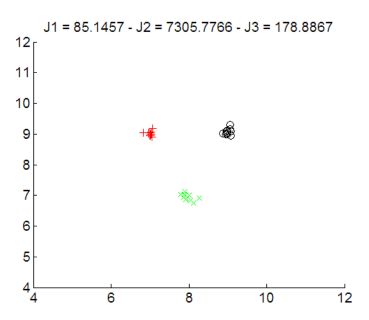
- Important properties:
 - trace(S_w) measures the dispersion of features inside the classes
 - trace(S_b) measures the dispersion of the class centers amongst them
 - ≡ All this permits comparing different features sets among them,
 - i.e. they are not absolute measures, but relative

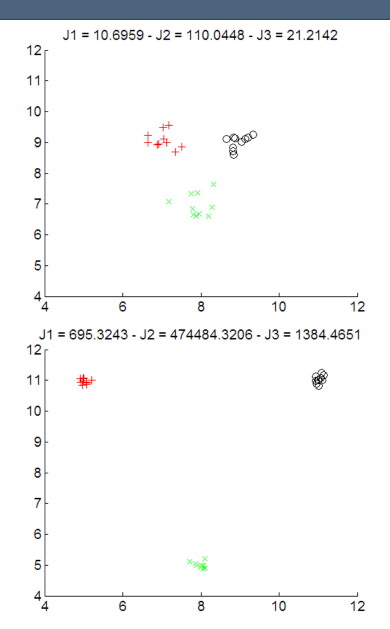
We can define the following measures:

$$J_1 = \frac{\operatorname{trace}(S_m)}{\operatorname{trace}(S_w)}$$

$$J_2 = \frac{|S_m|}{|S_w|} = |S_w^{-1} S_m|$$

$$J_3 = \operatorname{trace}(S_w^{-1} S_m)$$





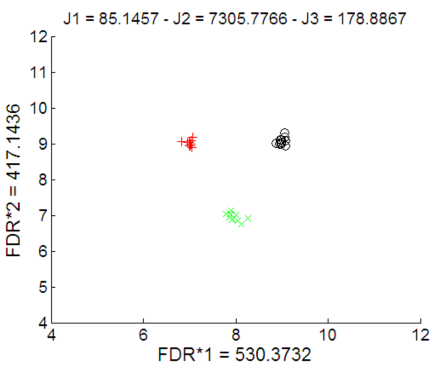
- 1D case (1 feature) and 2 equiprobable classes: (= feature by feature and every 2 classes)
 - $-S_w$ gets reduced to $\sigma_1^2 + \sigma_2^2$
 - $-S_b$ can be shown to be $\frac{1}{2}(\mu_1 \mu_2)^2$
- Following with this reasoning, we obtain the *Fisher's Discriminant Ratio* (FDR) for feature f and classes ω_1 and ω_2 :

$$FDR_f(\omega_1, \omega_2) = \frac{(\mu_1 - \mu_2)^2}{\sigma_1^2 + \sigma_2^2}$$

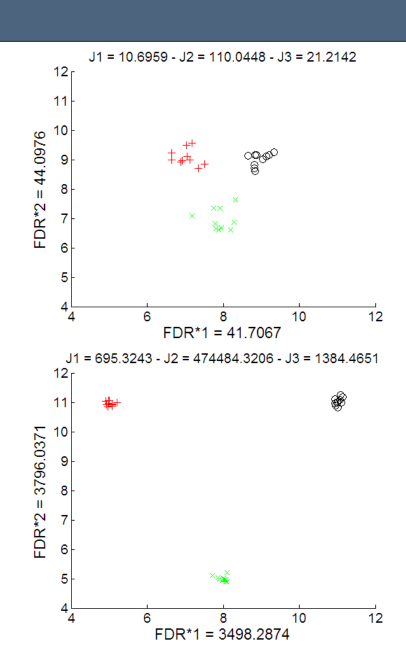
- FDR can be used to quantify the separability capacity of individual features
- Similar to the q-statistics (measures based on statistical hypothesis testing), but the FDR does not depend on the statistical distribution of the data!!
- Multiclass case, one feature f:

$$FDR_f^* = \sum_{k_1 \neq k_2} FDR_f(\omega_{k_1}, \omega_{k_2}), \quad FDR_f^+ = \min_{k_1 \neq k_2} \{FDR_f(\omega_{k_1}, \omega_{k_2})\}$$

• For the previous example:



$\Sigma_{f}FDR_{f}^{*}$	$\Sigma_{f}FDR_{f}^{+}$
947,5168	417,1436
85,8043	41,7067
7294,3245	3498,2874



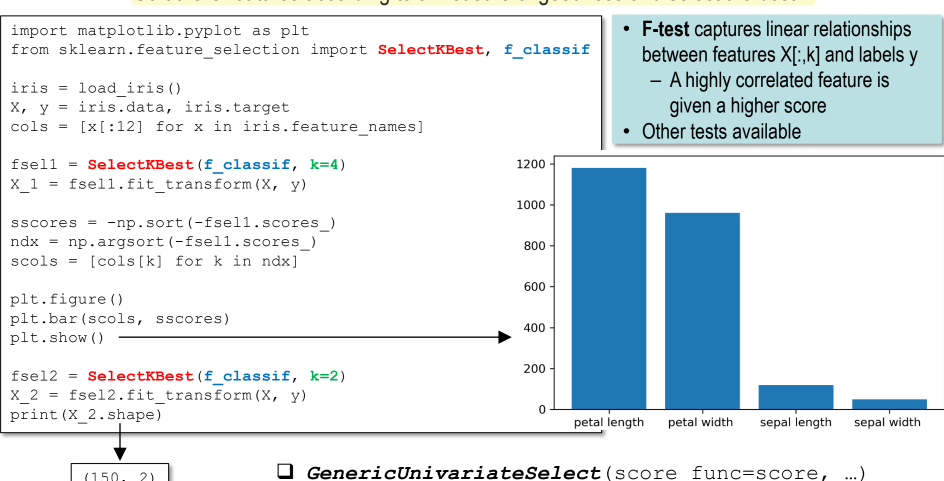
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- Given C features in total, this point is about selecting L as the most suitable subset
- Several approaches:
 - Isolated feature selection
 - Joint features selection

Isolated selection

Sort the **C** features according to a measure of goodness and select the best **L**

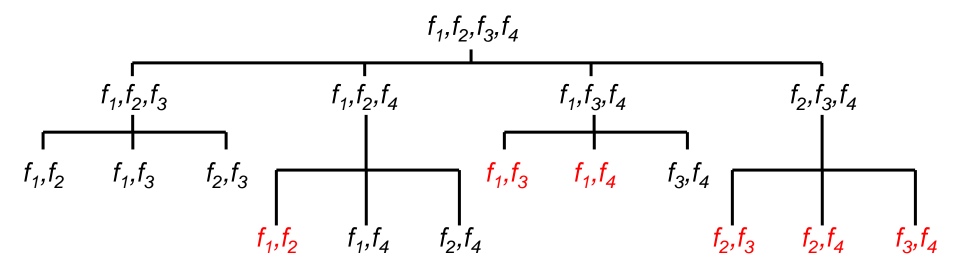


(150, 2)

Joint selection

Consider **subsets of L** features

- 1) Joint exhaustive selection: go through all combinations and select the best one
 - For example, let us suppose C = 4 and L = 2:



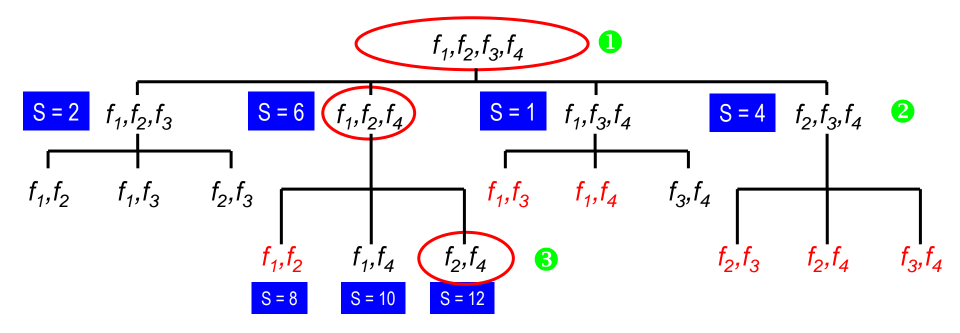
- 6 combinations in total in this case
- General case:

$$\binom{C}{L} = \frac{C!}{L!(C-L)!}$$
 combinations (e.g. $\binom{20}{5} = 15504$)

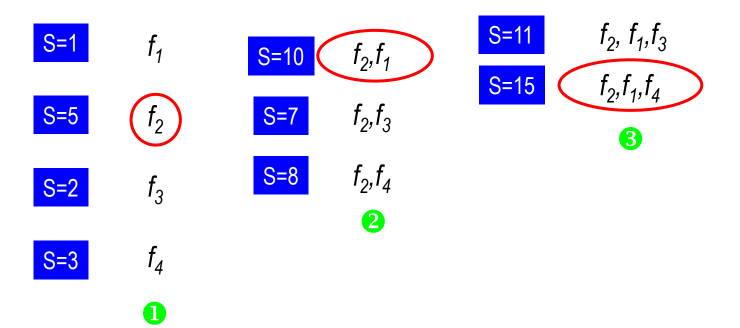
2) Joint suboptimal selection

- Go through a subset of combinations
- It does not guarantee to find the optimal selection but can provide an acceptable selection in less time
- Two variations:
 - Backward sequential selection
 - Forward sequential selection

- Suboptimal solutions: Backward sequential selection
 - Select a suitable score S, e.g. separability indices J_1 , J_2 or J_3 , etc.
 - Let us assume S gets higher as better is the combination
 - Starting from all the features, progressively remove features until reaching the required amount (of features)
 - At each step keep the combination with the largest score



- Suboptimal solutions: Forward sequential selection
 - Select a suitable score S, e.g. separability indices J_1 , J_2 or J_3 , etc.
 - Let us assume S gets higher as better is the combination
 - Starting with one feature, progressively add characteristics until the required number of features is reached
 - At each step keep the combination with the largest score



Example:

```
import numpy as np
from sklearn.datasets import load diabetes
diabetes = load diabetes()
X, y = diabetes.data, diabetes.target
print(diabetes.DESCR)
from sklearn.feature selection import SequentialFeatureSelector
from sklearn.linear model import RidgeCV
ridge = RidgeCV(alphas=np.logspace(-6, 6, num=5)).fit(X, y)
sfs forward = SequentialFeatureSelector(
    ridge, n features to select=2, direction="forward"
).fit(X, y)
sfs backward = SequentialFeatureSelector(
   ridge, n features to select=2, direction="backward"
).fit(X, y)
feature names = np.array(diabetes.feature names)
print(
    "Features selected by forward sequential selection: "
    f"{feature names[sfs forward.get support()]}"
print(
    "Features selected by backward sequential selection: "
    f"{feature names[sfs backward.get support()]}"
```

```
Diabetes dataset
```

Ten baseline variables, age, sex, body mass index, average blood pressure, and six blood serum measurements were obtained for each of n=442 diabetes patients, as well as the response of interest, a quantitative measure of disease progression one year after baseline.

```
:Number of Attributes: First 10 columns are numeric predictive values
```

```
:Target: Column 11 is a quantitative measure of disease progression one year after baseline
```

:Attribute Information:

- age age in years
- sex
- bmi body mass index
- bp average blood pressure
- s1 tc, total serum cholesterol
- s2 ldl, low-density lipoproteins
- s3 hdl, high-density lipoproteins
- s4 tch, total cholesterol / HDL
- s5 ltg, log of serum triglycerides level
- s6 glu, blood sugar level

```
Features selected by forward sequential selection: ['bmi' 's5']
```

Features selected by backward sequential selection: ['bmi' 's5']

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Next week, practical session on Thursday: L2

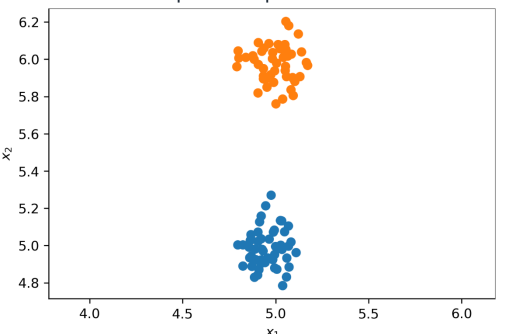
- Laptops available ?
- Description of work to do available on Monday in the webpage
- Revise the notes of lecture 2, try to reproduce the examples

Change in first exam date: L1, L2 - from 19/10 to 2/11/2023 (L3, L4, L6a - 21/12/2023 & L5, L6b - 1/02/2024)

- Dimensionality reduction (DR) refers to the transformation of the original data into a reduced-dimension space, i.e. a new set of variables leading to a reduced set of features
 - Reduce redundancy in the feature set, e.g. avoid correlated features
 - Provide a relevant set of features for a classifier, aiming at improved performance
 - Give rise to new meaningful variables underlying the data, leading to greater understanding of the dataset
- Because of some of the byproducts of DR, it is also termed as feature extraction
- One can find several DR methods in the literature:
 - Principal Component Analysis (PCA) and variants (Sparse PCA, Kernel PCA, etc.)
 - Other matrix factorizations:
 - Non-negative Matrix Factorization (NMF)
 - Independent Component Analysis (ICA)
 - Truncated Singular Value Decomposition
 - Multi-dimensional Scaling (MDS)
 - t-distributed Stochastic Neighbor Embedding (t-SNE) rather for visualizing highdimensional data

- PCA is a popular technique for analyzing large high-dimensional datasets to favour the interpretability of the data while preserving the maximum amount of information, and maybe enabling the visualization of multidimensional data
 - The aim is to derive new variables in decreasing order of importance that are linear combinations of the original variables and that are uncorrelated
 - The data is linearly transformed into a new coordinate system where most of the variation in the data can be described with fewer dimensions than the initial data
 - The principal components are a sequence of orthogonal unit vectors, i.e. an orthonormal basis in which the different individual dimensions of the data are linearly uncorrelated
 - It is a data-directed technique: it makes no assumptions of the existence of any kind of property in the data, e.g. clusters
 - Invented in 1901 by Karl Pearson and developed by Harold Hotellin in the 1930s
 - Also named as the discrete Karhunen-Loeve transform (KLT) in signal processing, proper orthogonal decomposition (POD) in mechanical engineering, etc.

• A simple example:



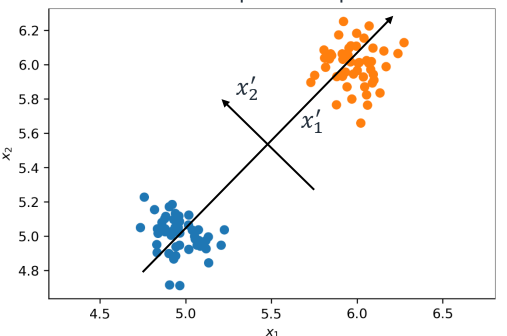
$$\sigma_{x_1}^2 = \frac{1}{N} \sum_{i=1}^N (x_{i1} - \mu_1)^2 = 0.008$$

$$\sigma_{x_2}^2 = \frac{1}{N} \sum_{i=1}^N (x_{i2} - \mu_2)^2 = 0.256$$

$$cov(X) = \frac{1}{N} \sum_{i=1}^{N} (x_i - \mu)(x_i - \mu)^T$$
$$= \begin{pmatrix} 0.008 & 0.012 \\ 0.012 & 0.256 \end{pmatrix}$$

- Which feature should we get rid of?
 - $-x_1$ is not useful from the discrimination point of view
 - $-x_2$ allows discriminating between the two classes
 - $\Rightarrow x_2$ carries more information than x_1 and besides $\sigma_{x_1}^2 < \sigma_{x_2}^2$
 - \Rightarrow if we have to choose, better to get rid of x_1 , the one with lowest variance

A more complex example:



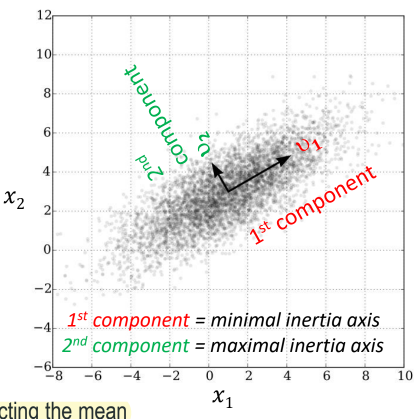
$$\sigma_{x_1}^2 = \frac{1}{N} \sum_{i=1}^{N} (x_{i1} - \mu_1)^2 = 0.280$$

$$\sigma_{x_2}^2 = \frac{1}{N} \sum_{i=1}^{N} (x_{i2} - \mu_2)^2 = 0.257$$

$$cov(X) = \frac{1}{N} \sum_{i=1}^{N} (x_i - \mu)(x_i - \mu)^T$$
$$= \begin{pmatrix} 0.280 & 0.254 \\ 0.254 & 0.257 \end{pmatrix}$$

- Which feature should we get rid of?
 - now is not so simple
 - better to consider a different set of features, x'_1 and x'_2 , to have more or less the same separation between classes (besides, we would see more or less the values as for the previous example in the covariance matrix)

- PCA can be thought of as fitting an
 L-dimensional hyperellipsoid to the data
 - Each axis of the ellipsoid represents a principal component
 - If some axis of the ellipsoid is short, it is because the variance along that axis is small
- x_1 and x_2 features are linearly correlated (when x_1 grows, x_2 grows proportionally)
 - but data points projected onto the resulting orthogonal basis are no longer correlated
- To find the axes of the ellipsoid:
 - center the values of each feature by subtracting the mean
 - compute the scatter or the covariance matrix
 - calculate the eigenvalues and eigenvectors of the resulting matrix
 - the eigenvectors constitute the orthonormal basis (= ellipsoid axes) and
 each eigenvalue is the variance along the respective axis, i.e. eigenvector



Given the data matrix X which has been **mean-centered** $(X = X_u - \overline{X})$, whose rows contain the data samples x_i and its columns are the feature values, we are looking for a set of vectors v_i that constitute an **orthonormal basis** where the data is going to be expressed in:

$$t_{i,k} = \mathbf{x}_i \cdot \nu_k$$
, $i = 1, \dots, N$ and axes $k = 1, \dots, L$

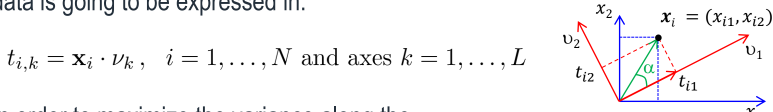
In order to maximize the variance along the first component we look for vector v_1 such that:

$$\nu_1 = \underset{\|\nu\|=1}{\text{arg max}} \left\{ \sum_i t_{i,\nu}^2 \right\} = \underset{\|\nu\|=1}{\text{arg max}} \left\{ \sum_i (\mathbf{x}_i \cdot \nu)^2 \right\}$$

In matrix form, this becomes:

$$\nu_1 = \underset{\|\nu\|=1}{\text{arg max}} \left\{ \|X\nu\|^2 \right\} = \underset{\|\nu\|=1}{\text{arg max}} \left\{ \nu^T X^T X \nu \right\}$$

where $A = X^T X$ is the **scatter matrix** of X_u (\equiv covariance matrix if divided by N-1).



$$\mathbf{x}_i \cdot \nu_1 = \|\nu_1\| \|x_i\| \cos \alpha$$
$$= \|x_i\| \cos \alpha = t_{i,1}$$

• To find the constrained maximization problem we build the **Lagrangian function** L(v) as follows: $\max v^T A v \Rightarrow \max L(v) = v^T A v - \lambda (v^T v - 1)$

• The solution is given by:
$$\frac{\partial L}{\partial \nu} = 2A\nu - 2\lambda\nu = 0 \Rightarrow A\nu = \lambda\nu$$

$$\frac{\partial L}{\partial \lambda} = \nu^T\nu - 1 = 0 \Rightarrow \nu^T\nu = 1$$

• Equation $A\upsilon = \lambda\upsilon$ has L solutions (υ_i, λ_i) for $A_{L\times L}$, i.e. $A\upsilon_i = \lambda_i\upsilon_i$, $\forall i$ which corresponds to the **eigendecomposition** of matrix $A_{L\times L}$, which in matrix form is given by:

$$A = V D V^{-1}, \text{ with } V = \begin{bmatrix} \uparrow & \uparrow \\ \nu_1 & \nu_2 \\ \downarrow & \downarrow \end{bmatrix} \dots \begin{bmatrix} \uparrow \\ \nu_L \\ \downarrow \end{bmatrix} \text{ and } D = \begin{bmatrix} \lambda_1 & 0 & \dots & 0 \\ 0 & \lambda_2 & \dots & 0 \\ & & \ddots & \\ 0 & 0 & \dots & \lambda_L \end{bmatrix}$$

• Linear algebra libraries typically return **unit eigenvectors**, so that all equations are satisfied, and also ordered from largest eigenvalue to lowest eigenvalue:

$$\lambda_1 \geq \lambda_2 \geq \cdots \geq \lambda_L \geq 0$$

- Then, we have to find the eigenvector υ that gives rise to $\max \nu^T A \nu = \max \nu^T V D V^{-1} \nu = \max \nu^T V D V^T \nu$
- Let us now suppose that $\upsilon = \upsilon_{\rm j}$. Then:

$$\nu^T V = \mathbf{e}_j$$
, where $\mathbf{e}_j = (0, \dots, 0, 1, 0, \dots, 0)$

$$\max \ \nu^T V D V^T \nu = \max \ \mathbf{e}_i^T D \mathbf{e}_i = \max \ \lambda_i$$

• The first component of PCA is therefore the eigenvector associated to the **largest** eigenvalue, and so $\upsilon = \upsilon_1$ if the eigenvalues are sorted.

and:

To find the second component, we have to maximize for the remaining variance:

$$\max_{\nu} |\nu^T A \nu - \nu_1^T A \nu_1$$

s.t. $||\nu|| = 1$ and $\nu^T \nu_1 = 0$

Then, the Lagrangian function becomes:

$$L(\nu) = \nu^T A \nu - \nu_1^T A \nu_1 - \lambda \left(\nu^T \nu - 1\right) - \mu \left(\nu^T \nu_1\right)$$
 and
$$\frac{\partial L}{\partial \nu} = 2A \nu - 2\lambda \nu - \mu \nu_1 = 0 \Rightarrow A \nu = \lambda \nu$$

$$\frac{\partial L}{\partial \lambda} = \nu^T \nu - 1 = 0 \Rightarrow \nu^T \nu = 1$$

$$\frac{\partial L}{\partial \mu} = \nu^T \nu_1 = 0$$

Referring to the first equation:

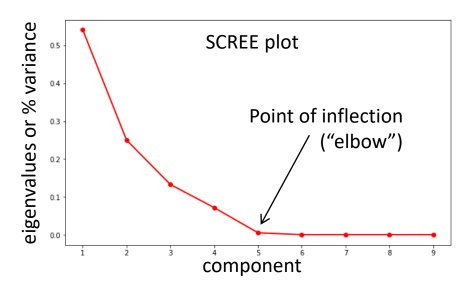
$$\nu_1^T (2A\nu - 2\lambda\nu - \mu\nu_1) = 2\nu_1^T A\nu - 2\lambda\nu_1^T \nu - \mu\nu_1^T \nu_1$$

$$= 2\nu_1^T A\nu - \mu = 0$$

$$\Rightarrow \mu = 2\nu_1^T A\nu = 2\nu^T A\nu_1 = 2\nu^T \lambda_1 \nu_1 = 0$$

• Therefore, the second component is another eigenvector of A and hence has to be the second eigenvector of A, $v = v_2$, since λ_2 is the second largest eigenvalue of A.

- The remaining components can be proved to be the remaining eigenvectors, ordered by the corresponding eigenvalue from higher to lower.
- Now that we know that the components are the eigenvectors of matrix X^TX , we have to deal with the **reduced-dimension representation**.
- To this end, we consider the **fraction of the total variance** that is accounted for by the first p components: $\frac{\sum_{i}^{p} \text{var}\left[\nu_{i}\right]}{\sum_{i}^{L} \text{var}\left[\nu_{i}\right]} = \frac{\sum_{i}^{p} \lambda_{i}}{\sum_{i}^{L} \lambda_{i}}$
 - We can specify a threshold τ on this ratio to choose the number of components necessary to account for at least a τ fraction of the total variance.
- We can also plot the eigenvalues in decreasing order (SCREE plot) and look for the component for which the accounted variance falls sharply:
 - eigenvalues
 - fraction of total variance: $\frac{\lambda_i}{\sum_i \lambda_i}$



• Once we have decided to make use of p components, we can find the reduced-dimensionality vectors/samples: (we do not refer to a particular sample x_i)

$$egin{array}{ccccc} \mathbf{centered\ data} & \mathbf{uncentered\ data} \ & & \chi_p = V_{:p}^T \mathbf{x} & \chi_p = V_{:p}^T (\mathbf{x} - \mu) \end{array}$$

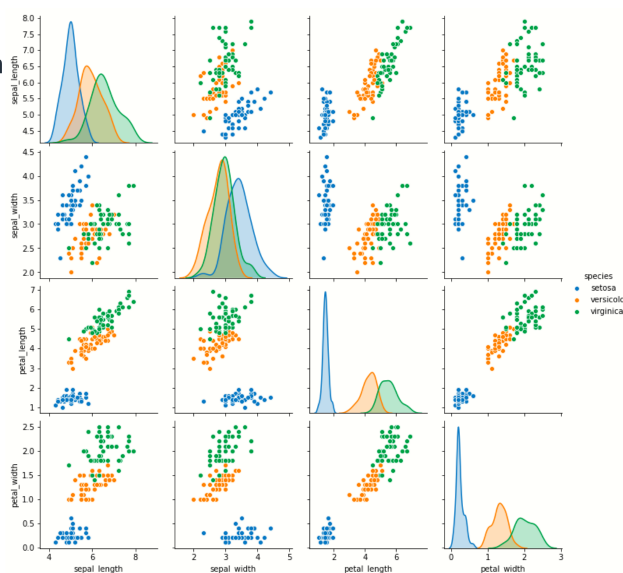
where χ_p is the reduced-dimension sample, μ is the mean and

$$V_{:p} = \left[\nu_1 \mid \nu_2 \mid \dots \mid \nu_p\right]_{L \times p}$$

• The dimensionality reduction operation gives rise to an **error of representation**. This makes interesting to know the representation x_p of χ_p in the original L-dimensional space:

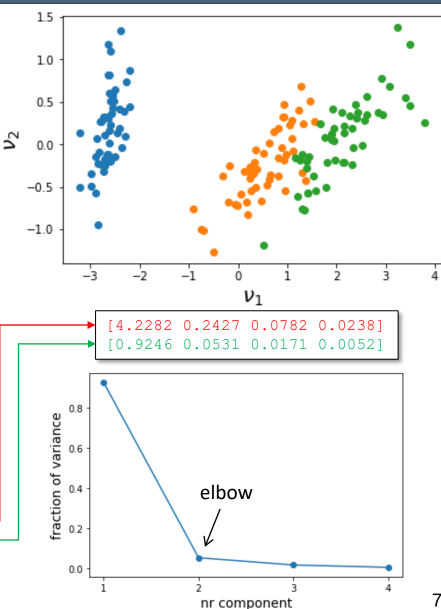
	centered data	uncentered data
to transformed space	$\chi = V^T \mathbf{x}$	$\chi = V^T(\mathbf{x} - \mu)$
to original space	$\mathbf{x} = V\chi$	$\mathbf{x} = V\chi + \mu$
reduced dimension, but in the original space	$\mathbf{x}_p = V \begin{pmatrix} \chi_p \\ 0 \end{pmatrix}$ $= V_{:p} \chi_p$ $= V_{:p} V_{:p}^T \mathbf{x}$	$\mathbf{x}_{p} = V \begin{pmatrix} \chi_{p} \\ 0 \end{pmatrix} + \mu$ $= V_{:p} \chi_{p} + \mu$ $= V_{:p} V_{:p}^{T} (\mathbf{x} - \mu) + \mu$

• Example 1: 7.5 -



Example 1:

```
import numpy as np
import matplotlib.pyplot as plt
from sklearn import datasets
from sklearn.decomposition import PCA
iris = datasets.load iris()
X = iris.data
y = iris.target
pca = PCA(n components=2)
# fit method already includes centering
Xr = pca.fit(X).transform(X)
plt.figure()
for c in range(3):
    i = np.where(y == c)[0]
    plt.scatter(Xr[i,0],Xr[i,1])
plt.show()
pca = PCA(n components=4)
pca.fit(X)
print(pca.explained variance )-
print(pca.explained variance ratio ) -
```



• <u>Example 1</u>:

```
(continued)
pca = PCA(n components=2)
pca.fit(X)
Xr = pca.transform(X)
X = pca.inverse transform(Xr)
import numpy as np
from math import sqrt
error matrix = X - X
error sq = np.sum(np.sum((error matrix)**2, axis=1))
error = sqrt(error sq)
N = X.shape[0]
print('total error = %f, total error/sample = %f' % (error, error / N))
m = np.min(np.abs(error matrix), axis=0)
print('min. errors: %f %f %f %f' % (m[0], m[1], m[2], m[3]))
m = np.max(np.abs(error matrix), axis=0)
print('max. errors: %f %f %f %f' % (m[0], m[1], m[2], m[3]))
m = np.min(X, axis=0)
print('min. values: %f %f %f %f' % (m[0], m[1], m[2], m[3]))
m = np.max(X, axis=0)
print('max. values: %f %f %f %f' % (m[0], m[1], m[2], m[3]))
```

```
total error = 3.899313, total error/sample = 0.025995
min. errors: 0.001556 0.001401 0.000492 0.000810
max. errors: 0.451606 0.463801 0.233806 0.591713
min. values: 4.300000 2.000000 1.000000 0.100000
max. values: 7.900000 4.400000 6.900000 2.500000
```

Example 2: Olivetti faces dataset,
 400 faces of 64 × 64 pixels,
 4096 dimensions

Faces from dataset





Eigenfaces - PCA using randomized SVD





Transformed faces (10 components)





Transformed faces (100 components)





Transformed faces (200 components)





Example 2: Olivetti faces dataset,
 400 faces of 64 × 64 pixels,
 4096 dimensions

Faces from dataset

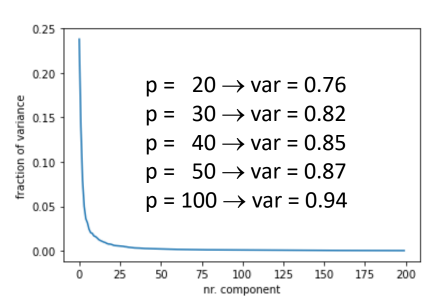




Eigenfaces - PCA using randomized SVD







Transformed faces (200 components)





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- Introduction
- Data exploration
- Data preprocessing
- Goodness measures
- Feature selection
- Dimensionality reduction
- Pipelines

Pipelines

8ρx • Scikit-learn allows chaining steps to transform data until reaching the final estimator:

```
from sklearn import datasets
from sklearn.decomposition import PCA
from sklearn.linear model import LogisticRegression
from sklearn.model selection import GridSearchCV
from sklearn.pipeline import Pipeline
from sklearn.preprocessing import StandardScaler
# Define a pipeline to search for the best combination of PCA truncation
scaler = StandardScaler() # mu-sigma scaler to normalize inputs
pca = PCA()
                          # dimensionality reduction
logistic = LogisticRegression(max iter=10000, tol=0.1) # classifier
pipe = Pipeline(steps=[("scaler", scaler), ("pca", pca), ("logistic", logistic)])
X digits, y digits = datasets.load digits(return X y=True)
# Parameters of pipelines can be set using ' ' separated parameter names:
param grid = {
        "pca n components": [10, 20, 30, 40, 50, 60],
        "logistic C": [0.01, 10, 100],
search = GridSearchCV(pipe, param grid, n jobs=-1)
search.fit(X digits, y digits)
print("Best configuration (CV score=%0.3f):" % search.best score )
print(search.best params )
```

```
Best configuration (CV score=0.917):
{'logistic_C': 10, 'pca_n_components': 50}
```

Lecture 2: Data analysis



Departament de Ciències Matemàtiques i Informàtica 11752 Aprendizaje Automático
11752 Machine Learning
Máster Universitario
en Sistemas Inteligentes

Alberto ORTIZ RODRÍGUEZ