# ICT for Health Laboratory # 3 ROC - Covid-19 serological tests

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#### - Description

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## **Description** [1]

Covid-19 can be diagnosed using naso-pharyngeal swabs or using serological tests (blood test). Serological tests, at least at the beginning, were faster, but naso-pharyngeal swabs were more reliable (higher sensitivity and specificity).

In the experiment, naso-pharyngeal and two serological tests were used on some subjects; for the serological tests the marker levels (IGg) were recorded, for the swab test only the result (positive/negative) was recorded.

Goal of the lab is to set the threshold of the marker above which each serological test is considered positive, **using ROC curves**, and compare the two serological tests, deciding which should be used.

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## **Data analysis**

The input file is covid\_serological\_results.csv and contains 3 columns:

- ► COVID\_swab\_res which is the swab result: 0 means negative, 2 means positive, 1 means uncertain result
- ► IgG\_test1\_titre which is the level of Immunoglobulin G (IgG) in test #1; values range from 2.5 to 314
- ► IgG\_test2\_titre which is the level of Immunoglobulin G (IgG) in test #2; values range from 0 to 9.71

The dataset includes 879 rows, without missing values. Note that the scale with which IgG is measured is somehow arbitrary and therefore each test will have its own threshold.

We will consider swab test result as ground truth (which might not be correct, since also the swab test has its own sensitivity and specificity), and we will **drop patients with uncertain (i.e. equal to 1) swab tests**.

Data analysis

## Read the data [1]

The first lines in the script are the following:

```
import pandas as pd
import numpy as np
import matplotlib.pyplot as plt
import sklearn.cluster as sk

plt.close('all')
xx=pd.read_csv("covid_serological_results.csv")
xx=xx=xx[xx.COVID_swab_res!=1]# remove rows with swab test = 1
xx.COVID_swab_res[xx.COVID_swab_res=2]=1# replace swab test = 2 with swab test = 1
swab=xx.COVID_swab_res.values
Test1=xx.lgG_Test1_titre.values
Test2=xx.lgG_Test2_titre.values
```

## Read the data [2]

 $Perform\ the\ usual\ data\ analysis\ using\ xx. \texttt{describe(),}\ pd. \texttt{plotting.scatter\_matrix(),}\ etc.$ 

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## Find sensitivity and specificity [1]

Let us focus on Test2 (you have to repeat everything for Test1). We arbitrarily set the threshold to 5, and we find sensitivity and specificity for this threshold.

```
10 x=Test2
y=swab
12 x0=x[swab==0] # test results for healthy patients
3 x1=x[swab==1] # test results for ill patients
14 Np=np.sum(swab==1) # number of ill patients
15 Nn=np.sum(swab==0) # number of healthy patients
16 thresh = 5 # example of threshold
17 n1=np.sum(x1>thresh) # number of true positives for the given thresh
18 sens=n1/Np # sensitivity
19 n0=np.sum(x0<thresh) # number of true negatives
20 spec=n0/Nn # specificity
```

Sensitivity is 0.48 (not nice) and specificity is 0.96 (good).

## Find sensitivity and specificity [2]

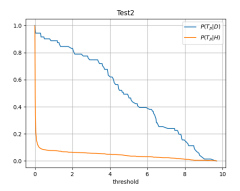
Write a method (def) that generates the values of sensitivity and specificity versus the threshold, considering a reasonable interval for the threshold. A good choice of the set of thresholds is made of the sorted values of Test2; otherwise use np.arange(start,stop,step).

Remember to include threshold zero.

The method should output thresholds, sensitivity and specificity.

Plot sensitivity and specificity versus the threshold (two curves in the same graph).

# Find sensitivity and specificity [3]



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## **ROC** [1]

ROC (Receiver Operating Curve) is sensitivity  $P(T_p|D)$  (y-axis) versus false positive rate  $P(T_p|H)$  (x-axis):

$$P(T_p|D)$$
 versus  $P(T_p|H)$ 

Therefore, to plot ROC, we are not interested in specificity, but in false positive rate, which is 1 minus specificity:

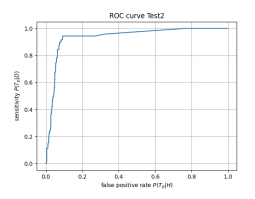
$$P(T_p|H) = 1 - P(T_n|H)$$

21 FA=1-spec

Generate the ROC curve by plotting sensitivity versus false positive rate. Set grid, the x and y labels, etc.

## **ROC [2]**

The ROC curve will be similar to the following:



The ideal working point is for false positive rate equal to 0 and sensitivity equal to 1, which is not reached by this test; however the results are not so bad, after all.

## **ROC** [3]

A parameter that is typically used to understand if a test is better than another is the area under the ROC curve (AUC): the higher the area, the better the test.

Write a method to find the area under a curve (input vectors x and y). Use the trapezoidal rule to evaluate the area.

#### Check

Use methods available in Scikit Learn to check your results:

```
from sklearn import metrics
from sklearn metrics import roc_auc_score, average_precision_score
fpr, tpr, thresh=metrics.roc_curve(swab, Test2, pos_label=1)
AUC_Test2=roc_auc_score(swab, Test2)
```

#### Threshold setting

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## How to set the threshold [1]

There are several ways to set the threshold above which the test is considered positive:

- 1. The required sensitivity is set, and the associated false positive rate (and therefore specificity) is obtained as a consequence.
- **2.** The required false positive rate is set, and the associated sensitivity is obtained as a consequence.
- **3.** Equal values of sensitivity and specificity are set (when sensitivity and specificity are both important at the same level).

Set the threshold according to method 3, but critically think if this choice is correct fro the COVID-19 test.

Threshold setting

## **Final activity**

Repeat the exercise for Test1.

Which of the two tests is better? or are they equivalent?