ICT for Health Laboratory # 1: Regression on Parkinson data

Prof. Monica Visintin - Politecnico di Torino



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Parkinson's disease

Laboratory #1

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To do

Parkinson's disease [1]

Very short description:

- Patients affected by Parkinson's disease cannot exactly control their **muscles**. In particular they show tremor, they walk with difficulties and, in general, they have problems in starting a movement. Many of them cannot **speak correctly**, since they cannot control the vocal chords and the vocal tract. It has been shown that they overcome the illness if they dance or have an **external clock** that gives the time.
- ▶ Levodopa is prescribed to the patients, but most of the medicine, which should be absorbed in the intestine, is absorbed by the stomach; as the movements become slower and slower, levodopa stays more and more in the stomach and cannot reach the intestine.
- ▶ The beneficial effects of levodopa last for some time, and then a new dose of levodopa should be taken. The neurologist decides when the patient should take levodopa and how much levodopa he/she should take, but it is difficult for the neurologist to **optimize the treatment**, because of the continuous progression of the illness.

Parkinson's disease [2]

- ▶ The severity of the illness is measured by neurologists, who judge the patients by asking them to perform many movements (for example tapping the other four fingers with the thumb, or rising from a chair, or walking a short distance, or saying some words) and judging the quality of their life (able to dress? able to prepare his/her own meals?). Adding together the scores gives the final grade, which is called total **UPDRS** (Unified Parkinson's Disease Rating Scale). The visit takes a lot of time, different neurologists may give slightly different scores.
- It would be useful to find an **automatic way** to give the patient an objective score, which can be measured **several times during the day** and help the neurologist to optimize the treatment.
- One possibility is to use parameters of voice to predict the total UPDRS: it is then sufficient to record voice samples (for example using a smartphone), generate these voice parameters (features) and then use a regression technique to predict UPDRS. Unfortunately, Parkinson's disease not always affects voice, and therefore the method can be used only for a subset of patients.
- ► Goal of the lab is to use linear regression to predict total UPDRS from a set of voice parameters and other features.

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To do

Prepare and analyze the data [1]

- ► From DropBox download file parkinsons_updrs_av.csv. This file was obtained by processing an original dataset available at (Irvine University, California) https://archive.ics.uci.edu/ml/datasets/Parkinsons+Telemonitoring
- ► The features available in the dataset are those described at https://archive.ics.uci.edu/ml/datasets/Parkinsons+Telemonitoring
- ▶ The data were obtained by some researchers who evaluated in the same day total UPDRS and motor UPDRS of some patients and recorded the speech of the patients to measure voice parameters. The patients were analyzed several times in 6 months and several records exist for each patient, at different times. The outcome is a matrix with many rows (one for each measurement) and many columns (one for each feature).

Prepare and analyze the data [2]

- A useful Python library that can be used to analyze data is Pandas: if you have not installed it yet, download it.
- ▶ Open Spyder or Pycharm or Jupyter or Colab. Start a new script in which you import pandas, matplotlib.pyplot, numpy:

```
1 import numpy as np
2 import matplotlib.pyplot as plt
3 import pandas as pd
```

Prepare and analyze the data [3]

► Read the Parkinson's data file using Pandas function pd.read_csv(''filename'').

```
4 X=pd.read_csv("parkinsons_updrs_av.csv")
```

This line works if your script and file parkinsons_updrs_av.csv are both in your working folder. If your working folder is different, you must specify the entire path of parkinsons_updrs_av.csv as argument of pd.read_csv.

► Search the web or ask ChatGPT for the usage of Pandas. The main things you have to know is that x is a **DataFrame**, and that many attributes and methods exist for DataFrames (see https://pandas.pydata.org/pandas-docs/stable/api.html#id2).

Prepare and analyze the data [4]

A dataframe is essentially an "organized" table/matrix. Command X.head() shows the first lines of dataframe X:

```
In [4]: X.head()
Out[4]:
    subject# age sex test_time ... HNR RPDE DFA PPE
0 1 72 0 5 ... 27.765 0.406 0.536 0.152
1 1 72 0 12 ... 25.900 0.443 0.539 0.191
2 1 72 0 19 ... 25.843 0.464 0.537 0.209
3 1 72 0 25 ... 28.148 0.422 0.553 0.173
4 1 72 0 33 ... 25.558 0.461 0.544 0.218

[5 rows x 22 columns]
```

Each column has the corresponding column name, all the rows have an index (integer from 0 onwards). You do not have to remember that feature age is the third column, you directly access to X.age or X['age'].

Prepare and analyze the data

Prepare and analyze the data [5]

Examples of methods associated with DataFrames:

```
features=list(X.columns)}#list with the names of the features

X.info()#gives you information about the data (number of valid values, type)

X.describe().T#descr. of dataset (min, max, mean, etc of each feat.)

X.plot.hist(bins=50)#plots the histograms of all the features

X.plot.scatter('a','b')#plots the scatter plot, i.e. x.b versus x.a

X.cov()}#gives the covariance matrix for the features in the columns

X.values()#gives the NumPy Ndarray with the data
```

Prepare and analyze the data [6]

- ▶ Start checking the data (mandatory step each time you work with a new dataset):
 - 1. Write in your code

```
1 X. describe ().T 2 X. info ()
```

and look at the printed values. Check that there are no major problems with the data (no missing values, no out-of-scale values, etc).

Laboratory # 1

Prepare and analyze the data

Prepare and analyze the data [7]

X.info() gives the following output

```
subject#
                         990 non-null
                                           int64
                         990 non-null
                                           int64
        age
        sex
                         990 non-null
                                           int64
                         990 non-null
                                           int64
        test_time
        motor_UPDRS
                         990 non-null
                                           float64
        total_UPDRS
                                           float64
                         990 non-null
        Jitter (%)
                         990 non-null
                                           float64
                                           float64
        Jitter (Abs)
                         990 non-null
        Jitter: RAP
                         990 non-null
                                           float64
    9
        Jitter:PPO5
                         990 non-null
                                           float64
                         990 non-null
    10
        Jitter:DDP
                                           float64
    11
        Shimmer
                         990 non-null
                                           float64
13
    12
        Shimmer (dB)
                         990 non-null
                                           float64
14
        Shimmer: APQ3
                                           float64
    13
                         990 non-null
15
                                           float64
    14
        Shimmer: APQ5
                         990 non-null
16
    15
                         990 non-null
                                           float64
        Shimmer: APQ11
    16
        Shimmer:DDA
                         990 non-null
                                           float64
18
    17
        NHR
                         990 non-null
                                           float64
19
        HNR
    18
                         990 non-null
                                           float64
    19
        RPDE
                         990 non-null
                                           float64
    20
        DFA
                         990 non-null
                                           float64
    21
        PPE
                                           float64
                         990 non-null
   dtypes: float64(18), int64(4)
```

which means that there are 990 rows with no missing values and the read values are float numbers.

Prepare and analyze the data [8]

2. Check the names/meanings of the available features:

```
features=list(x.columns)
print(features)
```

The list of features is 'subject#', 'age', 'sex', 'test_time', 'motor_UPDRS', 'total_UPDRS', 'Jitter(%)', 'Jitter(Abs)', 'Jitter:RAP', 'Jitter:PPQ5', 'Jitter:DDP', 'Shimmer', 'Shimmer(dB)', 'Shimmer:APQ3', 'Shimmer:APQ5', 'Shimmer:APQ11', 'Shimmer:DDA', 'NHR', 'HNR', 'RPDE', 'DFA', 'PPE'.

There are 22 features.

Note again that you can access the values of feature 'age' by writing X.age or X.['age'] (you get a Pandas "series", i.e. one column).

- Feature 'total_UPDRS' is the regressand.
- Using line

```
subj=pd.unique(X['subject#'])# existing values of patient ID
print("The number of distinct patients in the dataset is ",
len(subj))
```

we check that there are 42 patients with distinct IDs.

Features 'age' and 'sex' are obvious, they will be regressors.

Prepare and analyze the data [9]

- ▶ Jitter is the variation of the fundamental frequency (or, conversely, its period) in signals that should be periodic but are not (it is impossible that the frequency of a sinusoidal signal generated by an electronic equipment never changes; it is impossible that a human generated vocal signal like 'a' is perfectly periodic). Shimmer is the variation of amplitude in signals that should be periodic but are not. NHR is the noise to harmonics ratio; HNR is the harmonics to noise ratio. RPDE is Recurrence Period Density Entropy, DFA is the Detrended Fluctuation Analysis, PPE is Perceived Vocal Effort. All these features/parameters are related to voice and are automatically evaluated by specific software that uses a voice signal as input.
- ▶ Other features could be extracted from voice signals, for example the maximum value, the minimum value, the maximum of the absolute value of the FFT (Fast Fourier Transform) of the signal. The process of extracting relevant features is complex. In this lab we use the features available in the dataset.

Prepare and analyze the data [10]

3. Let us check if features are correlated. Method X.cov() gives the covariance of the dataset, but unfortunately feature 'test_time' has a large variance, that makes the other covariance values too small to be seen in an image. Therefore, it is necessary to first normalize the data, then use the DataFrame method cov to evaluate the covariance matrix and plot it using Matplotlib. In practice, instead of showing the covariance

$$C[i,j] = \mathbb{E}\{(X_i - \mu_i)(X_k - \mu_k)\}\$$

of the random variables X_i and X_k , we want to see the correlation coefficient

$$\rho[i,j] = \frac{\mathbb{E}\{(X_i - \mu_i)(X_k - \mu_k)\}}{\sigma_i \sigma_k}$$

 $(\mu_i, \mu_k, \sigma_i, \sigma_k)$ are the means and standard deviations of the random variables X_i and X_k , respectively).

Note that we are just observing the data, we are not yet performing regression and therefore we use the entire dataset.

Remember that the correlation coefficient can only take values in the range [-1, 1].

Prepare and analyze the data

Prepare and analyze the data [11]

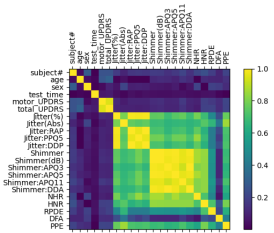
```
Xnorm=(X-X.mean())/X.std()#normalize the entire dataset
c=Xnorm.cov()#measure the covariance
plt.figure()
plt.matshow(np.abs(c.values),fignum=0)
plt.xticks(np.arange(len(features)), features, rotation=90)
plt.yticks(np.arange(len(features)), features, rotation=0)
plt.colorbar()
plt.colorbar()
plt.title('Correlation coefficients of the features')
plt.tight_layout()
plt.savefig('./corr_coeff.png')# save the figure
plt.show()
```

Laboratory # 1

Prepare and analyze the data

Prepare and analyze the data [12]

Correlation coefficients of the features

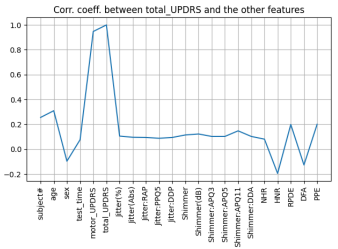


Prepare and analyze the data [13]

- 4. Note that motor and total UPDRS are highly correlated (obviously), but also the various jitter parameters are correlated among themselves, and the same occurs for shimmer parameters. This might give rise to collinearity or multicollinearity: one feature among the regressors can be linearly derived from other regressors, which means that many different vectors we exist that solve the problem with the same value of the objective function. In such cases it might be convenient to remove all but one of the linearly dependent features, selecting the one that has the highest correlation coefficient with the regressand (total UPDRS in our case). However we will first keep all the features to see what happens, and then we will remove some of them.
- 5. Look also at the values in DataFrame c (the matrix with the correlation coefficients) related to total UPDRS:

```
plt.figure()
c.total_UPDRS.plot()
plt.grid()
plt.xticks(np.arange(len(features)), features, rotation=90)
plt.title('corr. coeff. among total UPDRS and the other features')
plt.tight_layout()
plt.show()
```

Prepare and analyze the data [14]



Clearly motor UPDRS is correlated to total UPDRS (it is a part of it and we know), whereas the correlation between total UPDRS and voice features is not so large.

Prepare and analyze the data [15]

- 7. The subject ID is correlated to total UPDRS, but it is not logically correct to use it as regressor: a new patient will have a completely different ID, we cannot allow that his/her regressed total UPDRS depend on this (random) ID. We will later drop the subject ID from the list of regressors.
- **8.** This initial investigation takes time, it is annoying, but it is **mandatory**, otherwise big errors are possible. You must be aware of the meaning of the data you are processing. It is not a matter of blindly running an algorithm, people health is at stake.

Prepare and analyze the data [16]

Prepare a new DataFrame in which you **randomly permute** (**shuffle**) the rows of the original DataFrame. This operation avoids that the data of only the first patients appear in the training dataset. The rationale behind this operation is that we **pretend** that all the measurements are related to different patients at different stages in the illness evolution. We then take the first rows of the DataFrame to train/validate the regression model and the remaining rows to test the performance of the found model. It would be more correct to select some patients as training patients (and shuffle only the data of these patients, if necessary), and then use the remaining patients as test patients. **What is the relationship between shuffling and overfitting, in your opinion?** Shuffling is performed setting the seed, to have reproducibility of the script. Note that results might change by changing the seed.

Prepare and analyze the data

Prepare and analyze the data [17]

First solution to shuffle the data:

Second solution to shuffle the data:

```
43  Xsh=X. sample (fract=1, replace=False, random_state=101, axis=0,
44  ignore_index=True)
```

► The regressand will be **total UPDRS**, whereas all the other features (including motor UPDRS and excluding subject ID) will be regressors.

Perform regression [1]

▶ It is now time to start generating the regression model. We assume that random variable total_UPDRS linearly depends on the random variables sex, age, motor_UPDRS, shimmer, jitter, etc:

$$Y = w_1 X_1 + w_2 X_2 + \dots + w_{N_f} X_{N_f} + \nu$$

where $Y, X_1, \dots, X_{N_f}, \nu$ are all random variables and w_1, \dots, w_{N_f} are the weights to be found.

Perform regression [2]

▶ In the previous model we assume that all the random variables have zero mean. A more complete model is

$$Y = w_1 X_1 + w_2 X_2 + \dots + w_{N_f} X_{N_f} + C + \nu$$

but it is convenient to work with features with zero-mean and variance one, which typically reduces numerical problems and speeds up algorithm convergence (once you remove the mean from all the random variables, including the regressand, the model has obviously intercept equal to zero). Therefore we first perform **normalization (or standardization or scaling)** of the data, by removing the mean and divide by the standard deviation each random variable:

$$X_{norm} = \frac{X - \mu_X}{\sigma_X}; \ (X = \sigma_X X_{norm} + \mu_X)$$

In order to be consistent with our scenario, **the feature means and standard deviations can only be measured on the training dataset**: when we train the regression model we do not know the parameters of **future patients** that will appear in the test dataset, do we?

Perform regression [3]

▶ We will use the first 50% of the rows (495) of the shuffled matrix as **training** points (define the new DataFrame as X_tr), and the remaining 50% of the rows (495) for **testing** (define the new DataFrame as X_te). This is accomplished with the lines

```
At Ntr=int(Np*0.5) # number of training points

Nte=Np-Ntr # number of test points

X_tr=Xsh[0:Ntr]# dataframe that contains only the training data

mm=X_tr.mean()# mean (series) of the training data

ss=X_tr.std()# standard deviation (series) of the training data

my=mm['total_UPDRS']# mean of total UPDRS

sy=ss['total_UPDRS']# st.dev of total UPDRS
```

A series is substantially a DataFrame with just one column (not exact, just to give an idea).

Perform regression [4]

Now we can normalize the data and split them into training, validation, test subsets. At the moment only training and test subsets are needed (LLS solution). Remember that validation is a subset of the training dataset.

```
| Xsh_norm=(Xsh-mm)/ss# normalized data
| ysh_norm=Xsh_norm['total_UPDRS']# regressand
| Xsh_norm=Xsh_norm.drop(['total_UPDRS','subject#'],axis=1)# regressors
| Xtr_norm=Xsh_norm[0:Ntr]# training regressors
| X_te_norm=Xsh_norm[Ntr:]# test regressors
| y_tr_norm=ysh_norm[0:Ntr]# training regressand
| y_te_norm=ysh_norm[Ntr:]# test regressand
```

Perform regression [5]

- ► Regression procedure:
 - DataFrames X_tr_norm and Y_tr_norm can be directly used to find (no need to move to Numpy NDarrays):

$$\hat{\mathbf{w}} = \arg\min \|\mathbf{X}\mathbf{w} - \mathbf{y}\|^2 / N$$

 $(X_{tr}_norm=X, Y_{tr}_norm=y, Ntr=N)$. With LLS:

60 w_hat=np.linalg.inv(X_tr_norm.T@X_tr_norm)@(X_tr_norm.T@y_tr_norm)

Perform regression [6]

However slicing (selection of subsets of rows/columns) with Pandas is sometimes more complex than in Numpy (for which slicing is obvious), so we suggest to use Numpy, as follows:

```
Xsh_norm=(Xsh-mm)/ss# normalized data
ysh_norm=Xsh_norm['total_UPDRS']# regressand
Xsh_norm=Xsh_norm.drop(['total_UPDRS', 'subject#'], axis=1)# regressors
Xsh_norm=Xsh_norm.values # Xsh_norm is now a Numpy NDarray (matrix)
ysh_norm=ysh_norm.values # ysh_norm is now a Numpy NDarray (vector)
X_tr_norm=Xsh_norm[0:Ntr]# training regressors
X_te_norm=Xsh_norm[Ntr:]# test regressors
y_tr_norm=ysh_norm[0:Ntr]# training regressand
y_te_norm=ysh_norm[Ntr:]# test regressand
w_hat=np.linalg.inv(X_tr_norm.T@X_tr_norm)@(X_tr_norm.T@y_tr_norm)
```

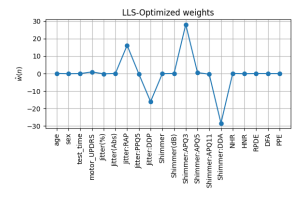
Of course you can use the classes you wrote during the first laboratory to perform the regression.

Perform regression

Perform regression [7]

2. Once you find $\hat{\mathbf{w}}$, you must plot it in order to find potential problems (again, you can use the method we wrote in laboratory 0):

Perform regression [8]



The effect of **collinearity** can be seen since, for example, the weights associated with 'Shimmer:APQ3' and 'Shimmer:DDA' are very large and with opposite signs (the same occurs with 'Jitter:RAP' and 'Jitter:DDP'). For the moment we keep all the regressors and we finish the analysis, but you must drop the two features Jitter:DDP and Shimmer:DDA in your final version of the script.

Perform regression [9]

3. Then, having found $\hat{\mathbf{w}}$, the test dataset will be used to evaluate

$$\hat{\mathbf{y}} = \mathbf{X}\hat{\mathbf{w}}$$

 $(X_{te}_{morm}=X)$. Goodness of $\hat{\mathbf{w}}$ will be measured by comparing $\hat{\mathbf{y}}$ and Y_{te}_{morm} . However, we are interested also in the training dataset performance to find out a possible overfitting phenomenon:

If the mean square error for the training dataset is much smaller than that for the test dataset, then overfitting has occurred.

Perform regression

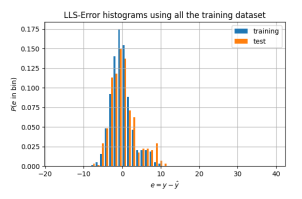
Perform regression [10]

4. Note that y_te_norm and y_hat_te_norm are **normalized**, and therefore the value of MSE (mean Square Error) does not say much to a medical doctor (what is the unit of measurement of the normalized mean square error?), and it is necessary to first de-normalize \hat{y} to get a meaningful mean square error:

Performance figures [1]

1. You must check if the regression error shows peculiar trends, which might reveal an error in the script. A histogram of the error $Y - \hat{Y}$ is useful.

Performance figures [2]



Clearly, the error does not have a Gaussian pdf, it looks like a mixture of two Gaussian pdfs, and there is not much difference in the training and test subsets (which means that there is no **overfitting**). What do you expect this figure to be like, if overfitting were present?

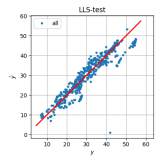
Perform regression

Performance figures [3]

2. You must compare the true and the regressed value of y by plotting \hat{y} versus y (regression line). Note: when you plot a versus b, a is on the y-axis and b is on the x-axis.

```
95  y_hat_te=(X_te_norm@w_hat)*sy+my
96  y_te=y_te_norm*sy+my
97  plt.figure(figsize=(6,4))
98  plt.plot(y_te,y_hat_te,'.')
99  v=plt.axis()
100  plt.plot([v[0],v[1]],[v[0],v[1]],'r',linewidth=2)
101  plt.xlabel(r'$y$')
102  plt.ylabel(r'$\y\$')
103  plt.grid()
104  plt.title('LLS-test')
105  plt.savefig('./LLS-yhat_vs_y.png')
106  plt.savefig('./LLS-yhat_vs_y.png')
107  plt.show()
```

Performance figures [4]



The estimated values \hat{y} are close to the true values y apart from some cases in which y is very large and \hat{y} takes smaller values with an error around 7-8 UPDRS points (which justifies the second Gaussian-like part of the histogram for error values around 8). Also there is a point with y=42 and \hat{y} close to zero.

Performance figures [5]

3. Other important parameters are min, max, mean, standard deviation, mean square value of the error in each of the subsets, R^2 (coefficient of determination), correlation coefficient:

```
108 \mid E_{tr}_{min} = E_{tr}_{min}()
109 \mid E_{tr_max} = E_{tr_max}()
110 \mid E_{tr_mu} = E_{tr.mean}
111 \mid E_{tr} = sig = E_{tr} \cdot std()
112 E_{tr}MSE=np.mean(E_{tr}**2)
113 R2_tr=1-E_tr_MSE/(np.var(v_tr))
114 c_{tr} = np.mean((y_{tr} - y_{tr}.mean())*(y_hat_{tr} - y_hat_{tr}.mean()))
115 c_{tr} = c_{tr}/(y_{tr}.std()*y_{hat_{tr}.std()})
116 E_{te_min} = E_{te_min}()
117 E<sub>te</sub>max=E<sub>te</sub> \frac{max}{max}
118 \mid E_{te_mu} = E_{te_mean}
119 E_{te}=sig=E_{te}. std ()
120 E_te_MSE=np. mean (E_te **2)
121 R2_te=1-E_te_MSE/(np.var(v_te))
     c_t = np.mean((y_t = -y_t e.mean())*(y_h at_t = -y_h at_t e.mean()))
|c_{te}| = c_{te} / (v_{te} \cdot std() * v_{hat} = .std())
```

Performance figures [6]

To show the results in a better form, a DataFrame can be generated and printed:

The printed output is

```
R^2
             min
                     max
                                mean
                                         std
                                                  MSE
                                                               corr coeff
Training -7.504
                  10.469 -8.198e-14
                                       2.886
                                                8.326
                                                       0.922
                                                                    0.960
          -7.539
                  40.810
                           3.786e - 01
                                       3.905
                                               15.390
                                                       0.873
                                                                   0.9351
test
```

Performance figures [7]

Conclusions that can be drawn looking at the above results are:

- ▶ The error mean in the training subset is zero (as it should be); the error mean is slightly positive in the test subset because the mean of total UPDRS were evaluated using only the training subset, and it is therefore possible that total UPDRS means in the test subset are not exactly zero, after normalization.
- The error for the test dataset has one "outlier": the regressed value is much lower than the true value and the maximum error is 40.8. This only large error accounts for the higher error standard deviation and mean square value for the test dataset with respect to the training dataset.
- The coefficient of determination R^2 is close to around 0.9, as the correlation coefficient, which means that the regression is pretty good, in spite of the error at the single point.
- Since the error standard deviation is around 3-4 UPDRS points, this means that most of the times the regression error is around 6-8 points (twice the standard deviation, see also the histogram), which might still be accepted by a medical doctor. Regression is not very precise, but having an error of 6 points when total_UPDRS is equal to 50 points is still reasonable. Note that the regressand (UPDRS) has standard deviation equal to 10.34 points and mean value 28.50 points, therefore without regression we should predict a UPDRS value of 28.50 and we would have an error standard deviation equal to 10.34 points, which is about 2.5 times what the regression model provides.

Perform regression

Performance figures [8]

It has to be noticed that the model was obtained in the presence of collinearity and the performance might improve by removing it. On the contrary, total UPDRS was not regressed from voice parameters only, but also from **motor UPDRS**: if it is dropped then the performance is much worse. Using motor UPDRS to predict total UPDRS makes sense only if motor UPDRS can be automatically measured using sensors. If, on the other hand, a medical doctor is needed to measure motor UPDRS, then he/she can complete the visit to provide total UPDRS too, and the regression is useless.

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Parkinson's disease

Laboratory #1

Prepare and analyze the data Perform regression

To do

What you have to do [1]

- 1. Use your ID/matricola number as random seed.
- **2.** Drop features Jitter:DDP and Shimmer:DDA.
- 3. Add regression (exactly the same steps shown for LLS) using steepest descent, with a suitable stopping condition (you are responsible of your decisions). Try and use classes and inherited methods or methods (for example to plot histograms, to plot \hat{y} versus y, to generate the DataFrame with the results, etc), exploit the code you already wrote. Compare results obtained with steepest descent with those obtained using LLS (same input dataset, i.e. drop Jitter:DDP and Shimmer:DDA also for LLS).
- **4.** Then drop also Motor_UPDRS so that only voice parameters are used for regression. What happens?
- 5. Organize your software so that by changing the value of one variable (flag) you run your code with or without Motor_UPDRS. You will be asked to run the code with or without Motor_UPDRS.
- **6.** Try and see what happens if you do not shuffle the data. Possibly use another flag to turn on or off the shuffling.

Some final comments

- ▶ VERY IMPORTANT: in the applications analyzed in this course **execution speed** is not an issue (we are not dealing with big data); algorithm **complexity** is not an issue (think of the cost of a "simple" ultrasound machine, around 60-100 kEuros, and the cost of a "good" server with CUDAs etc, let's say 5-10 KEuros). If you need more CPUs or CUDAs, you simply buy them. The true issue is **reliability** and goodness of the result. As already remarked, patient's health is at stake.
- If an algorithm gives you $R^2 = 0.986$ and another one gives you $R^2 = 0.987$, the two algorithms are equivalent, you cannot say that the first is better than the second because R^2 has a lower third decimal digit, such a difference is irrelevant from an engineering point of view.
- ➤ You might be asked to run your script during the oral exam, explain what you did and why you did it and comment the results. Therefore it is important that you appropriately store your software in organized folders.

Next lab

In the next lab we will improve the results of this lab by using only nearest neighbours to perform regression.