Lab on Regression

[ credits: A. Geron, “Hands-On Machine Learning With Scikit-Learn and Tensorflow” ]
Steps to follow

Steps:

• frame your problem
• select a performance measure
• get the data
• descriptive statistics → discover and visualise the data to gain insights
• data pre-processing → prepare the data for ML algos
• model selection, model training
• model fine-tuning
• solution presentation
• (launch, monitor, maintain your newly deployed system)
The goal

Build a model of housing prices in California using the California census data.

Pretend to be a recently-hired data scientist in a real estate company in California, and you are asked to predict the price of a house given various parameters, having at your disposal the California Housing Prices dataset:


From this dataset, you know:

- population, median income, median housing price, much more.. for each block group (of 600-3000 people) - called “districts”

  - caveats: not updated (data from the 90s) and minor mods (added a categorical attribute, removed a few features for teaching purposes)

Your model should learn from this data and be able to predict the median housing price in any district, given all the other metrics
Building a model is not the goal. **Ask for the goal(s).** You do not work in an isolated system: stand up and ASK.

Good questions are:

- "**"what is my model being used for, eventually?"**
  
  ❖ this tells you how you concretely organise the **approach** to the problem, what **algorithms** you will select, what **performance** measure you will use to evaluate your model, how much **effort** you should invest in each (sub-)part of the work

- "**"what the current status of study of this problem is (if any)?"**
  
  ❖ this gives you a reference performance, as well as insights on how to solve the problem

- "**"what the expected full data pipeline which my solution will insert in?"**
  
  ❖ Data pipeline as a sequence of data processing components. Very common in ML. Async and self-contained components, data store as the only interface, different teams on different components, tactics for broken components, monitoring, etc
Frame the problem / make assumptions - step 2

Is it Supervised, Unsupervised, or Reinforcement Learning? Is it a classification task or a regression task? Should you use batch learning or online learning techniques?

(... think ...)
Frame the problem / **make assumptions** - step 2

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It is a **supervised** learning task..

- you are given labeled training examples: each instance comes with the expected output, i.e. the district’s median housing price
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It is a **univariate regression** task..

- you are asked to predict a value, and a single one per district
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- you are asked to predict a value, and a single one per district

**Batch learning** techniques should work just fine..

- data is small enough to fit in memory, there is no continuous flow of data coming in the system, there is no particular need to adjust to rapidly changing data
“I have the feeling I did nothing so far...”
“Quite the opposite! you did something already!”

I studied the problem.
Now I know where I am headed to.
Frame the problem / check assumptions - step 3

After knowing the full pipeline (from external input). now recheck your assumptions.

So far: **Supervised. Univariate regression. Batch learning**

- “is any other components in the overall work pipeline making my assumptions unnecessary or tactically wrong?”

**Example:**

Suppose your value predictions are going to be clustered into coarse-grain categories (e.g. just “cheap”, “medium”, “expensive”). Then, getting the price perfectly right is not important at all, you just need to get the category right, and your task should have been framed as a **classification task** instead!
**Notation**

- $m$: number of **instances**, i.e. examples in the training dataset
- $\mathbf{x}$: "input" variables, or "**features**" (*) (a vector per example)
- $y$: "output" variable, or "**label**" (a.k.a "target") per example
- $(\mathbf{x}^{(i)}, y^{(i)})$: the single $i^{th}$ training example ($i^{th}$ row)

Example:

\[
\mathbf{x}^{(1)} = \begin{pmatrix}
-118.29 \\
33.91 \\
1,416 \\
38,372
\end{pmatrix},
\]

\[
y^{(1)} = 156,400
\]

(*) strictly speaking, attribute \(\neq\) feature ..
Both are ways to measure the distance between two vectors (predictions and labels). Various distance measures, or norms, are possible:

• **RMSE** → root of sum of squares → Euclidean norm, or \( \ell_2 \) norm, often noted \( || \cdot ||_2 \)

• **MAE** → sum of absolutes → Manhattan norm, or \( \ell_1 \) norm, often noted \( || \cdot ||_1 \)

In general:

• \( \ell_k \) norm of a vector \( \mathbf{v} \) containing \( n \) elements: \( ||\mathbf{v}||_k = (|v_0|^k + |v_1|^k + \ldots + |v_n|^k)^{1/k} \)

• \( \ell_0 \) gives the # non-zero elements; \( \ell_\infty \) gives the max absolute value in the vector

Which one?

• The higher the norm index, the more it focuses on large values and neglects small ones. This is why RMSE is more sensitive to outliers than the MAE (if you have outliers, use MAE; when outliers are exponentially rare (like in a bell-shaped curve), RMSE performs very well and is generally preferred

A typical performance measure for regression problems is the Root Mean Square Error (**RMSE**)

- it gives an idea of how much error the system typically makes in its predictions: the smaller it is the better

Or the Mean Absolute Error (**MAE**) - aka Average Absolute Deviation

\[
\text{RMSE} (\mathbf{X}, h) = \sqrt{\frac{1}{m} \sum_{i=1}^{m} (h(x^{(i)}) - y^{(i)})^2}
\]

\[
\text{MAE} (\mathbf{X}, h) = \frac{1}{m} \sum_{i=1}^{m} |h(x^{(i)}) - y^{(i)}|
\]

\( h \to \) my hypothesis for \( y \)

Select a **performance measure**
You should start your code with a set-up part.

If you are trying all this real-time during the workshop, this means you should go to colab.
Then, it is time to **import the data** and check if this worked.
Inspect the data

Once data source has been downloaded, inspect it straight

```
l'head -20 datasets/housing/housing.csv
```

```bash
<table>
<thead>
<tr>
<th>longitude</th>
<th>latitude</th>
<th>housing_median_age</th>
<th>total_rooms</th>
<th>total_bedrooms</th>
<th>population</th>
<th>households</th>
<th>median_income</th>
<th>median_house_value</th>
<th>ocean_proximity</th>
</tr>
</thead>
<tbody>
<tr>
<td>-122.23</td>
<td>37.88</td>
<td>41.0</td>
<td>880.0</td>
<td>129.0</td>
<td>322.0</td>
<td>126.0</td>
<td>8.3252</td>
<td>452600.0</td>
<td>NEAR BAY</td>
</tr>
<tr>
<td>-122.22</td>
<td>37.86</td>
<td>21.0</td>
<td>7099.0</td>
<td>1106.0</td>
<td>2401.0</td>
<td>1138.0</td>
<td>8.3014</td>
<td>358500.0</td>
<td>NEAR BAY</td>
</tr>
<tr>
<td>-122.24</td>
<td>37.85</td>
<td>52.0</td>
<td>1467.0</td>
<td>190.0</td>
<td>496.0</td>
<td>177.0</td>
<td>7.2574</td>
<td>352100.0</td>
<td>NEAR BAY</td>
</tr>
<tr>
<td>-122.25</td>
<td>37.85</td>
<td>52.0</td>
<td>1274.0</td>
<td>235.0</td>
<td>558.0</td>
<td>219.0</td>
<td>5.6431</td>
<td>341300.0</td>
<td>NEAR BAY</td>
</tr>
<tr>
<td>-122.25</td>
<td>37.85</td>
<td>52.0</td>
<td>1627.0</td>
<td>280.0</td>
<td>565.0</td>
<td>259.0</td>
<td>3.8462</td>
<td>342200.0</td>
<td>NEAR BAY</td>
</tr>
<tr>
<td>-122.25</td>
<td>37.85</td>
<td>52.0</td>
<td>919.0</td>
<td>213.0</td>
<td>413.0</td>
<td>193.0</td>
<td>4.0368</td>
<td>269760.0</td>
<td>NEAR BAY</td>
</tr>
<tr>
<td>-122.25</td>
<td>37.84</td>
<td>52.0</td>
<td>2535.0</td>
<td>489.0</td>
<td>1094.0</td>
<td>514.0</td>
<td>3.6591</td>
<td>299200.0</td>
<td>NEAR BAY</td>
</tr>
<tr>
<td>-122.25</td>
<td>37.84</td>
<td>52.0</td>
<td>3104.0</td>
<td>687.0</td>
<td>1157.0</td>
<td>647.0</td>
<td>3.12</td>
<td>241400.0</td>
<td>NEAR BAY</td>
</tr>
<tr>
<td>-122.26</td>
<td>37.84</td>
<td>42.0</td>
<td>2555.0</td>
<td>665.0</td>
<td>1206.0</td>
<td>595.0</td>
<td>2.0804</td>
<td>226700.0</td>
<td>NEAR BAY</td>
</tr>
<tr>
<td>-122.25</td>
<td>37.84</td>
<td>52.0</td>
<td>3549.0</td>
<td>707.0</td>
<td>1551.0</td>
<td>714.0</td>
<td>3.6912</td>
<td>261100.0</td>
<td>NEAR BAY</td>
</tr>
<tr>
<td>-122.26</td>
<td>37.85</td>
<td>52.0</td>
<td>2202.0</td>
<td>434.0</td>
<td>910.0</td>
<td>402.0</td>
<td>3.2031</td>
<td>281500.0</td>
<td>NEAR BAY</td>
</tr>
<tr>
<td>-122.26</td>
<td>37.85</td>
<td>52.0</td>
<td>3503.0</td>
<td>752.0</td>
<td>1504.0</td>
<td>734.0</td>
<td>3.2705</td>
<td>241800.0</td>
<td>NEAR BAY</td>
</tr>
<tr>
<td>-122.26</td>
<td>37.85</td>
<td>52.0</td>
<td>2491.0</td>
<td>474.0</td>
<td>1098.0</td>
<td>468.0</td>
<td>3.075</td>
<td>213500.0</td>
<td>NEAR BAY</td>
</tr>
<tr>
<td>-122.26</td>
<td>37.84</td>
<td>52.0</td>
<td>696.0</td>
<td>191.0</td>
<td>345.0</td>
<td>174.0</td>
<td>2.6736</td>
<td>191300.0</td>
<td>NEAR BAY</td>
</tr>
<tr>
<td>-122.26</td>
<td>37.85</td>
<td>52.0</td>
<td>2643.0</td>
<td>626.0</td>
<td>1212.0</td>
<td>620.0</td>
<td>1.9167</td>
<td>159200.0</td>
<td>NEAR BAY</td>
</tr>
<tr>
<td>-122.26</td>
<td>37.85</td>
<td>50.0</td>
<td>1120.0</td>
<td>283.0</td>
<td>697.0</td>
<td>264.0</td>
<td>2.125</td>
<td>140000.0</td>
<td>NEAR BAY</td>
</tr>
<tr>
<td>-122.27</td>
<td>37.85</td>
<td>52.0</td>
<td>1966.0</td>
<td>347.0</td>
<td>793.0</td>
<td>331.0</td>
<td>2.775</td>
<td>152500.0</td>
<td>NEAR BAY</td>
</tr>
<tr>
<td>-122.27</td>
<td>37.85</td>
<td>52.0</td>
<td>1228.0</td>
<td>293.0</td>
<td>648.0</td>
<td>303.0</td>
<td>2.1202</td>
<td>155500.0</td>
<td>NEAR BAY</td>
</tr>
<tr>
<td>-122.26</td>
<td>37.84</td>
<td>50.0</td>
<td>2239.0</td>
<td>455.0</td>
<td>990.0</td>
<td>419.0</td>
<td>1.9911</td>
<td>158700.0</td>
<td>NEAR BAY</td>
</tr>
</tbody>
</table>
```

Mostly numbers, but also text (and repetitive..)

Or use pandas to deal with a DataFrame object - much easier to view and manipulate:

```
housing = load_housing_data()
housing.head()
```
Inspect the data further:

- use the pandas `info()` method on your dataframe (tip: it is called “housing”)
- use `value_counts()` to inspect special attributes (tip: “ocean proximity”)
- use `describe()` to get a summary of the numerical attributes
- plot all attributes using `matplotlib`
- review all the output you got, and try to extract interesting observations
Inspect the data

Each row represents one district. There are 10 attributes (columns):

- longitude, latitude, housing_median_age, total_rooms, total_bedrooms, population, households, median_income, median_house_value, ocean_proximity

The pandas `info()` method is useful to get a quick description of the data:

- total number of rows and columns, each attribute’s type, # non-null values

```
In[1]: housing.info()
```

```
<class 'pandas.core.frame.DataFrame'>
RangeIndex: 20640 entries, 0 to 20639
Data columns (total 10 columns):
longitude    20640 non-null float64
latitude     20640 non-null float64
housing_median_age 20640 non-null float64
total_rooms  20640 non-null float64
total_bedrooms 20433 non-null float64
population   20640 non-null float64
households   20640 non-null float64
median_income 20640 non-null float64
median_house_value 20640 non-null float64
ocean_proximity 20640 non-null object
dtypes: float64(9), object(1)
memory usage: 1.6+ MB
```

20,640 instances in the dataset

`total_bedrooms` attribute has only 20,433 non-null values, meaning that 207 districts are missing this feature
Inspect the data

All attributes numerical, except \texttt{ocean_proximity}. Its type is object, so it could hold any kind of py object, but you loaded from a CSV file so you know that it must be a text attribute. Probably categorical: use \texttt{value_counts()}

The \texttt{describe()} method shows a summary of the numerical attributes

null values are ignored
Inspect the data

Notice a few things in these histograms:

- **[A]** attribute not expressed in the **expected units** (USD). Ask who gave you the data..
  - it is in tens of thousands of USD

- **[B]** **[C]** were **capped**. Note you have a label here. It could be a problem. Your ML algo may learn that prices never go beyond that limit, which is wrong. Check if you need precise predictions also in those ranges. If yes, either you collect proper labels for the districts whose labels were capped, or you remove those examples from the training set (and the test set)

- The attributes (e.g. **[A]** **[C]**) have very **different scales** → **feature scaling**

- Many histograms are **skewed**: this may make it a bit harder for some ML algos to detect patterns → **transform attributes** to get symmetric distributions

- ...

**You are not wasting time!** Some steps here go towards a better understanding (and preprocessing) of the data you are feeding to ML
Train / Test splitting

Split into train and test set:

- in the standard way
- (NOT MANDATORY) using stratification
Create a test set

Take a subset of your data and put it aside.

Why? Because your brain is an amazing pattern detection system and you should avoid it to trick you!

• i.e. your brain is highly prone to overfitting. Looking at the test set, you might see some patterns and be biased towards some ML model. Then, when you estimate the generalisation error using the test set, your estimate will be too optimistic and you will launch a system that will eventually perform on new data much worse than expected → "data snooping bias"

```python
from sklearn.model_selection import train_test_split
train_set, test_set = train_test_split(housing, test_size=0.2, random_state=42)
```

>90% of ML practitioners do this.. but..
**Sampling: purely random vs stratified**

**Purely random sampling** works OK only if the dataset is large enough. If not, you risk to introduce a significant sampling bias.

Best is to use **stratified sampling**

- as the population is divided into homogeneous subgroups called **strata**, sample **the right number of instances from each stratum** to guarantee that the test set you build is **representative of the overall population**

E.g. if you are told that median income is an important attribute to predict house prices in a district, make sure you represent all categories of salary in your test set

- create e.g. 5 categories, assign examples to each
- then, use the sklearn’s **StratifiedShuffleSplit** class
- compare the income category proportions in the overall dataset, in the test set generated with stratified sampling, and in a test set generated using purely random sampling
Visualize to gain insight:

- hint: use the previous dataframe, and `.plot()` with proper options
  - e.g. scatter kind, and x and y should be the GPS coordinates

Practice R5

3 minutes

Time to code!
Visualising Geographical Data

For geolocated data, this often gives useful insights.

```
housing.plot(kind="scatter", x="longitude", y="latitude")
```
Visualising Geographical Data

Use `matplotlib` features to highlight high density patterns.

```python
housing.plot(kind="scatter", x="longitude", y="latitude", alpha=0.1)
```
Visualising Geographical Data

Display population by circles’ size and house price by colour on a heatmap.

Housing prices are very much related to location (e.g. close to the ocean) and to the population density.

Use a clustering algo to detect the main clusters?

Careful about ocean proximity attribute: perhaps useful but works different North vs South, so not a simple rule..
Visualising Geographical Data

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Careful about ocean proximity attribute: perhaps useful but works different North vs South, so not a simple rule..
Correlations

Inspect correlations

• hint: start with the housing dataframe, and explore .corr()

Code available also to do it visually, afterwards

• here, try to run the code, create the plots and examine them

Practice R6  3 minutes  Time to code!
Looking for **correlations** (numerically)

Dataset not huge $\rightarrow$ compute the **standard correlation coefficient** (aka **Pearson's r**) between every pair of attributes

- e.g. check how much each attribute correlates with the **median house value**

```
[53] corr_matrix = housing.corr()
```

<table>
<thead>
<tr>
<th>Correlation</th>
<th>Median house value</th>
<th>Median income</th>
<th>total_rooms</th>
<th>housing_median_age</th>
<th>households</th>
<th>total_bedrooms</th>
<th>population</th>
<th>longitude</th>
<th>latitude</th>
</tr>
</thead>
<tbody>
<tr>
<td>median_house_value</td>
<td>1.000000</td>
<td>0.687160</td>
<td>0.135097</td>
<td>0.114110</td>
<td>0.064506</td>
<td>0.047689</td>
<td>-0.026920</td>
<td>-0.047432</td>
<td>-0.142724</td>
</tr>
</tbody>
</table>

Median house value tends to go up when the median income goes up

Small negative correlation between the latitude and the median house value (i.e. prices have a slight tendency to go down when you go north)
The correlation coefficient only measures linear correlations

- it may completely miss out on nonlinear relationships (e.g., “if x is close to zero then y generally goes up”).

+1 or -1, and nothing to do with the slope

0, but axes clearly non independent.. (clearly, non-linear relationship)
Looking for **correlations** (visually)

11 attributes $\rightarrow 11^2$ plots, here focussing on just 4 $\rightarrow 16$ plots

Correlation is strong..

We see the cap at 500k

And we see more if we zoom..

(next)
You may want to try removing the corresponding districts to prevent your algos from learning to reproduce these “data quirks”
Explore attribute combinations

You need to clean

• here you can follow on slides and code

Practice R7  5 minutes  Time to code!
Explore attribute combinations

Create and add more meaningful attributes. And re-check correlation matrix.

Not bad:

- one of the new variables (bedrooms per room) is more anti-correlated to median house value than other old variables like the total number of rooms or bedrooms. Apparently houses with a lower bedroom/room ratio tend to be more expensive.

- The number of rooms per household is also more informative than the total number of rooms in a district - obviously the larger the houses, the more expensive they are.
Data preparation for ML algos: **missing features**

You need to prepare the data for ML algos, and one action is to attack the issues of attributes that miss the corresponding feature.

- here you can follow on slides and code

**Practice R8**

**7 minutes**

**Time to code!**
Data preparation for ML algos: missing features

Deal with **missing features**

- remember `total_bedrooms`, that had missing entries?

3 options:

1. Get rid of the corresponding districts → drop rows
2. Get rid of the whole attribute → drop a single column
3. Set the values to some value (zero, the mean, the median, etc.) → save it as you will need it for the test set too, or if/when the system goes live to replace new missing values
You need to prepare the data for ML algos, and another action is to **attack categorical attributes**

- here you can follow on slides and code
Data prep for ML: Handling Text and Categorical Attributes

Need to convert categories from text to numbers.

```python
from sklearn.preprocessing import OrdinalEncoder

ordinal_encoder = OrdinalEncoder()
housing_cat_encoded = ordinal_encoder.fit_transform(housing_cat)
housing_cat_encoded[:10]

array([[0.],
       [0.],
       [0.],
       [1.],
       [1.],
       [0.],
       [1.],
       [0.],
       [0.],
       [0.]])

ordinal_encoder.categories_

array([['<1H OCEAN', 'INLAND', 'ISLAND', 'NEAR BAY', 'NEAR OCEAN'],
       dtype=object])
```

This representation has the issue that ML algos will assume that 2 nearby values are more similar than 2 distant values.

```python
cat_encoder = OneHotEncoder(sparse=False)
housing_cat_1hot = cat_encoder.fit_transform(housing_cat)
housing_cat_1hot

array([[1., 0., 0., 0., 0.],
       [1., 0., 0., 0., 0.],
       [0., 0., 0., 0., 1.],
       ...
       [0., 1., 0., 0., 0.],
       [1., 0., 0., 0., 0.],
       [0., 0., 0., 1., 0.]])
```

A common solution is to create one binary attribute per category. This is called one-hot encoding, because only one attribute will be equal to 1 (hot), while the others will be 0 (cold).
Additional code available also to add new attributes, make a pipeline, transformations, etc.:

- check the code (it is abundant!)
Data preparation for ML algos: **Feature scaling**

ML algos don’t perform well when the input numerical attributes have very different scales

There are 2 common ways to get all attributes to have the same scale:

- **min-max scaling** (aka **normalisation**): values are shifted and rescaled so that they end up all ranging from 0 to 1 (or any other similar range)
  - useful e.g. for NN

- **standardisation**: it subtracts the mean value (so standardised values always have a zero mean), and then it divides by the standard deviation so that the resulting distribution has unit variance
  - much less affected by outliers w.r.t. normalisation

They can be used altogether.
Data preparation for ML algos: **Transformation pipelines**

Many data transformation steps that need to be executed in sequence and in the right order: sklearn provides the **Pipeline** class to help with such sequences of transformations.

```python
from sklearn.pipeline import Pipeline
from sklearn.preprocessing import StandardScaler

num_pipeline = Pipeline([  
    ('imputer', SimpleImputer(strategy="median")),
    ('attrs_adder', CombinedAttributesAdder()),
    ('std_scaler', StandardScaler()),
])

housing_num_tr = num_pipeline.fit_transform(housing_num)
```
Here we are!

**Practice R11**  **10 minutes**  **Time to code!**
Select and Train a Model

Here we are!

• I framed the problem
• I got the data and explored it
• I sampled a training set and a test set
• I wrote transformation pipelines to clean up and prepare data for ML

Now, data is ready, and next is to select and train a ML model

• start simple: a linear model

```python
def from sklearn.linear_model import LinearRegression
lin_reg = LinearRegression()
lin_reg.fit(housing_prepared, housing_labels)
```

Done! It is that simple!
I hope this gave you a feeling as of how much time to spend on **input data preprocessing** w.r.t **model selection**

“90% of my time in careful data preparation for ML” is probably a decent guess.
Select and Train a Model

How is the model working? Let’s try some predictions!

```python
# let's try the full preprocessing pipeline on a few training instances
some_data = housing.iloc[:5]
some_labels = housing_labels.iloc[:5]
some_data_prepared = full_pipeline.transform(some_data)

print("Predictions:", lin_reg.predict(some_data_prepared))
```

```
('Predictions:', array([210644.60459286, 317768.80697211, 210956.43331178, 59218.98886849, 189747.55849879]))
```

```python
[121] print("Labels:", list(some_labels))
```

```
('Labels:', [286600.0, 340600.0, 196900.0, 46300.0, 254500.0])
```

It works, although the predictions are not exactly accurate (e.g. the first is off by close to 40%!). Measure the RSME on the whole training set:

```python
[123] from sklearn.metrics import mean_squared_error

housing_predictions = lin_reg.predict(housing_prepared)
lin_mse = mean_squared_error(housing_labels, housing_predictions)
lin_rmse = np.sqrt(lin_mse)
lin_rmse
```

```
68628.19819848923
```

well.. I get this, as a typical prediction error, when median housing values range between $120,000 and $265,000.

Clearly not a great score (underfitting) but it is a start! (and it was relatively quick!)
Select and Train a Model

This point is one of the most tricky you will encounter in project on real-world datasets

- i.e. “I need to make a choice. What do I try next?”
- data science is general is done via experienced trials.. no recipes written in stones on most aspects.. And all may largely be dependant on your dataset..

Common practices help you, though. This case:

- **symptoms of underfitting**: this is already A LOT to drive your next choice!
- main ways to fix underfitting are:
  - select a more powerful model
  - feed the training algorithm with better features
  - reduce the constraints on the model

What would you choose?
Select and Train a Model

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• data science is general is done via experienced trials.. no recipes written in stones on most aspects.. And all may largely be dependant on your dataset..

Common practices help you, though. This case:

• symptoms of underfitting: this is already A LOT to drive your next choice!

• main ways to fix underfitting are:
  ❖ select a more powerful model → you can try, and much cheaper
  ❖ feed the training algorithm with better features → you can try, but expensive..
  ❖ reduce the constraints on the model → non regularized, so this is ruled out
Are you able to try a DecisionTreeRegressor yourself?

- HINT: sklearn documentation, StackOverflow, …
Select and Train a Model

Try a decision tree.

- Because it is a powerful model, capable of finding complex nonlinear relationships in the data

```python
from sklearn.tree import DecisionTreeRegressor

tree_reg = DecisionTreeRegressor(random_state=42)
tree_reg.fit(housing_prepared, housing_labels)
```

Now that the model is trained, evaluate it on the training set:

```python
housing_predictions = tree_reg.predict(housing_prepared)
tree_mse = mean_squared_error(housing_labels, housing_predictions)
tree_rmse = np.sqrt(tree_mse)
tree_rmse
```

WHAT?!? (see next)
Wait, what!? No prediction error AT ALL!? Is it PERFECT!?

- much more likely that the model has badly overfit the data.. I need to be sure though.. How?

I can’t touch the test set (until I am ready to test a final model I am decently confident about, and see how it may perform in production). So I need to only use a sub-part of the training set for training, and use the remaining data for.. model validation
Check how it is done in the code.
Model Evaluation w/ k-fold Cross-Validation

You can (statically):

- split the original training set into train and validation sets, train your model on the (smaller) training sub-set and evaluate it against the validation sub-set

Or (more dynamically)...

- randomly split the training set into k distinct subsets called "folds"
  - or think of strata if you think it is the case...
- permute and pick k-1 fields for training and evaluate on the remaining 1, i.e. train and evaluate your model k times
- the result is an array containing the k evaluation scores (you will average..)

```python
from sklearn.model_selection import cross_val_score

scores = cross_val_score(tree_reg, housing_prepared, housing_labels,
                          scoring="neg_mean_squared_error", cv=10)
tree_rmse_scores = np.sqrt(-scores)
```

Practical note: sklearn CV approach expects a utility function (greater is better) rather than a cost function (lower is better), so the scoring function is actually the opposite of the MSE (i.e. a negative value), which is why the code above computes -scores before calculating the square root.
Model Evaluation w/ k-fold Cross-Validation

Let's look at results:

Observation:

- **the good**: I have k models, so this technique gives me a standard deviation too
  - it came at the cost of multiple trainings, you cannot afford it always..

- **the bad**: the score with a more complex model is **worse** than that with a simpler one..
  - no progress?
    - wait - we are comparing LinearRegression w/o CV (RMSE 68628) with DecisionTreeRegressor w CV (RMSE 71407). For a fair comparison, to be sure, run CV also for LinearRegression.. you get 69052 with a std deviation of 2731.. (see next for reference)

**Not getting better**: it is comparably bad, even with a more complex model and with proper CV..
(the right comparison discussed in previous slide)

```python
lin_scores = cross_val_score(lin_reg, housing_prepared, housing_labels,
                            scoring="neg_mean_squared_error", cv=10)
lin_rmse_scores = np.sqrt(-lin_scores)
display_scores(lin_rmse_scores)

('Scores:', array([66782.73843989, 66960.118071 , 70347.95244419, 74739.57052552,
                  68031.13388938, 71193.84183426, 64969.63056405, 68281.61137997,
                  71552.91566558, 67665.10082067]))
('Mean:', 69052.46136345083)
('Standard deviation:', 2731.674001798344)
```
Are you able to try a RandomForestRegressor yourself?

- HINT: sklearn documentation, StackOverflow, …
Try one more model: Random Forest.

- it is an Ensemble Learning technique: build a model on top of many other models
  - it works by training many Decision Trees on random subsets of the features, then averaging out their predictions

```python
from sklearn.ensemble import RandomForestRegressor

forest_reg = RandomForestRegressor(n_estimators=100, random_state=42)
forest_reg.fit(housing_prepared, housing_labels)

housing_predictions = forest_reg.predict(housing_prepared)
forest_mse = mean_squared_error(housing_labels, housing_predictions)
forest_rmse = np.sqrt(forest_mse)
forest_rmse
```

1) this is MUCH better: Random Forests look very promising.

```python
from sklearn.model_selection import cross_val_score

forest_scores = cross_val_score(forest_reg, housing_prepared, housing_labels,
                                scoring="neg_mean_squared_error", cv=10)
forest_rmse_scores = np.sqrt(-forest_scores)
display_scores(forest_rmse_scores)
```

2) score on the training set is still << than on the validation sets: the model is still overfitting the training set.
Have you noted the execution of last cells of code became much slower over last few steps?

Have you understood:

- which step precisely?
- why?

*The timing is NOT going to get better.. note it down..*
Model Evaluation w/ k-fold Cross-Validation

Possible solutions:

• simplify the model
• constrain it (i.e. regularize it)
• get a lot more training data

What would you choose?
Towards fine-tuning..

Possible solutions:

• simplify the model → we just moved to a more complex.. perhaps try others?
• constrain it (i.e. regularize it) → this is fine tuning of hyperparameters
• get a lot more training data → this (alone) might work only in some cases..
Towards fine-tuning..

Possible solutions:

• simplify the model → we just moved to a more complex.. perhaps try others?
• constrain it (i.e. regularize it) → this is fine tuning of hyper-parameters
• get a lot more training data → this (alone) might work only in some cases..

Try out many other models from various categories of ML algos

• e.g. Support Vector Machines with different kernels
• e.g. possibly a Neural Network..

w/o spending too much time (yet) tweaking the hyper-parameters

The goal is to shortlist a few (5ish?) promising models, and continue the work in parallel with them altogether

• yes, the work is self-organising in various branches.. to be constantly compared..
Let’s assume that I have now a shortlist of (few) promising models. I need to fine-tune them.

One way to do that would be to fiddle with the hyper-parameters manually, until you find a great combination of their values that “magically” works

• this would be very tedious and time-consuming..

There are various ways to automatically do so:

• GridSearch
• Randomised Search
• Ensemble Methods
Grid Search does the search (via CV) of the best parameters across all permutations in the parameters’ grid

- all you need to do is tell which hyper-parameters you want it to experiment with, and what values to try out

```python
from sklearn.model_selection import GridSearchCV

param_grid = [
    # try 12 (3×4) combinations of hyperparameters
    {'n_estimators': [3, 10, 30], 'max_features': [2, 4, 6, 8]},
    # then try 6 (2×3) combinations with bootstrap set as False
    {'bootstrap': [False], 'n_estimators': [3, 10], 'max_features': [2, 3, 4]},
]

forest_reg = RandomForestRegressor(random_state=42)
# train across 5 folds, that's a total of (12+6)×5=90 rounds of training
grid_search = GridSearchCV(forest_reg, param_grid, cv=5,
                          scoring='neg_mean_squared_error',
                          return_train_score=True)
grid_search.fit(housing_prepared, housing_labels)
```

Note: do not worry about what these params mean: these are for RandomForest Regressor, others do have different ones..

Focus on the fact that this GridSearch launches 90ish training passes in one go!

The timing is NOT going to get better.. as expected..
Let's look at the score of each hyperparameter combination tested during the grid search:

```python
for mean_score, params in zip(cvres['mean_test_score'], cvres['params']):
    print(np.sqrt(-mean_score), params)
```

The best hyper-parameters combination found is this one.

```python
{'max_features': 8, 'n_estimators': 30}
```

The RMSE score for this combination is 49,682 (code not shown): slightly better than the score you got earlier using the default hyperparameter values (it was 50,182).

Some fine tuning worked!
Would you stop here!?

• *Hint: the chosen parameters happens to be the **maximum** values that were evaluated..*
Randomized Search

Grid Search is fine when exploring relatively few combinations. Move to **Randomized Search** if you want a larger hyperparameter search space.

Instead of trying out all possible combinations, it evaluates a given number of random combinations by selecting a random value for each hyperparameter at every iteration.

- if you let the randomised search run for, say, 1000 iterations, this approach will explore 1,000 different values for each hyperparameter (instead of just a few values per hyperparameter as with GridSearch).

```python
from sklearn.model_selection import RandomizedSearchCV
from scipy.stats import randint

param_distributions = {
    'n_estimators': randint(low=1, high=200),
    'max_features': randint(low=1, high=8),
}

forest_reg = RandomForestRegressor(random_state=42)
rnd_search = RandomizedSearchCV(forest_reg, param_distributions, n_iter=10, cv=5, scoring='neg_mean_squared_error', random_state=42)
rnd_search.fit(housing_prepared, housing_labels)
```

Not bad at all..
Ensemble Methods

This is another way to fine-tune your system: try to combine the models that perform best, as the group (or “ensemble”) will often perform better than the best individual model.

- we saw this already: RandomForest performed better than the individual DecisionTrees it relied on.
- especially good if the individual models make very different types of prediction errors.
Analyze the best models and their prediction errors

Crucial to understand why a model is working better than others

- “who drove this model to the point it performs the best?”

**Feature importance**: e.g. RandomForestRegressor can indicate the relative importance of each attribute for making accurate predictions

```python
feature_importances = grid_search.best_estimator_.feature_importances_
array([7.334432355e-02, 6.29090705e-02, 4.11437985e-02, 1.46726854e-02, 1.41064935e-02, 1.48742809e-02, 1.42575993e-02, 3.66150989e-01, 5.64191792e-02, 1.08792957e-01, 5.33510773e-02, 1.03114883e-02, 1.64780994e-01, 6.028803867e-05, 1.96041560e-03, 2.85647466e-03])
```

You may want to try dropping some of the less useful features

(e.g. apparently only one ocean_proximity category is really useful, so you could drop the others)

But do more!

- add extra features or, on the contrary, get rid of uninformative ones, cleaning up outliers, etc
OK.

You eventually have a system that performs sufficiently well. Now is the time to evaluate the final model on the test set.
Evaluate Your System on the Test Set

Easy, and nothing technical different wrt what we did already

- get features and labels from your test set, now
- run your full pipeline to transform the data \texttt{(transform(), not fit\_transform())}
- evaluate the final model on the test set

```
[150] final_model = grid_search.best_estimator_

X_test = strat_test_set.drop("median_house_value", axis=1)
y_test = strat_test_set["median_house_value"].copy()

X_test_prepared = full_pipeline.transform(X_test)
final_predictions = final_model.predict(X_test_prepared)

final_mse = mean_squared_error(y_test, final_predictions)
final_rmse = np.sqrt(final_mse)
```

Not bad. Let’s stop here.

Communicate out. You got an idea of features importances \texttt{(median income as main predictor)}, you studied and excluded some features \texttt{(e.g. some of the ocean vicinity ones)}, plenty of lesson learned \texttt{(what worked and what not)}, you got a performance that can be compared with others

Depending on the case, \texttt{(your boss will) consider to switch the production system to this one.} \texttt{(then, plenty of monitor, re-checks, etc..)}
That’s it, for our Lab on Regression