

# **Impact of Individual Attributes**

We may be interested in understanding the impact of individual attributes



■ E.g. is the price more impacted by age, distance from MRT stations, or what?

# Let's start by repeating our basic linear regression approach

```
In [2]: import numpy as np
        import pandas as pd
        from sklearn.linear model import LinearRegression
        from sklearn.model selection import train test split
        from sklearn.metrics import r2 score
        data = pd.read csv('data/real estate.csv', sep=',')
        cols = data.columns
       X = data[cols[:-1]]
       y = np.log(data[cols[-1]])
       X tr, X ts, y tr, y ts = train test split(X, y, test size=0.34, random state=42)
        m = LinearRegression()
       m.fit(X tr, y tr);
        y pred tr, y pred ts = m.predict(X tr), m.predict(X ts)
        print(f'R2: {r2 score(y tr, y pred tr):.3} (training), {r2 score(y ts, y pred ts):.3} (test)')
        R2: 0.691 (training), 0.645 (test)
```

- We load the dataset, separate input & output, split the training and test set
- The we fit the model and evaluate the results

We can think of looking at the coefficients

# We can think of looking at the coefficients

Intuitively:

- The higher the (absolute) weight, the more important the attribute
- A positive coefficient corresponds to a positive correlation (w.r.t. the price)
- A negative coefficient corresponds to a negative correlation

# We can obtain the coefficient from the LinearRegression object

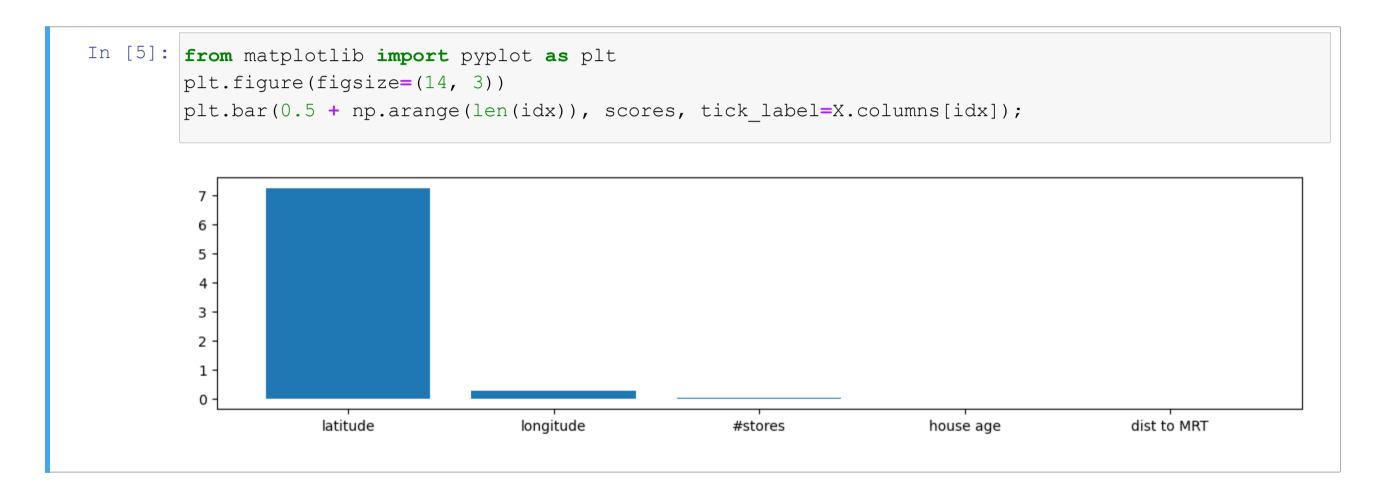
Then we can sort them by decreasing absolute value:

```
In [4]: idx = np.argsort(-np.abs(coeffs)) # Sort indexes by decreasing absolute coefficient
    scores = m.coef_[idx] # Coefficients, in the order defined by idx
```

### We can show the results via a bar chart

```
In [5]: from matplotlib import pyplot as plt
        plt.figure(figsize=(14, 3))
        plt.bar(0.5 + np.arange(len(idx)), scores, tick_label=X.columns[idx]);
          6
          5 -
                      latitude
                                         longitude
                                                                               house age
                                                                                                  dist to MRT
                                                             #stores
```

### We can show the results via a bar chart



- The plot seems to suggest that "latitude" has by far the largest impact
- ...But that does not match with our original dataset inspection!

# There is a (huge) pitfall with this approach

- Each coefficient depends on the attribute importance, but also on its scale
- I.e. an attribute with larger values may end up with a smaller coefficient

### A possible solution consists in preprocessing the data

...So that all inputs have similar scale

■ A first common approach: normalization (all values in the 0-1 range)

$$x' = \frac{x - \min(x)}{\max(x) - \min(x)}$$

■ A second one: standardization (the mean becomes 0, the st. dev. becomes 1)

$$x' = \frac{x - \text{mean}(x)}{\text{std}(x)}$$

# Normalization/Standardization

# scikit-learn provides simple tools to take care of this step

The are called MinMaxScaler and StandardScaler:

```
In [6]: from sklearn.preprocessing import StandardScaler # for standardization
    scaler = StandardScaler()
    X_tr_s = scaler.fit_transform(X_tr)
```

### On the training set:

- We calibrate the scaler (i.e. compute the mean and st. dev.)
- The we apply the transformation

```
In [7]: X_ts_s = scaler.transform(X_ts)
```

# On the test set we simply apply the transformation

We do not want information about the test data to "leak" into the model

# The Correct Approach

# We can then repeat the process using the scaled data

```
In [8]: m = LinearRegression()
    m.fit(X_tr_s, y_tr);

y_pred_ts = m.predict(X_ts_s)
    print(f'R2 on the test set: {r2_score(y_ts, y_pred_ts):.3}')

R2 on the test set: 0.645
```

- The quality of the predictions is the same as before
- ...Since we have applied only a linear transformation to the input

# The Correct Approach

# Now we can build and inspect the same plot as before

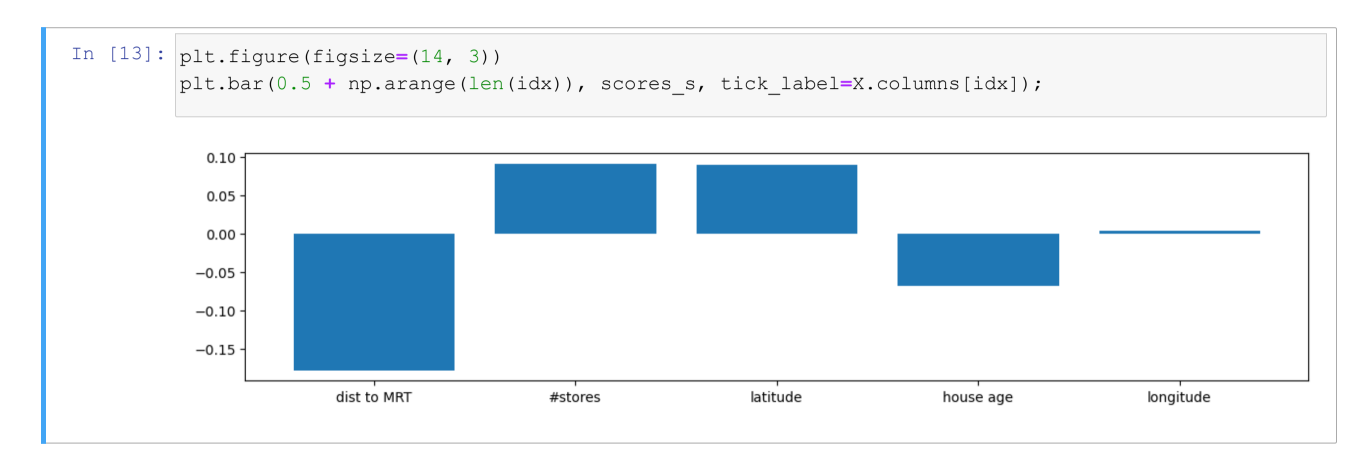
```
In [12]: | idx = np.argsort(-np.abs(m.coef))
          scores s = m.coef [idx]
          plt.figure(figsize=(14, 3))
          plt.bar(0.5 + np.arange(len(idx)), scores s, tick label=X.columns[idx]);
             0.10
             0.05
             0.00
            -0.05
            -0.10
            -0.15
                         dist to MRT
                                              #stores
                                                                  latitude
                                                                                    house age
                                                                                                        longitude
```

■ The order of the attributes has changed!

# LASSO

## **Attribute Selection**

# Let's look again at the last plot



- Is it worth keeping "longitude", given its low importance?
- Collecting useless attributes has still a cost!

# **Attribute Selection**

# Choosing which attributes to keep is known as attribute selection

There are multiple solution approaches

- An exact (but costly) one: consider all possible sets and repeat training
- A cheap (but heuristic) one: remove attributes with small (abs.) coefficients

### In the case of linear regression, we have a third option

We can act on the structure of the loss (error) function used during training:

$$L(w) = MSE(\theta) + \alpha \|\theta\|_1$$

- Where  $\|\theta\|_1 = \sum_{j=1}^n |\theta_j|$
- lacktriangle The additional term  $\|w\|$  is introduced to make mode model "well behaved"
- ...And for this reason it is called a reguralizer
- In particular, since we are using the L1 norm, we call it an L1 regularizer

# **LASSO**

### At training time, we will need to solve

$$\operatorname{argmin}_{\theta} MSE(\theta) + \alpha \|\theta\|_1$$

For reasons not discussed in detail:

- If moving a coefficient away from 0 does not provide enough error improvement
- ...Then the attribute will not be used at all
- lacktriangle What "enough" means in practice depends on the value of lpha

# As a result, we naturally end-up doing attribute selection

The approach is known as Least Absolute Shrinkage and Selection Operator

...Or LASSO for short :-)

# **Using LASSO**

# Scikit-learn provides a convenient LASSO implementation

```
In [14]: from sklearn.linear_model import Lasso
m2 = Lasso(alpha=1.0)
```

- lacktriangle When building the model, we need to pick a value for lpha
- lacktriangle Higher lpha values promote using fewer attributes
- ...But they also reduce the prediction quality

# For now, we will fix $\alpha$ via trial-and-error approach:

```
In [15]: m2.fit(X_tr_s, y_tr);
y_pred_tr2 = m2.predict(X_tr_s)
print(f'R2 on the training set: {r2_score(y_tr, y_pred_tr2):.3}')
R2 on the training set: 0.0
```

■ This is too low, and therefore  $\alpha = 1.0$  is too large

# Choosing $\alpha$

```
In [16]: m2 = Lasso(alpha=0.1)
    m2.fit(X_tr_s, y_tr);
    y_pred_tr2 = m2.predict(X_tr_s)
    print(f'R2 on the training set: {r2_score(y_tr, y_pred_tr2):.3}')

R2 on the training set: 0.549
```

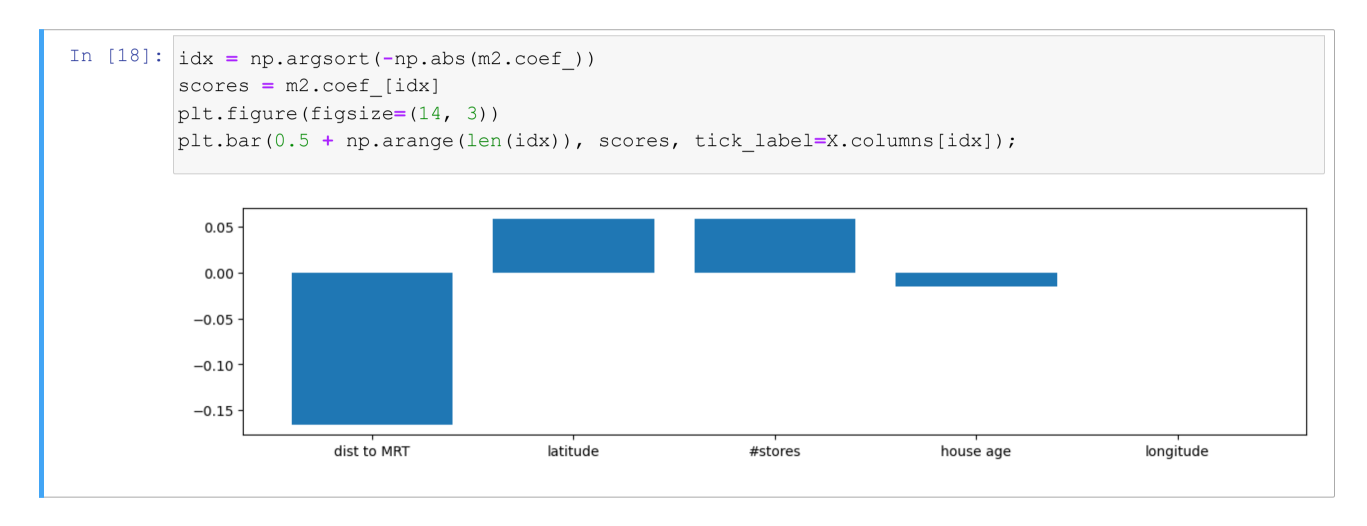
■ The prediction quality is still much lower than the original one

```
In [17]: m2 = Lasso(alpha=0.05)
    m2.fit(X_tr_s, y_tr);
    y_pred_tr2 = m2.predict(X_tr_s)
    print(f'R2 on the training set: {r2_score(y_tr, y_pred_tr2):.3}')
R2 on the training set: 0.643
```

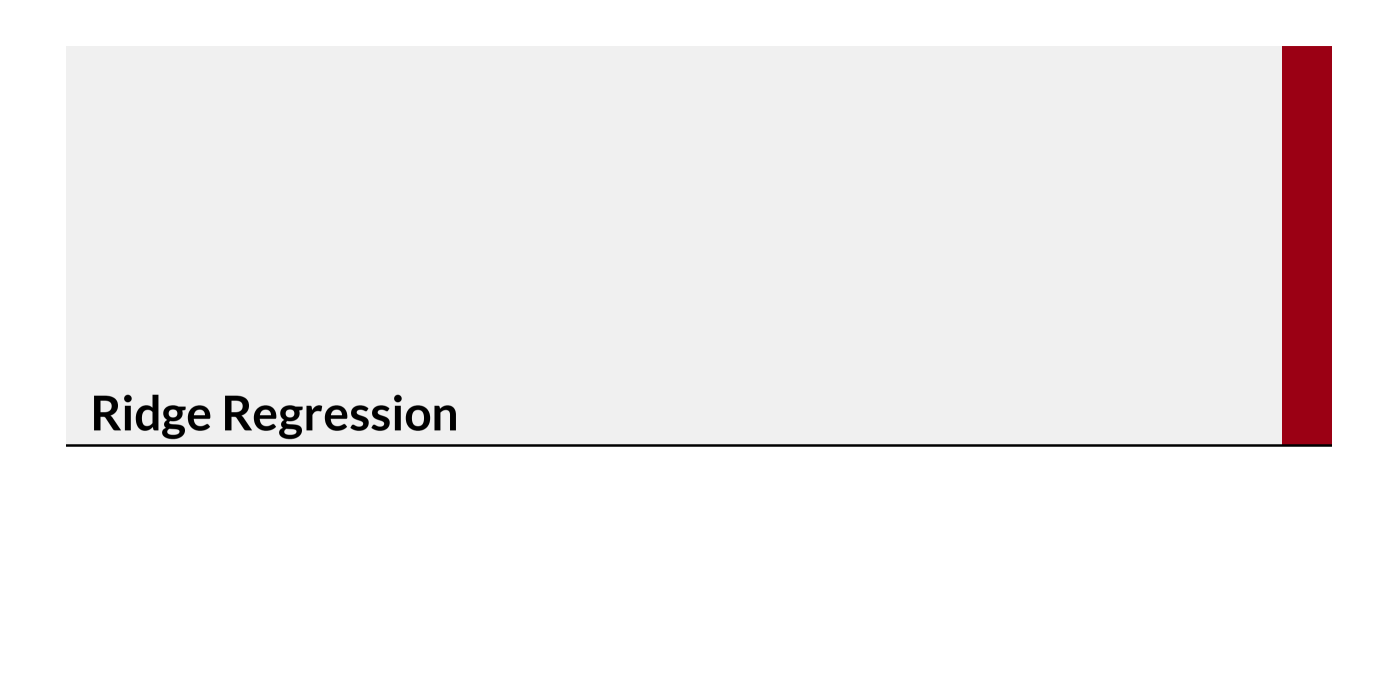
■ Now we are back to the original value (more or less)!

# **Impact of Each Attribute**

### Once we are satisfied with the $\alpha$ values, we can inspect the coefficients



- The "longitude" attribute has been dropped
- When there are many attributes, the LASSO method can be extremely useful



# **Overfitting**

# We have already observed how our model has a modest degree of overfitting

```
In [19]: print(f'R2 on the training set (linear regression): {r2_score(y_tr, y_pred_tr):.3}')
    print(f'R2 on the test set (linear regression): {r2_score(y_ts, y_pred_ts):.3}')

R2 on the training set (linear regression): 0.691
R2 on the test set (linear regression): 0.645
```

### Overfitting arises when:

- The model learns patterns in the training set
- ...That do not repeat on unseen data (e.g. since they due to random sampling)

# This becomes more critical with many attributes and scarce data

- A linear regression model with n coefficients
- $\blacksquare$  ...Can interpolate perfectly any collections of n examples
- ...But typically, that means we are just learning noise :-(

# First, Let's Make It Worse

# First, let's amplify the problem, so as to make any change more significant

In particular, we will reduce the amount of available training data

```
In [20]: X_tr2, X_ts2, y_tr2, y_ts2 = train_test_split(X, y, test_size=0.938, random_state=42)

scaler2 = StandardScaler()
X_tr_s2 = scaler2.fit_transform(X_tr2)
X_ts_s2 = scaler2.transform(X_ts2)
print(f'Training set size: {len(X_tr_s2)}')

m3 = LinearRegression()
m3.fit(X_tr2, y_tr2)

y_pred_tr3, y_pred_ts3 = m3.predict(X_tr2), m3.predict(X_ts2)
print(f'R2: {r2_score(y_tr2, y_pred_tr3):.3} (training), {r2_score(y_ts2, y_pred_ts3):.3} (test)

Training set size: 25
R2: 0.628 (training), 0.509 (test)
```

- Having access to little data is not uncommon
- Input are easy to obtain, but targets are often expensive to collect

# Ridge Regression

# We will show how to reduce overfitting by using ridge regression

...Which is obtained from linear regression by adding a quadratic reguralizer:

$$\operatorname{argmin}_{\theta} MSE(\theta) + \alpha \|\theta\|_{2}^{2}$$

where 
$$\|\theta\|_{2}^{2} = \sum_{j=1}^{2} \theta_{j}^{2}$$

- Since the  $||w||_2$  is an L2 norm, this is called an L2 regularizer
- It looks similar to the L1 regularizer in the LASSO approach
- ...But it has a very different effect

# This regularizer prevents weights from having a large absolute value

- Smaller (absolute) weights result in smoother behavior
- ...Which typically leads to better generalization

# **Using Ridge Regression**

# Scikit-learn provides an implementation for Ridge Regression

```
In [21]: from sklearn.linear_model import Ridge
m4 = Ridge(alpha=1.0)
m4.fit(X_tr_s2, y_tr2);
```

It works similarly to the Lasso class

- lacksquare When we build the model, we need to specify a value for lpha
- ...Then we train the model as usual

```
In [22]: y_pred_tr4 = m4.predict(X_tr_s2)
    y_pred_ts4 = m4.predict(X_ts_s2)
    print(f'R2 on the training set: {r2_score(y_tr2, y_pred_tr4):.3}')
    print(f'R2 on the test set: {r2_score(y_ts2, y_pred_ts4):.3}')

R2 on the training set: 0.627
    R2 on the test set: 0.534
```

 $\blacksquare$  The default value for  $\alpha$  is 1.0

# **Cross-Validation and Hyper-Parameter Tuning**

# This time, we will follow a systematic approach to pick a value for $\alpha$

- Our goal is to improve the model results on unseen examples
- lacksquare So we can choose lpha to minimize the error on examples not in the training set

# It would be tempting to optimize $\alpha$ for the test set...

- ...But it would also be very wrong
- The test set is our ultimate approach to assess generalization
- For this, its content stay unseen whenever we calibrate parameters

# We can however split yet another set on this purpose

- This would not be used to train the model
- ...But only for tuning the training parameters (also called hyper-parameters)

### This is called a validation set

■ But there is a drawback: we have even less data available for training

# **Cross-Validation and Hyper-Parameter Tuning**

### As an alternative, we can use k-fold cross-validation:

- lacksquare We partition the training data in k sets, called folds
- For i = 1..k:
  - $\blacksquare$  We separate the i-th fold
  - We train a model on the remaining data
  - We evaluate the prediction quality on the separated fold
  - Then we put the fold back

During each iteration, the examples in the i-th folder are effectively unseen

### At the end of the process we have:

- k models (that typically we don't use)
- lacktriangleright k prediction quality evaluations (of which we can take the mean)

We can use cross validation to assess how good a model is with unseen data

# **Applying Cross-Validation and Hyper-Parameter Tuning**

# Scikit learn allows us to automate this process with a few lines of code

- First we specify which parameters we want to tune
- ...And which range we want to investigate

We do so by building a dictionary:

```
In [23]: param_ranges = {'alpha': np.logspace(0, 2, 100)}
```

- The dictionary keys are the parameters to be tuned
- The entries are the values to be tested

Then we build a GridsearchCV object:

```
In [24]: from sklearn.model_selection import GridSearchCV

gscv = GridSearchCV(estimator=Ridge(), param_grid=param_ranges, cv=5)
```

■ The cv parameter specifies the number of folds

# **Applying Cross-Validation and Hyper-Parameter Tuning**

# The we can "train" the GridSearchCv object

- This will try all combinations of the values in parameter ranges
- ...And pick the parameters leading to the best cross-validation score

```
In [25]: gscv.fit(X_tr_s2, y_tr2);
```

The best parameters can be accessed via the best\_params\_ field:

```
In [26]: gscv.best_params_
Out[26]: {'alpha': 27.185882427329414}
```

# **Applying Cross-Validation and Hyper-Parameter Tuning**

### By default, the GridSearchCV object:

- Trains an estimator with the best parameters on all data
- This is the model that we are going to use

Calling predict on Gridsearchev calls the same method on the best estimator:

```
In [27]: y_pred_tr4 = gscv.predict(X_tr_s2)
    y_pred_ts4 = gscv.predict(X_ts_s2)
    print(f'R2 on the training set: {r2_score(y_tr2, y_pred_tr4):.3}')
    print(f'R2 on the test set: {r2_score(y_ts2, y_pred_ts4):.3}')

R2 on the training set: 0.516
    R2 on the test set: 0.638
```

- This is close to our original approach on unseen data
- ...Even if we are using only 25 data points!

# **Conclusions and Take-Home Messages**

- Attribute normalization to improve interpretability
  - Linear Regression is a rather interpretable ML model...
  - ...Provided that attributes are processed to have a similar scale
- L1 regularization
  - L1 regularization can be employed for automatic attribute selection
  - ...And it also helps reducing overfitting
- L2 regularization
  - L2 regularization is mostly about reducing overfitting
  - ...But for that it works better than L1 regularization
- (Cross)-validation
  - Validation data can be used to calibrate hyper-parameters
  - ...Such as the regularization weights
  - Cross-validation enables a more efficient use of data