Linear Methods

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Overview

- Preliminaries
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- Model selection
 - Loss functions and robustness

Preliminaries

- We will go very deep into linear models.
- Most of you probably have seen linear models in some form, but we will start from scratch
 to further illustrate key concepts such as bias and variance.
- Using linear examples, we will discuss the basic machine learning concepts of model selection, cross-validation, and loss functions.

Notation

- Capital letters, e.g., X denote variables.
- Lower-case letters e.g., x, denote observations.
- Dummy index j denotes different variables, e.g., X_j
- Dummy index *i* denotes different observations, e.g., *x_i*
- Bolded variables are vector/matrices, e.g., y, X

Linear Regression

Linear Regression

Simplest possible model between target and feature

$$Y = f(X_1, X_2, ..., X_p) = \beta_0 + \sum_{j=1}^{p} \beta_j X_j$$

X_i can be:

- Quantitative inputs
- Transformations of quantitative inputs, e.g., log, exp, powers, etc. Basis expansions, e.g., $X_2 = X_1^2, X_3 = X_1^3$
- Interactions between variables, e.g., X_1X_2
- Encoding of levels of inputs

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Supervised learning

- Given a set of paired observations $\{x_{ij}, y_i\}$, what are the model parameters (in this case, the coefficients β_i) that are "optimal"?
- "Optimal" is typically defined as minimization of some loss function (also known as cost function) that measures the error of the model.

Least squares regression

Consider the simple case of

$$Y = \beta_0 + \beta_1 X_1$$

In least squares regression, the loss function is defined as the sum squared error given the Nobservations:

$$L(Y, \hat{f}(X)) = \sum_{i=1}^{N} (y_i - f(x_i))^2$$
$$= \sum_{i=1}^{N} (y_i - \beta_0 - \beta_1 x_{i1})^2$$

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What are the optimal parameters β_0 and β_1 ?

$$\frac{\partial L}{\partial \beta_0} = \sum_{i=1}^{N} 2(y_i - \beta_0 - \beta_1 x_{i1})(-1) = 0$$

$$\implies \sum_{i=1}^{N} y_i = N\beta_0 + \sum_{i=1}^{N} \beta_1 x_{i1}$$

$$\implies \beta_0 = \bar{y} - \beta_1 \bar{x}_1$$

$$\frac{\partial L}{\partial \beta_1} = \sum_{i=1}^{N} 2(y_i - \beta_0 - \beta_1 x_{i1})(-x_{i1}) = 0$$

$$\implies \beta_1 = \frac{\sum_{i=1}^{N} x_{i1} y_i - N\bar{x}_1 \bar{y}}{\sum_{i=1}^{N} x_{i2}^2 - N\bar{x}_1^2}$$

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Reformulating the general multiple linear regression as a vector equation...

Considering N observations of

$$y_i = \beta_0 + \beta_1 x_{i1} + \beta_2 x_{i2} + \dots + \beta_p x_{ip}$$

Let

$$\mathbf{y} = \begin{pmatrix} y_1 \\ y_2 \\ \dots \\ y_n \end{pmatrix}, \boldsymbol{\beta} = \begin{pmatrix} \beta_0 \\ \beta_1 \\ \dots \\ \beta_p \end{pmatrix}, \mathbf{X} = \begin{pmatrix} 1 & x_{11} & x_{12} & \dots & x_{1p} \\ 1 & x_{21} & x_{22} & \dots & x_{2p} \\ \vdots & & & & \\ 1 & x_{N1} & x_{N2} & \dots & x_{Np} \end{pmatrix},$$

So,

$$y = X\beta$$

Note that **y** is a $N \times 1$ vector, β is a $(p+1) \times 1$ vector, and **X** is a $N \times (p+1)$ matrix.

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Reformulating the general multiple linear regression as a vector equation...

$$L = RSS = (\mathbf{y} - \mathbf{X}\boldsymbol{\beta})^T (\mathbf{y} - \mathbf{X}\boldsymbol{\beta})$$

Assuming (for the moment) that X has full column rank, and hence X^TX is positive definite, It can be shown using the same principles that the following unique solution for β is:

$$\hat{\boldsymbol{\beta}} = (\mathbf{X}^T \mathbf{X})^{-1} \mathbf{X}^T \mathbf{y}
\hat{\mathbf{y}} = \mathbf{X} \hat{\boldsymbol{\beta}} = \mathbf{X} (\mathbf{X}^T \mathbf{X})^{-1} \mathbf{X}^T \mathbf{y}$$

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Graphic representation of MLR with two dependent variables

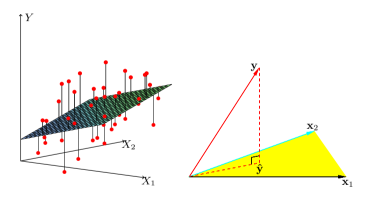


Figure: MLR minimizes sum square of residuals. The projection $\hat{\mathbf{y}}$ represents the vector of the least squares predictions onto the hyperplane spanned by the input vectors \mathbf{x}_1 and \mathbf{x}_2 . [1].

Validity of least squares criterion

- Observations are independently drawn at random.
- Variance of **y** is constant given by σ^2 .

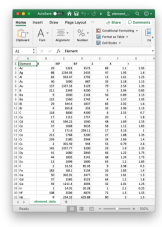
$$\operatorname{var}(\hat{\boldsymbol{\beta}}) = (\mathbf{X}^T \mathbf{X})^{-1} \sigma^2$$

 \bullet and σ is estimated using:

$$\sigma^2 = \frac{1}{N - p - 1} \sum_{i=1}^{N} (y_i - \hat{y}_i)^2$$

Example materials data

- Target: Bulk modulus of elements (from Materials Project)
- Candidate features:
 - Melting point (MP)
 - Boiling point (MP)
 - Atomic number (Z)
 - Electronegativity (χ)
 - Atomic radius (r)
- Question: Why these features?
- We will add some transformations of these inputs as well, i.e., the square and square root of the electronegativity and atomic radius.



Using pandas for easy data manipulation

```
import pandas as pd
# Read in data and set first column as index.
data = pd.read_csv("element_data.csv", index_col=0)
# Generate transformations as additional columns.
data["X^2"] = data["X"] ** 2
data["sqrt(X)"] = data["X"] ** 0.5
data["r^2"] = data["r"] ** 2
data["sqrt(r)"] = data["r"] ** 0.5
# Define our features, which is all the columns
# excluding K. which is the target.
features = [c for c in data.columns if c != "K"]
x = data[features]
v = data["K"]
```

MLR in scikit-learn

```
from sklearn import linear_model
reg = linear_model.LinearRegression()
reg.fit(x, y)
print(ref.coef_)
print(reg.intercept_)
```

- Note that x should contain the features only there is no need to add a 1 column for the intercept. By default, the parameter fit_intercept in sklearn.linear_model.LinearRegression is True. You can set it to False to do a MLR without intercept.
- Documentation: link.

Hypothesis Testing for Coefficients

- To derive insights into a model, we often want to know which of the input parameters are the most relevant to the target.
- Under assumptions of the errors in y follow a Gaussian distribution $N(0, \sigma^2)$, the errors in $\hat{\beta}$ also have a Gaussian distribution $N(\beta, (\mathbf{X}^T\mathbf{X})^{-1}\sigma^2)$
- Hypothesis testing can be carried out for whether a particular β_j is 0 using the following test statistic:

$$t_j = rac{\hat{oldsymbol{eta_j}}}{\sigma \sqrt{oldsymbol{v_j}}}$$

where v_j is the jth diagonal element of $(\mathbf{X}^T\mathbf{X})^{-1}$. t_j has a t distribution with N-p-1 degrees of freedom (dof).

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Hypothesis Testing for Groups of Coefficients

- More often, we want to test groups of coefficient for significance. E.g., to the k levels of a categorical variable.
- We will use the following *F* statistic:

$$F = \frac{(\text{RSS}_0 - \text{RSS}_1)/(p_1 - p_0)}{\text{RSS}_1/(N - p_1 - 1)}$$

where RSS_0 is the RSS of the larger model with p_0+1 parameters and RSS_1 is the RSS of the smaller model with p_1+1 parameters with p_0-p_1 parameters set to zero. The F statistic has a distribution of $F_{p_1-p_0,N-p_1-1}$.

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• Consider the estimator $\hat{\theta}$ for a variable θ .

MSE =
$$E(\hat{\theta} - \theta)^2$$

= $var(\hat{\theta}) + [E(\hat{\theta}) - \theta]^2$

 The MSE can be broken down into the variance of the estimate itself and the square of the bias.

Gauss-Markov Theorem

The least squares estimator has the smallest variance among all linear unbiased estimators.

• However, there can be estimators that are biased with smaller MSE.

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Model selection

Model selection

Model performance

- We will take a brief digression into model assessment and selection before continuing on to other linear methods.
- Model performance is related to its performance on *independent test data*, i.e., one cannot simply report a model's performance on training data alone.
- Note that this section is deliberately limited to high level concepts that are needed to continue further in exploration of linear methods. A more detailed discussion will be performed in later lectures.

Typical measures of model performance

Mean squared error (MSE):

$$L(Y, \hat{f}(X)) = \frac{1}{N} \sum_{i=1}^{N} (y_i - f(x_i))^2$$

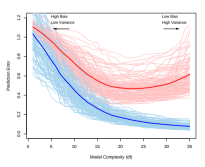
• Mean absolute error (MAE):

$$L(Y, \hat{f}(X)) = \frac{1}{N} \sum_{i=1}^{N} |y_i - f(x_i)|$$

- Test error: *L* over independent test set.
- Training error: L over training set.

Training and test errors with model complexity

- Model complexity increases as the number of parameters increases (e.g., number of independent variables in MLR).
- Training errors always decrease with increasing model complexity.
- However, test errors do not have a monotonic relationship with model complexity. Test
 errors are high when model complexity is too low (underfitting) or too high (overfitting).



Under-fitting versus over-fitting

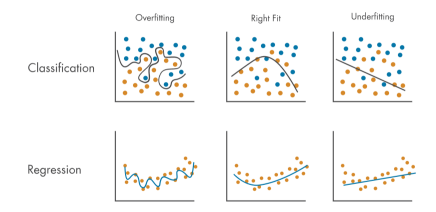


Figure: Source: Mathworks

Training, validation and test data

- Model selection: estimating the performance of different models in order to choose the best one.
- Model assessment: having chosen a final model, estimating its prediction error (generalization error) on new data.
- Ideal data-rich situation: Divide data into three parts:
 - Training set: For training the model.
 - Validation set: For estimating prediction error to select the model.
 - Test set: For assessing the generalization error of the final model.
- Typical training:validation:test split is 50:25:25 or 80:10:10, or in very data-poor situations, maybe even 90:5:5.
- Note that at no point in the model fitting process should the test set be "seen".

K-fold cross validation (CV)

- Simplest and most widely used approach for model validation.
- Data set is split into K buckets (usually by random).
- Typical values of K is 5 or 10. K = N is known as "leave-one-out" CV.

• CV score is computed on the validate data set after training on the train data:

$$CV(\hat{f}^{-k(i)}, \alpha) = \frac{1}{N_{k(i)}} \sum_{i=1}^{N_{k(i)}} L(y_i, \hat{f}^{-k(i)}(x_i, \alpha))$$

• assuming the k^{th} data bucket has $N_{k(i)}$ data points and $\hat{f}^{-k(i)}$ refers to the model fitted with the k^{th} data left out $(N - N_{k(i)})$ data in fitting).

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CV in scikit-learn

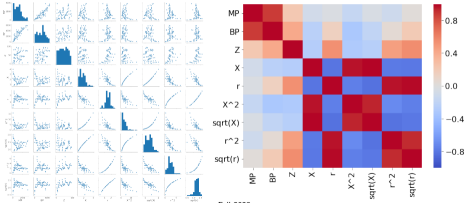
```
    kfold = KFold(n_splits=5, shuffle=True, random_state=42)
    cv_results = cross_validate(ridge, z, y, cv=kfold)
    Note that we have customized the KFold object passed to the cross_validate method. The
```

- Note that we have customized the KFold object passed to the cross_validate method. The
 reason is that our element data is non-random by default. So we want to perform
 shuffling prior to doing the splits.
- Documentation: link.

from sklearn.model_selection import cross_validate, KFold

Characteristics of the example materials dataset

- Before proceeding further, let us try to tease out some aspects of the dataset.
- Quite clearly, there are correlations between some sets of variables.
- In other words, the input features are **non-orthonormal** with each other.



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Loss functions for regression

- We have thus far focused on the squared error loss $L(y, f(x)) = (y f(x))^2$
- Another common loss function is the absolute error L(y, f(x)) = |y f(x)|
- MSE penalizes outliers with large observed residuals severely, and hence is less robust in data with long-tailed distributions.
- MAE is more robust against outliers.
- Other criteria include the Huber loss:

$$L(y, f(x)) = \begin{cases} (y - f(x))^2 & |y - f(x)| \le \delta \\ 2\delta(y - f(x) - \delta^2) & \text{otherwise} \end{cases}$$

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The End