# Introduction to ML and Regression Statistical Natural Language Processing

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# Why machine learning?

- Majority of the modern computational linguistic tasks and applications are based on machine learning
  - Tokenization
  - Part of speech tagging
  - Parsing
  - ...
  - Speech recognition
  - Named Entity recognition
  - Document classification
  - Question answering
  - Machine translation
  - ...

# Machine learning is ...

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Statistical learning refers to a vast set of tools for understanding data.

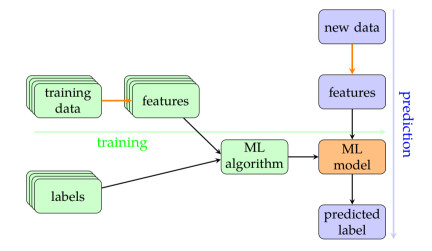
—James et al. (2013)

# Supervised or unsupervised

- Machine learning methods are often divided into two broad categories: *supervised* and *unsupervised*
- Supervised methods rely on labeled (or annotated) data
- Unsupervised methods try to find regularities in the data without any (direct) supervision
- Some methods do not fit any (or fit both):
  - *Semi-supervised* methods use a mixture of both
  - *Reinforcement learning* refers to the methods where supervision is indirect and/or delayed

In this course, we will mostly discuss/use supervised methods.

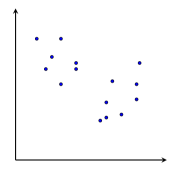
# Supervised learning



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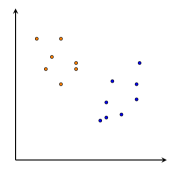
# Unsupervised learning

- In unsupervised learning we do not have any labels
- The aim is discovering some 'latent' structure in the data
- Common examples include
  - Clustering
  - Density estimation
  - Dimensionality reduction
- The methods that do not require (manual) annotation are sometimes called unsupervised



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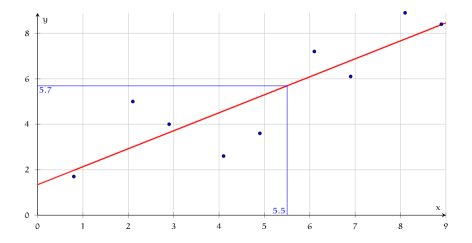
# Supervised learning

two common settings

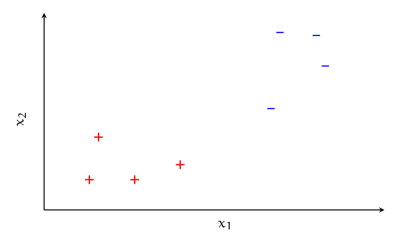
#### A supervised ML method is called

*regression* if the outcome to be predicted is a numeric (continuous) variable *classification* if the outcome to be predicted is a categorical variable

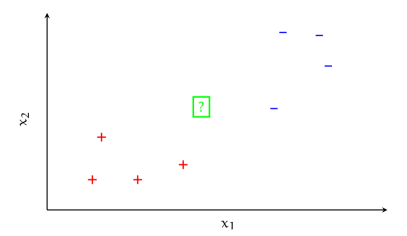
Regression



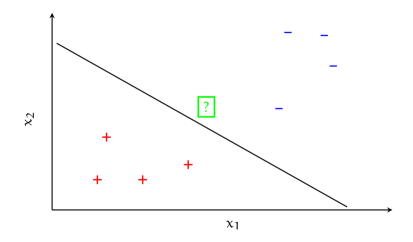
## Classification



## Classification



## Classification



# ML topics we will cover in this course

- (Linear) Regression (today)
- Classification (perceptron, logistic regression, ANNs)
- Evaluating ML methods / algorithms
- Unsupervised learning
- Sequence learning

# Machine learning and statistics

- The methods largely overlap (it was even suggested that both should be collectively called 'data science')
- Aims differ
  - In statistics (used as in experimental sciences) aim is making inferences using the models
  - In machine learning correct predictions are at the focus
- A more diverse set of models/methods are used in ML

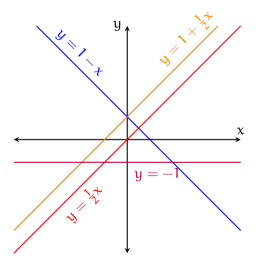
# Machine learning and models

- A machine learning method makes its predictions based on a model
- The models are often parametrized: a set of parameters defines a model
- Learning can be viewed as finding the 'best' model among a family of models (that differ based on their parameters)

## The linear equation: the regression model

y = a + bx

- a (intercept) is where the line crosses the y axis.
- b (slope) is the change in y as x is increased one unit.



# The simple linear model some terminology

#### $y_i = a + bx_i$

- y is the *outcome* (or response, or dependent) variable. The index i represents each unit observation/measurement (sometimes called a 'case')
- x is the *predictor* (or explanatory, or independent) variable
- a is the *intercept* (called *bias* in the NN literature)
- b is the *slope* of the regression line.
- a and b are called *coefficients* or *parameters*
- a + bx is the model's prediction of y ( $\hat{y}$ ), given x

 $y_i = a + bx_i$ 

 $y_i = \alpha + \beta x_i$ 

- Sometimes, Greek letters  $\alpha$  and  $\beta$  are used for intercept and the slope, respectively

 $y_i = \beta_0 + \beta_1 x_i$ 

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- In machine learning it is common to use *w* for all coefficients (sometimes you may see b used instead of w<sub>0</sub>)

 $y_i = \hat{w}_0 + \hat{w}_1 x_i$ 

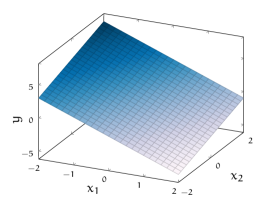
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 $y_i = w x_i$ 

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- Sometimes coefficients wear hats, to emphasize that they are estimates
- Often, we use the vector notation for both input(s) and coefficients:  $w = (w_0, w_1)$  and  $x_i = (1, x_i)$

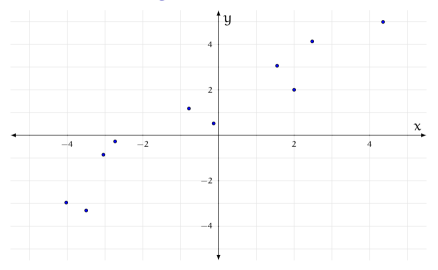
# Regression models with multiple predictors

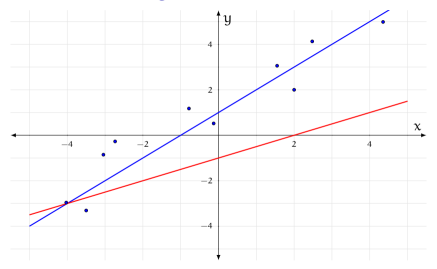
- The equation defines a (hyper)plane
- With 2 predictors:  $y = w_0 + w_1 x_1 + w_2 x_2$
- With more predictors it is more convenient to use the vector notation: y = wx

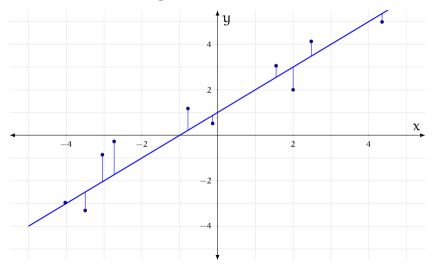


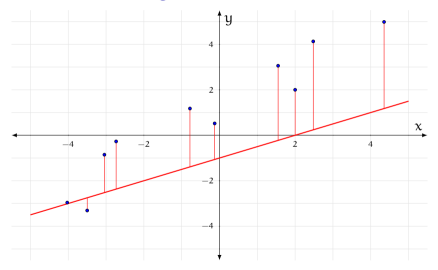
#### Parameter estimation

- In ML, we are interested in finding the best model based on data
- Learning is selecting a model from a family of models that differ in their parameters
- Typically, we seek the parameters that reduce the prediction error on a training set
- Ultimately, we seek models that do not only do well on the training data, but also new, unseen instances



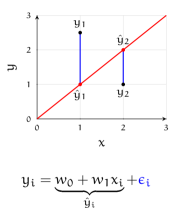






## Estimating regression parameters

- We view learning as a search for the regression equation with least error
- The error terms are also called *residuals*
- We want error to be low for the whole training set: average (or sum) of the error has to be reduced
- Can we minimize the sum of the errors?



## Least-squares regression

• Find  $w_0$  and  $w_1$ , that minimize the sum of the squared errors (SSE)

$$\mathsf{E}(\boldsymbol{w}) = \sum_{i} \varepsilon_{i}^{2} = \sum_{i} (y_{i} - \hat{y}_{i})^{2} = \sum_{i} (y_{i} - (w_{0} + w_{1}x_{i}))^{2}$$

• We can minimize E(w) analytically

$$w_1 = \frac{\sigma_{xy}}{\sigma_x^2} = r \frac{sd_y}{sd_x} \qquad \qquad w_0 = \bar{y} - w_1 \bar{x}$$

# Short digression: minimizing functions

In least squares regression, we want to find  $w_0$  and  $w_1$  values that minimize

$$E(\boldsymbol{w}) = \sum_{i} (y_i - (w_0 + w_1 x_i))^2$$

- Note that E(w) is a *quadratic* function of  $w = (w_0, w_1)$
- As a result, E(w) is *convex* and have a single extreme value
  - there is a unique solution for our minimization problem
- In case of least squares regression, there is an analytic solution
- Even if we do not have an analytic solution, if the error function is convex, a search procedure like *gradient descent* can still find the *global minimum*

## What is special about least-squares?

- Minimizing MSE (or  $SS_R)$  is equivalent to MLE estimate under the assumption  $\varepsilon\sim \mathcal{N}(0,\sigma^2)$
- Working with 'minus log likelihood' is more convenient

$$\mathsf{E}(w) = -\log \mathcal{L}(w) = -\log \prod_{i} \frac{e^{-\frac{(\mathsf{y}_{i} - \hat{\mathsf{y}}_{i})^{2}}{2\sigma^{2}}}}{\sigma\sqrt{2\pi}}$$

$$\hat{\boldsymbol{w}} = \operatorname*{arg\,min}_{\boldsymbol{w}} (-\log \mathcal{L}(\boldsymbol{w})) = \operatorname*{arg\,min}_{\boldsymbol{w}} \sum_{i} (y_{i} - \hat{y}_{i})^{2}$$

- There are other error functions, e.g., absolute value of the errors, that can be used (and used in practice)
- One can also estimate regression parameters using Bayesian estimation

# Regression with multiple predictors

$$y_{i} = \underbrace{w_{0} + w_{1}x_{i,1} + w_{2}x_{i,2} + \ldots + w_{k}x_{i,k}}_{\hat{y}} + \epsilon_{i} = wx_{i} + \epsilon_{i}$$

 $w_0$  is the intercept (as before).

- $w_{1..k}$  are the coefficients of the respective predictors.
  - $\epsilon$  is the error term (residual).
  - using the vector notation the equation becomes:

$$y_i = wx_i + \epsilon_i$$

where  $\boldsymbol{w} = (w_0, w_1, \dots, w_k)$  and  $\boldsymbol{x_i} = \left(1, x_{i,1}, \dots, x_{i,k}\right)$ 

It is a generalization of simple regression with some additional power and complexity.

### Evaluating machine learning systems

- Any (machine learning) system needs a way to measure its success
- For measuring success (or failure) in a machine learning system we need quantitative measures
- Remember that we need to measure the success outside the training data

#### Measuring success in Regression

• *Root-mean-square error* (RMSE)

$$\text{RMSE} = \sqrt{\frac{1}{n}\sum_{i}^{n}(y_{i} - \hat{y}_{i})^{2}}$$

measures average error in the units compatible with the outcome variable.

• Another well-known measure is the *coefficient of determination* 

$$R^{2} = \frac{\sum_{i}^{n} (\hat{y}_{i} - \bar{y})^{2}}{\sum_{i}^{n} (y_{i} - \bar{y})^{2}} = 1 - \left(\frac{RMSE}{\sigma_{y}}\right)^{2}$$

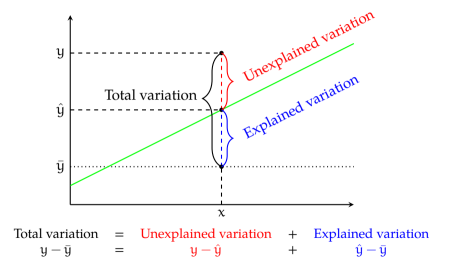
# Assessing the model fit: $R^2$

We can express the variation explained by a regression model as:

$$\frac{\text{Explained variation}}{\text{Total variation}} = \frac{\sum_{i=1}^{n} (\hat{y}_{i} - \bar{y})^{2}}{\sum_{i=1}^{n} (y_{i} - \bar{y})^{2}}$$

- In simple regression, it is the square of the correlation coefficient between the outcome and the predictor
- The range of  $\mathbb{R}^2$  is [0, 1]
- +  $100 \times R^2$  is interpreted as 'the percentage of variance explained by the model'
- $R^2$  shows how well the model fits to the data: closer the data points to the regression line, higher the value of  $R^2$

#### **Explained variation**



# Dealing with non-linearity

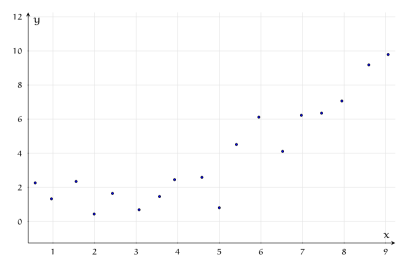
- Least-squares estimation works because the regression equation is linear with respect to parameters *w* (error function is quadratic)
- Introducing non-linear combinations of inputs does not affect the estimation procedure. The following are still linear models

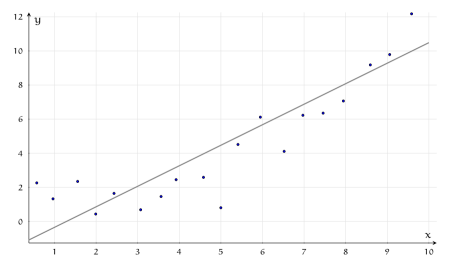
$$y = w_0 + w_1 x^2 + \epsilon$$
  

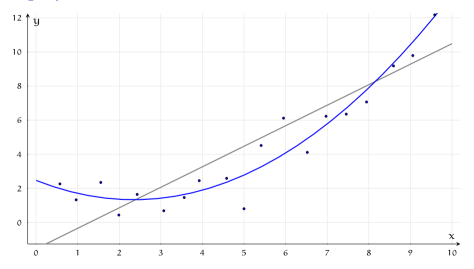
$$y = w_0 + w_1 \log(x) + \epsilon$$
  

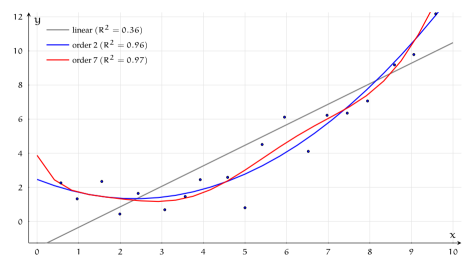
$$y = w_0 + w_1 x_1 + w_2 x_2 + w_3 x_1 x_2 + \epsilon$$

- In general, we can replace input x by a function of the input(s)  $\Phi(x)$ .  $\Phi()$  is called a *basis function*
- Basis functions allow linear models to model non-linear relations by *transforming* the input variables







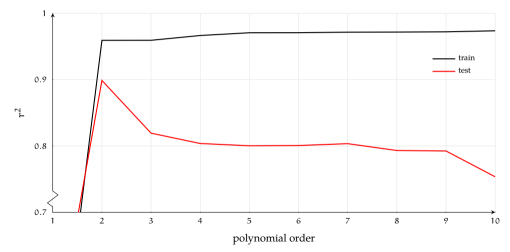


## Overfitting

- *Overfitting* is an important problem in ML, happens when the model learns peculiarities/noise in the training data
- An overfitted model will perform well on training data, but worse on new/unseen data
- Typically 'more complex' models are more likely to overfit

Overfitting

demonstration through polynomial regression



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### Preventing overfitting

- A straightforward approach is to chose a simpler model (family), e.g., by reducing the number of predictors
- More training data helps: it is less likely to overfit if number of training instances are (much) larger than the paramters
- There are other methods (one is coming on the next slide)
- We will return to this topic frequently during later lectures

#### Regularized parameter estimation

- *Regularization* is a general method for avoiding overfitting
- The idea is to constrain the parameter values in addition to minimizing the training error
- For example, the regression estimation becomes:

$$\hat{\boldsymbol{w}} = \operatorname*{arg\,min}_{\boldsymbol{w}} \sum_{i} (y_{i} - \hat{y}_{i})^{2}$$

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$$\hat{\boldsymbol{w}} = \operatorname*{arg\,min}_{\boldsymbol{w}} \sum_{i} (y_{i} - \hat{y}_{i})^{2} + \lambda \sum_{j=1}^{k} w_{j}^{2}$$

- The new part is called the regularization term,
- $\lambda$  is a *hyperparameter* that determines the strength of the regularization
- In effect, we are preferring small values for the coefficients
- Note that we do not include w<sub>0</sub> in the regularization term

### L2 regularization

The form of regularization, where we minimize the regularized cost function,

 $E(\boldsymbol{w}) + \lambda \|\boldsymbol{w}\|_2$ 

is called L2 regularization.

- Note that we are minimizing the L2-norm of the weight vector
- In statistic literature L2-regularized regression is called *ridge regression*
- The method is general: it can be applied to other ML methods as well
- The choice of  $\lambda$  is important
- Note that the scale of the input also becomes important

### L1 regularization

In L1 regularization we minimize

$$E(\boldsymbol{w}) + \lambda \sum_{j=1}^{k} |w_j|$$

- The additional term is the L1-norm of the weight vector (excluding  $w_0$ )
- In statistics literature the L1-regularized regression is called lasso
- The main difference from L2 regularization is that L1 regularization forces some values to be 0 the resulting model is said to be 'sparse'

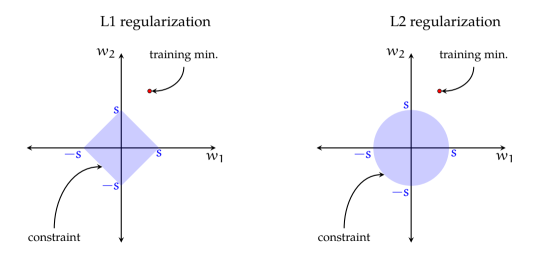
### Regularization as constrained optimization

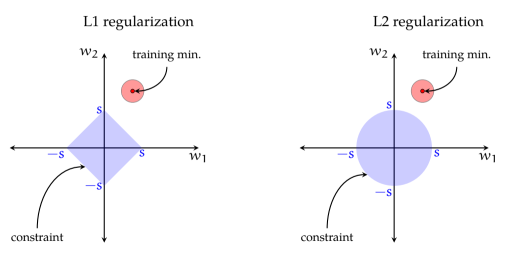
L1 and L2 regularization can be viewed as minimization with constraints L2 regularization

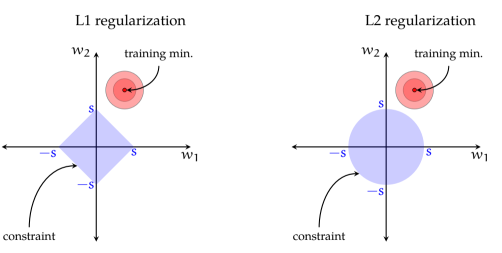
Minimize E(w) with constraint ||w|| < s

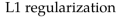
L1 regularization

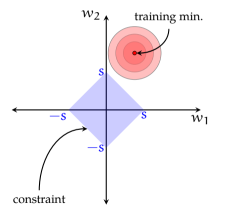
Minimize E(w) with constraint  $||w||_1 < s$ 

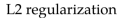


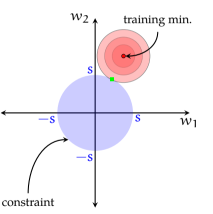


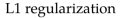


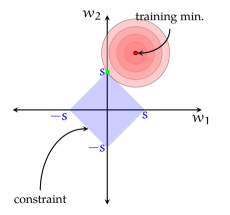


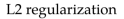


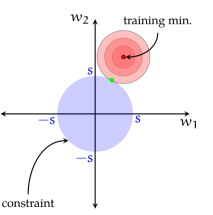












#### Regularization: some remarks

- Regularization prevents overfitting
- The *hyperparameter*  $\lambda$  needs to be determined
  - best value is found typically using a grid search, or a random search
  - it is tuned on an additional partition of the data, development set
  - development set cannot overlap with training or test set
- The regularization terms can be interpreted as *priors* in a Bayesian setting
- Particularly, L2 regularization is equivalent to a normal prior with zero mean

#### Gradient descent for parameter estimation

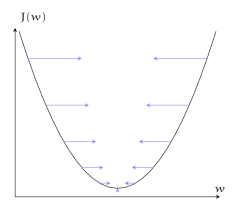
- In many ML problems, we do not have a closed form solution for finding the minimum of the error function
- In these cases, we use a search strategy
- *Gradient descent* is a search method for finding a minimum of a (error) function
- The general idea is to approach a minimum of the error function in small steps

$$\boldsymbol{w} \leftarrow \boldsymbol{w} - \eta \nabla J(\boldsymbol{w})$$

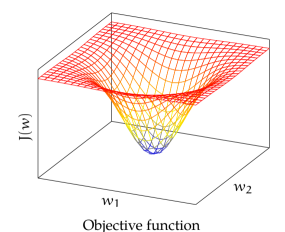
- $\nabla J\,$  is the gradient of the loss function, it points to the direction of the maximum increase
  - $\eta~$  is the learning rate

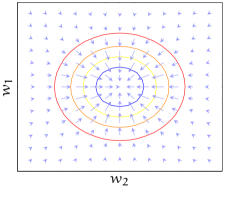
## Gradient descent with single parameter

- For a single parameter, gradient is a one-dimensional vector
- The direction of gradient is towards the maximum increase
- We take steps proportional to  $-\nabla J(w)$
- Steeper the curve, the larger the parameter update



## Gradient descent with single parameter





Negative gradients

# Categorical predictors

- Categorical predictors are represented as multiple binary coded input variables
- For a binary predictor, we use a single binary input. For example, (1 for one of the values, and 0 for the other)

$$\mathbf{x} = \begin{cases} \mathbf{0} & \text{ for male} \\ \mathbf{1} & \text{ for female} \end{cases}$$

• For a categorical predictor with k values, we use one-hot encoding (other coding schemes are possible)

$$\mathbf{x} = \begin{cases} (0,0,1) & \text{neutral} \\ (0,1,0) & \text{negative} \\ (1,0,0) & \text{positive} \end{cases}$$

# Summary

#### What to remember:

- Supervised vs. unsupervised learning
- Regression vs. classification
- Linear regression equation
- Least-square estimate

#### Next:

Wed classification

Fri first lab session

Mon classification

- MSE, R<sup>2</sup>
- non-linearity & basis functions
- L1 & L2 regularization (lasso and ridge)

# Additional reading, references, credits

- Hastie, Tibshirani, and Friedman (2009) discuss introductory bits in chapter 1, and regression on chapter 3 (sections 3.2 and 3.4 are most relevant to this lecture)
- Jurafsky and Martin (2009) has a short section (6.6.1) on regression
- You can also consult any machine learning book (including the ones listed below)



Barber, David (2012). Bayesian Reasoning and Machine Learning. Cambridge University Press. ISBN: 9780521518147.



i.

Hastie, Trevor, Robert Tibshirani, and Jerome Friedman (2009). The Elements of Statistical Learning: Data Mining, Inference, and Prediction. Second. Springer series in statistics. Springer-Verlag New York. ISBN: 9780387848587. URL: http://web.stanford.edu/-hastie/ElemStatLearn/.

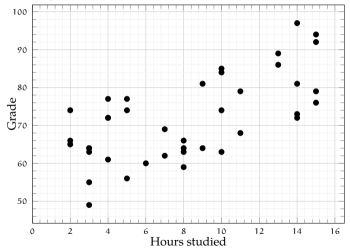


Jurafsky, Daniel and James H. Martin (2009). Speech and Language Processing: An Introduction to Natural Language Processing, Computational Linguistics, and Speech Recognition. second. Pearson Prentice Hall. 158N: 978-0-13-504196-3.

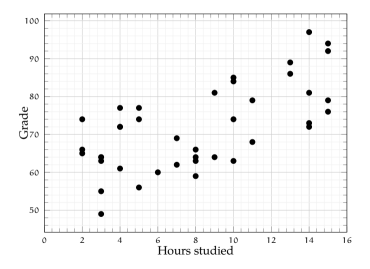
Mitchell, Thomas (1997). Machine Learning. 1st. McGraw Hill Higher Education. ISBN: 0071154671,0070428077,9780071154673,9780070428072.

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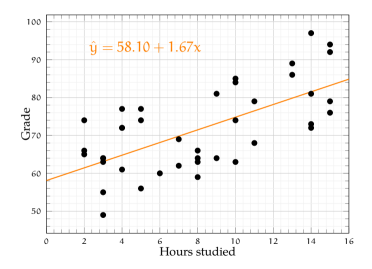
#### Draw a regression line over the plot



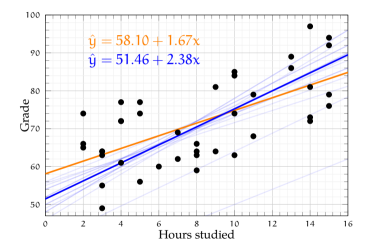
#### The regression line



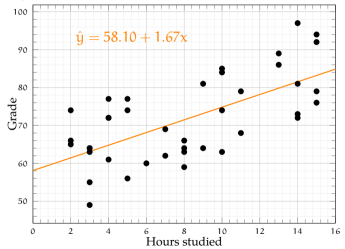
The regression line



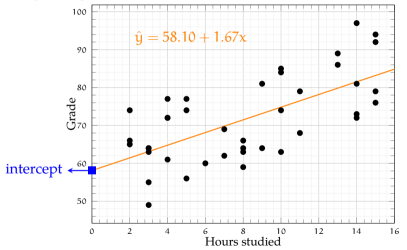
#### Your estimates



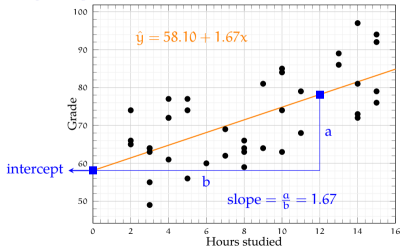
How to calculate the regression parameters



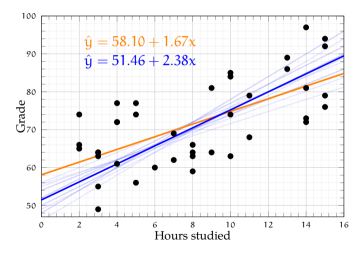
How to calculate the regression parameters



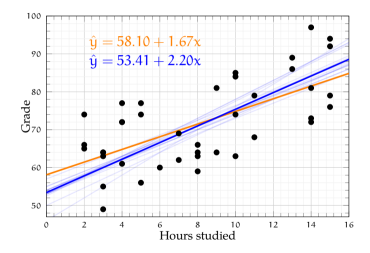
How to calculate the regression parameters



Your estimates (some removed)



Estimates from 2020



Regression estimates 2019–2021

