IB031 Úvod do strojového učení

Tomáš Brázdil

Course Info

Resources:

- Lectures & tutorials (the **main** source)
- Many books, few perfect for introductory level
 One relatively good, especially the first part:
 A. Géron. Hands-On Machine Learning with Scikit-Learn, Keras, and TensorFlow: Concepts, Tools, and Techniques to Build Intelligent Systems. O'Reilly Media; 3rd edition, 2022
- (Almost) infinitely many online courses, tutorials, materials, etc.

Evaluation

The evaluation is composed of three parts:

- Mid-term exam: Written exam from the material of the first half of the semester.
- End-term exam: The "big" one containing everything from the semester (with possibly more stress in the second half).
- Projects: During tutorials, you will work on larger projects (in pairs).

Each part contributes the following number of points:

▶ Mid-term exam: 25

► End-term exam: 50

▶ Project: 25

To pass, you need to obtain at least 60 points.

Distinguishing Properties of the Course

- Introductory, prerequisites are held to a minimum
- Formal and precise: Be prepared for a complete and "mathematical" description of presented methods.

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I assume that you have basic knowledge of

- Elementary understanding of mathematical notation (operations on sets, logic, etc.)
- Linear algebra: Vectors in \mathbb{R}^n , operations on vectors (including the dot product). Geometric interpretation!
- Calculus: Functions of multiple real variables, partial derivatives, basic differential calculus.
- Probability: Notion of probability distribution, random variables/vectors, expectation.

What Is Machine Learning?

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5

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Here is a slightly more general definition:

Arthur Samuel, 1959

Machine learning is the field of study that allows computers to learn without being explicitly programmed.

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Machine learning is the field of study that allows computers to learn without being explicitly programmed.

And a more engineering-oriented one:

Tom Mitchell, 1997

A computer program is said to learn from experience E concerning some task T and some performance measure P if its performance on T, as measured by P, improves with experience E.

Example

In the context of spam filtering:

- The task T is to flag spam in new emails.
- ► The experience *E* is represented by a set of emails labeled either spam or ham by hand (the training data).
- ▶ The performance measure *P* could be the accuracy, which is the ratio of the number of correctly classified emails and all emails.

There are many more performance measures; we will study the basic ones later.

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In the context of housing price prediction:

- ► The task *T* is to predict prices of new houses based on their basic parameters (size, number of bathrooms, etc.)
- ► The experience E is represented by information about existing houses.
- ► The performance measure *P* could be, e.g., an absolute difference between the predicted and real price.

Examples (cont.)

In the context of game playing:

- ▶ The task *T* is to play chess.
- ► The experience *E* is represented by a series of self-plays where the computer plays against itself.
- ► The performance measure *P* is winning/losing the game. Here, the trick is to spread the delayed and limited feedback about the result of the game throughout the individual decisions in the game.

Examples (cont.)

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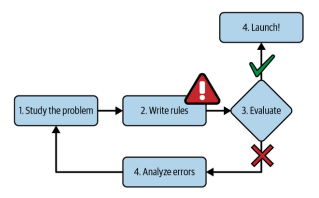
In the context of customer behavior:

- ► The task *T* is to group customers with similar shopping habits in an e-shop.
- ► The experience *E* consists of lists of items individual customers bought in the shop.
- The performance measure P? Measure how "nicely" the customers are grouped. (whether people with similar habits, as seen by humans, fall into the same group).

Comparison of Programming and Learning

How to code the spam filter?

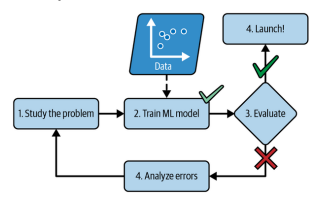
- Examine what spam mails typically contain: Specific words ("Viagra"), sender's address, etc.
- ▶ Write down a rule-based system that detects specific features.
- ► Test the program on new emails and (most probably) go back to look for more spam features.



Comparison of Programming and Learning

The machine learning way:

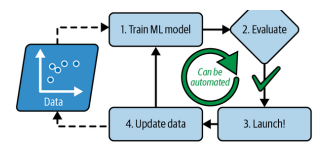
- Study the problem and collect lots of emails, labeling them spam or ham.
- Train a machine learning model that reads an email and decides whether it's spam or ham.
- Test the model and (most probably) go back to collect more data and adjust the model.



ML Solutions are Adaptive

Spam filter: Authors of spam might and will adapt to your spam filter (possibly change the wording to pass through).

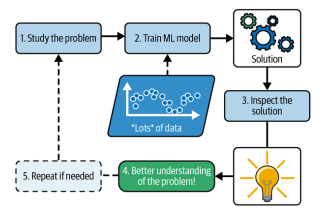
ML systems can be adjusted to new situations by retraining on new data (unless the data becomes ugly).



ML for Human Understanding

Spam filter: A trained system can be inspected for notorious spam features.

Some models allow direct inspection, such as decision trees or linear/logistic regression models.



Usage of Machine Learning

Machine learning suits various applications, especially where traditional methods fall short. Here are some areas where it excels:

- Solving complex problems where fine-tuning and rule-based solutions are inadequate.
- Tackling complex issues that resist traditional problem-solving approaches.
- Adapting to fluctuating environments through retraining on new data.
- Gaining insights from large and complex datasets.

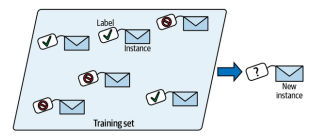
In summary, machine learning offers innovative solutions and adaptability for today's complex and ever-changing problems, (sometimes) providing insights beyond the reach of traditional approaches.

Types of Learning

There are main categories based on information available during the training:

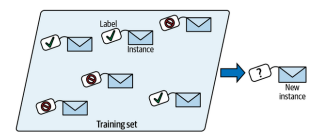
- Supervised learning
- Unsupervised learning
- Semi-supervised learning
- Self-supervised learning
- Reinforcement learning

Supervised Learning



Labels are available for all input data.

Supervised Learning

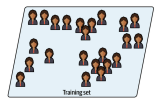


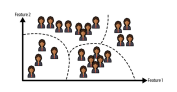
Labels are available for all input data.

Typical supervised learning tasks are

- Classification where the aim is to classify inputs into (typically few) classes
 - (e.g., the spam filter where the classes are spam/ham)
- Regression where a numerical value is output for a given input (e.g., housing prices)

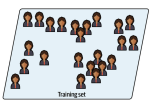
Unsupervised Learning

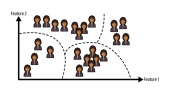




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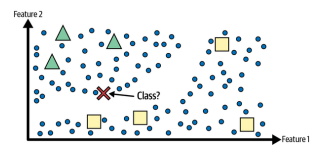
Typical unsupervised learning tasks are

 Clustering where inputs are grouped according to their features

(e.g., clients of a bank grouped according to their age, wealth, etc.)

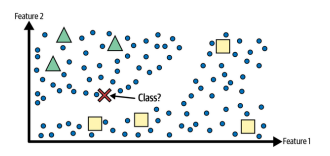
- Association where interesting relations and rules are discovered among the features of inputs
 - (e.g., market basket mining where associations between various types of goods are being learned from the behavior of customers)
- Dimensionality reduction reduce high-dimensional data to few dimensions (e.g., images to few image features)

Semi-Supervised Learning



Labels for some data.

Semi-Supervised Learning

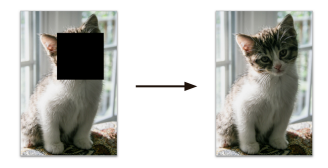


Labels for some data.

For example, Medical data, where elaborate diagnosis is available only for some patients.

Combines supervised and unsupervised learning: e.g., clusters all data and labels the unlabeled inputs with the most common labels in their clusters.

Self-Supervised Learning

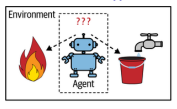


Generate labels from (unlabeled) inputs.

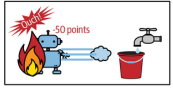
The goal is to learn typical features of the data.

It can be later modified to generate images, classify, etc.

Reinforcement Learning



- Observe
- 2 Select action using policy



- 3 Action!
- 4 Get reward or penalty



- 5 Update policy (learning step)
- 6 Iterate until an optimal policy is found

Learn from performing actions and getting feedback from environment.

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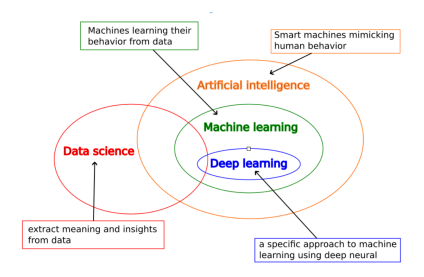
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- ▶ Various "table" data processing in finance, management, etc.
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 - Essential but not fancy
- ► Game playing: More fancy than useful, learning models beating humans in several difficult games.

ML in Context



Supervised Learning

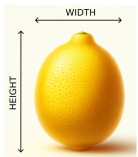
Example - Fruit Recognition

The goal: Create an automatic system for fruit recognition, concretely apple, lemon, and mandarin.

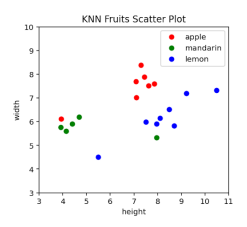
Inputs: Measures of *height* and *width* of each fruit.

Suppose we have a dataset of dimensions of several fruits labeled with the correct class.





Data



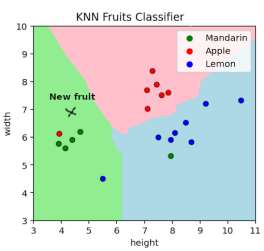
Use similarity to solve the problem.

	height	width	fruit
0	3.91	5.76	Mandarin
1	7.09	7.69	Apple
2	10.48	7.32	Lemon
3	9.21	7.20	Lemon
4	7.95	5.90	Lemon
5	7.62	7.51	Apple
6	7.95	5.32	Mandarin
7	4.69	6.19	Mandarin
8	7.50	5.99	Lemon
9	7.11	7.02	Apple
10	4.15	5.60	Mandarin
11	7.29	8.38	Apple
12	8.49	6.52	Lemon
13	7.44	7.89	Apple
14	7.86	7.60	Apple
15	3.93	6.12	Apple
16	4.40	5.90	Mandarin
17	5.50	4.50	Lemon
18	8.10	6.15	Lemon
19	8.69	5.82	Lemon

KNN Classification

Given a new fruit. What is it?

Find five closest examples



Where is the machine learning?

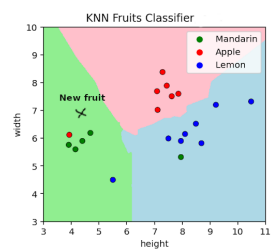
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Among the five closest:

- ightharpoonup M = 4 mandarins
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KNN Classification

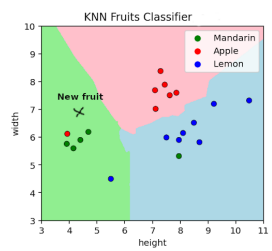
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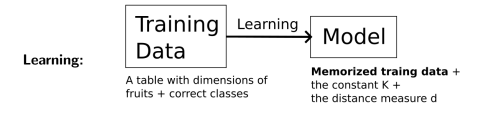
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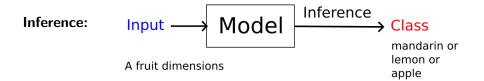
It is a mandarin!



Where is the machine learning?

Learning in Fruit Classification with KNN





Fruit Classification Algorithm

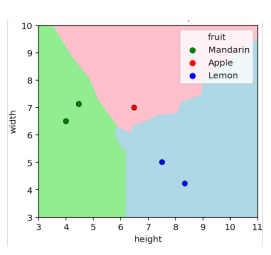
```
Input: A fruit F with dimensions height, width
Output: mandarin, lemon, apple
 1: Find K examples \{E_1, \ldots, E_K\} in the dataset whose
    dimensions are closest to the dimensions of the fruit F
 2: Count the number of examples of each class in \{E_1, \ldots, E_K\}
         M mandarins in \{E_1,\ldots,E_K\}
         L lemons in \{E_1,\ldots,E_K\}
         A apples in \{E_1,\ldots,E_K\}
 3: if M \ge L and M \ge A then return mandarin
 4: else if L \ge A then return lemon
 5: else return apple
 6: end if
```

Does it work?

Testing the Model for Fruit Classification

Consider a test set of new instances (K = 5, d is Euclidean):

height	width	fruit	
4.0	6.5	Mandarin	
4.47	7.13	Mandarin	
6.49	7.0	Apple	
7.51	5.01	Lemon	
8.34	4.23	Lemon	



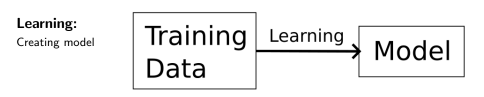
Perfect classification of new data! Just deploy and sell!!

K Nearest Neighbors

...on ideal data

Learning and Inference

Two crucial components of machine learning are the following:





Training Data

Assume table training data, i.e., of the form

Formally, we define training dataset

$$\mathcal{T} = \{ (\vec{x}_k, c_k) \mid k = 1, \dots, p \}$$

Here each $\vec{x}_k \in \mathbb{R}^n$ is an input vector and $c_k \in C$ is the correct class.

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$$\dots\}$$

KNN: Learning

Consider the training set:

$$\mathcal{T} = \{ (\vec{x}_k, c_k) \mid k = 1, \dots, p \}$$

and memorize it exactly as it is.

Store in a table.

Possibly use a clever representation allowing fast computation of nearest neighbors such as KDTrees (out of the scope of this lecture).

Also,

- ▶ determine the number of neighbors $K \in \mathbb{N}$,
- and the distance measure d.

Inference in KNN

Assume a KNN "trained" by memorizing

 $\mathcal{T} = \{(\vec{x}_k, c_k) \in \mathbb{R}^n \times C \mid k = 1, ..., p\}$, a constant $K \in \mathbb{N}$ and a distance measure d.

For d, consider Euclidean distance, but different norms may also be used to define different distance measures.

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Input: A vector $\vec{z} = (z_1, \dots, z_n) \in \mathbb{R}^n$

Output: A class from C

1: Find K indices of examples $X = \{i_1, \ldots, i_K\} \subseteq \{1, \ldots, p\}$ with minimum distance to \vec{z} , i.e., satisfying

$$\max \left\{ d(\vec{z}, \vec{x}_{\ell}) \mid \ell \in X \right\} \leq \min \left\{ d(\vec{z}, \vec{x}_{\ell}) \mid \ell \in \{1, \dots, p\} \setminus X \right\}$$

- 2: For every $c \in \mathcal{C}$ count the number #c of elements ℓ in X such that $c_\ell = c$
- 3: Return some

$$c_{max} \in \underset{c \in C}{\operatorname{arg max}} \# c$$

A class $c_{max} \in C$ which maximizes #c.

The resulting model

What exactly constitutes the model? The model consists of

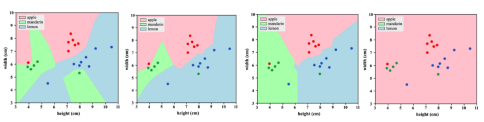
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- ► The *hyperparameters* set "from the outside": In this case, the number of neighbors *K* and the distance measure *d*.

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Note that different settings of K lead to different classifiers (for the same d):



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 - Deal with issues in the data
 - Data almost always comes in weird formats, with inconsistencies, missing values, wrong values, etc.
 - Data rarely have the ideal form for a given learning model.

We need to ingest, validate, and preprocess the data.

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- Deal with the wrong model by testing and validation in as realistic conditions as possible.
- Deal with deployment real-world application issues involving, e.g., implementation in embedded devices with limited resources.

Models Considered in This Course

Throughout this course, we will meet the following models:

- KNN (already did)
- Decision trees
- ► (Naive) Bayes classifier
- Clustering: K-means and hierarchical
- Linear and logistic regression
- Support Vector Machines (SVM)
- Kernel linear models
- Neural networks (light intro to feed-forward networks)
- Ensemble methods + random forests
- (maybe some reinforcement learning)

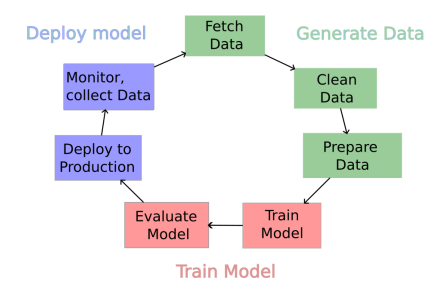
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... but first, let us see the whole machine learning pipeline.

Machine Learning Pipeline



Always start with

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- ▶ Integrate data from various sources. A serious diagnostic system must be trained/tested on data from many hospitals. You must blend the data from various sources (different formats, etc.).

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Data Separation

At this point, you should randomize the ordering of the data and select a test set to be used in model evaluation!

The test data are supposed to simulate the actual conditions, i.e., they should be "unseen".

For simple "toy" machine learning projects, you may fetch prepared datasets from various databases on the internet.

The data should be stored in an identified location and versioned. You will probably keep adding data and training models on the ever-changing datasets. You have to be able to keep track of the changes and map training data to particular models.

Tools such as ML Flow or Weights & Biases might be helpful.

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Data Exploration

Compute basic statistics to identify missing values, outliers, etc.

Clean Data

The cleaning usually comprises the following steps:

- Fix or remove incorrect or corrupted values.
- ▶ Identify outliers and decide what to do with them. Outliers may harm some training methods and are not "representative". However, sometimes, they naturally belong to the dataset, and expert insight is needed.
- Fix formatting.
 For example, the Date may be expressed in many ways, and a simple Yes/No answer.
- Resolve missing values (by either removing the whole examples or imputing)
 - Many methods have been developed for missing values imputation. It is a susceptible issue because new values may strongly bias the model.
- Remove duplicates.

The above steps often affect the training and need expertise in the application domain.

Later in this course, we will discuss techniques for data cleaning.

ID	Age	Income	Gender	Customer_Satisfaction
1	38	46641.356413713	nan	Unsatisfied
2	42	49129.0615585107	female	Neutral
3	18	119965.049731014	Male	nan
4	18	66828.0762224329	nan	very unsatisfied
5	58	57422.2721106762	female	very unsatisfied
6	28	59502.8174855665	Other	Satisfied
7	18	42659.6675768587	Other	Neutral
8	18	54019.1173206374	Other	Satisfied
9	40	25429.1604541137	female	Unsatisfied
10	21	15595.5862129548	Other	Satisfied
11	18	58094.2328460069	Other	very unsatisfied
12	18	39097.3278583155	female	Very Satisfied
13	30		Other	Satisfied
14	50	30617.3914472273	Female	Very Satisfied
15	18		nan	Neutral
16	34	39902.4430953214	male	nan
17	49	68381.6997683133	Female	Very Satisfied
18	33	44796.0962271524	Other	Very Satisfied
19	47	39218.9560738814	Female	very unsatisfied
20		14544.9226784447	Other	Satisfied

Prepare Data

Unlike cleaning, which is application-dependent, data preparation/transformation is model-dependent. This usually subsumes:

- Scaling: Settings values of inputs to a similar range.
 Some models, especially those utilizing distance, are sensitive to large differences between input sizes.
- ► Encoding: Encode non-numeric data using real-valued vectors.

 Many models, especially those based on geometry, work only with numeric data. Non-numeric data such as Yes/No, Short/Medium/Long must be encoded appropriately.
- ▶ **Binning or Discretization** Convert continuous features into discrete bins to capture patterns in ranges.

Comment: Sometimes **Normalization**, that is changing the distribution of inputs to resemble the normal distribution, is mentioned. However, this step is typically not essential for machine learning itself. However, it is important to use statistical inference to test the significance of learned parameters.

Prepare Data

Feature selection Throw out input features that are too "similar" to other features.

For example, if the temperature is measured both in Celsius and in Kelvin, keep one of them. The relationship can, of course, be a more complex (non-linear) correlation.

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▶ **Dimensionality reduction** Transforming data from \mathbb{R}^n to \mathbb{R}^m where m << n.

Growing dimension means growing difficulty of training for all models. Some models cease to work for high-dimensional data. The reduction typically searches for a few important characteristic features of inputs.

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► **Feature aggregation** Introducing new features using operations on the original ones.

We will see kernel transformations later in this course, allowing simple models to solve complex problems.

Train Model

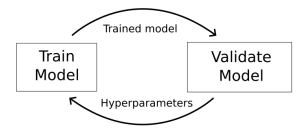
Now the dataset has been cleaned; we may train a model.

Train Model

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Before training, we should split the dataset into

- training dataset on which the model will learn
- validation dataset on which we fine-tune hyperparameters



The resulting model is obtained after several iterations of the above process.

Evaluate Model

Here, we use the test set that we separated during data fetching.

In some cases, a brand new test set can be generated.

patients are examined regularly, creating new records continuously.

In some cases, it is tough to obtain new data.

For example, new expensive and difficult measurements are needed to obtain new data.

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Critical issue: Make sure that you are truly testing exactly the whole inference process.

Often, just a model is tested, and the testing and production inference engines are separated. This leads to truly nasty errors in the production!

We will discuss various generic metrics helpful in measuring the quality of the resulting model.

Deployment of machine learning models is a complex question, application dependent.

The recently emerging area of MLOps is concerned with the engineering side of the model deployment.

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The recently emerging area of MLOps is concerned with the engineering side of the model deployment.

From the technical point of view, the typical issues solved by ML Ops teams are

- how to extract/process data in real-time
- how much storage is required
- how to store/collect model (and data) artifacts/predictions
- how to set up APIs, tools, and software environments
- What the period of predictions (instantaneous or batch predictions) should be
- how to set up hardware requirements (or cloud requirements for on-cloud environments) by the computational resources required
- how to set up a pipeline for continuous training and parameter tuning

From the user's point of view:

- How to get a sensible and valuable user output?
 - ► Al researchers will be satisfied with tons of running text in terminals.
 - "Normal" people need a graphical interface with understandable output.
 - Experts working in other domains typically demand speed and clarity at the extreme.

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 - Experts working in other domains typically demand speed and clarity at the extreme.
- ▶ How do you persuade users that the AI is working for them?
 - Especially if safety is at stake, you need to have outstanding arguments and explanations ready for end-users
 - In many areas, the devices need to be certified (medicine, automotive) for ML-based systems.

This complex subject will be only touched on in this course.

Monitor, collect Data

Deployed machine learning models must be constantly monitored.

Because of the influx of new data, ML models work in highly dynamic environments.

For example, an image-processing medical diagnostic model suddenly misdiagnosed a patient because a nurse marked the sample with a marker pen.

Every customer has a different infrastructure and may produce data slightly differently.

Data for retraining and improvement should be stored.

Also, many areas allow the *active learning* where users provide feedback for (continuous) retraining of the models.

Data

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After a few days, you have trained a model that predicts numbers resembling the ones in the table.

You contact the medical researcher and discuss the results.

Researcher: So, you got the data for all the patients?

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Researcher: Well, first, there is field 5, the variable we want to predict. It's common knowledge among people who analyze this type of data that results are better if you work with the log of the values, but I didn't discover this until later. Was it mentioned to you?

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Data Miner: No.

Researcher: But surely you heard about what happened to field 4? It's supposed to be measured on a scale from 1 to 10, with 0 indicating a missing value, but because of a data entry error, all 10's were changed into 0's. Unfortunately, since some of the patients have missing values for this field, it's impossible to say whether a 0 in this field is a real 0 or a 10. Quite a few of the records have that problem.

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Data Miner: Interesting. Were there any other problems? **Researcher:** Yes, fields 2 and 3 are basically the same, but I assume that you probably noticed that.

Data Miner: Yes, but these fields were only weak predictors of field 5.

Researcher: Anyway, given all those problems, I'm surprised you were able to accomplish anything.

Data Miner: True, but my results are really quite good. Field 1 is a very strong predictor of field 5. I'm surprised that this wasn't noticed before.

Researcher: What? Field 1 is just an identification number. **Data Miner:** Nonetheless, my results speak for themselves.

Researcher: Oh, no! I just remembered. We assigned ID numbers after we sorted the records based on field 5. There is a strong

connection, but it isn't very sensible. Sorry.

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connection, but it isn't very sensible. Sorry.

OK, what's the point?

You have to

Understand the task you want to solve and the data!

Data Objects

Data objects represent entities we work with (e.g., classify them).

For example, in cancer prediction, the data objects are patients. In fruit classification, the data objects are individual fruits.

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For example, in cancer prediction, the data objects are patients. In fruit classification, the data objects are individual fruits.

Data objects are described by attributes (or features or variables).

For example, the age, weight, genetic profile, and other patient characteristics. Or the width and height of a fruit.

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So, the following names are usually used as synonyms:

- Attributes used mostly by database and data mining experts.
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One may make some distinctions

- ► Attributes represent information about the object without any additional assumptions.
- Features assume that their values are somewhat characteristic of the object.
- Variables assume that there is some process behind them (typically a random process in the case of statistics).

Data Types - Categorical Attributes

Categorical attributes (nominal attributes) are symbols or names of things.

- Each value represents some kind of category, code, or state.
- ► Values are not ordered and should not be used quantitatively (in computer science, the values are known as enumerations).

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Examples:

```
\begin{split} &\mathsf{hair\_color} \in \{\mathsf{black}, \mathsf{brown}, \mathsf{blond}, \mathsf{red}, \mathsf{auburn}, \mathsf{gray}, \mathsf{white}\} \\ &\mathsf{marital\_status} \in \{\mathsf{single}, \mathsf{married}, \mathsf{divorced}, \mathsf{widowed}\} \\ &\mathsf{customer\_ID} \in \{0, 1, 2, \ldots\} \end{split}
```

Even though the last one is usually expressed using numbers, it should not be used quantitatively.

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Binary attributes are categorical attributes with only two values.

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Ordinal attribute is an attribute with values that have a meaningful order or ranking among them.

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drink\_size \in \{small, medium, large\}

grades \in \{A, B, C, D, E, F\}
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It can also be obtained by discretizing numeric quantities into series of intervals.

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Categorical and ordinal attributes are called *qualitative* attributes.

Next, we look at numeric, i.e., quantitative attributes.

Data Types - Numeric Attributes

Numeric attributes are quantities represented by numbers.

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Distinguish two types: Interval-scale and ratio-scale.

	INTERVAL SCALE	RATIO SCALE	
Measurement	Equal intervals between	Equal intervals with	
interval	consecutive points.	the presence of a true zero.	
Absolute	Lacks a true zero point.	Possesses a true	
zero	Lacks a true zero point.	zero point.	
Statistical	Limited to addition	Allows for meaningful	
analysis	and subtraction	multiplication and division.	
Meaningful	Ratios are not meaningful	Ratios are meaningful	
ratios	due to the lack of zero.	due to the presence of zero.	
Examples	IQ scores, Celsius temperature, NPS data, etc.	Height, weight, income, etc.	

Discrete vs Continuous Attributes

Often, two kinds of numeric attributes are distinguished:

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A finite or countably infinite range of values, i.e., integers may represent the values.

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► Continuous

An uncountably infinite range of values, typically an interval. There are several more or less formal definitions of continuous attributes in the literature. For example:

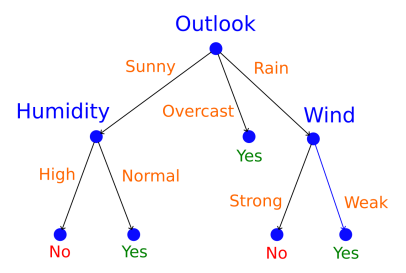
- ► All non-discrete variables.
- ► Have an infinite number of values between any two values.
- ► Their values are measured (??).

Deeper characteristics of data (statistical properties, etc.) will be examined at tutorials.

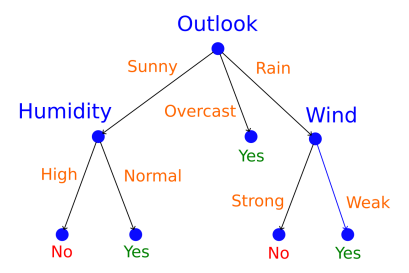
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- ➤ We will consider the ID3 algorithm. Quinlan, 1979
- ▶ Various adjustments that appear in C4.5, CART, etc.

Consider the weather forecast for tennis playing. How would you decide whether to play today?



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How do we obtain such a tree based on experience/data?

Consider data represented as follows:

- ▶ A finite set of *attributes* $A = \{A_1, ..., A_n\}$.
- ▶ Each attribute $A \in A$ has its set of values V(A).

We start with trees on discrete datasets, that is, assume V(A) finite for all $A \in A$.

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Objects to be classified are described by vectors of values of all attributes:

$$\vec{x} = (x_1, \dots, x_n) \in V(A_1) \times \dots \times V(A_n)$$

Given \vec{x} and an attribute A_k we denote by $A_k(\vec{x})$ the value x_k of the attribute A_k in \vec{x} .

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Consider a set C of classes.

We consider a multiclass classification in general, i.e., ${\it C}$ is an arbitrary finite set.

The tennis problem:

► The attributes are:

$$A_1 = Outlook, A_2 = Temperature, A_3 = Humidity, A_4 = Wind$$

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► *C* = { *Yes*, *No*}

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A decision tree is

- ightharpoonup a tree $\mathcal{T} = (T, E)$ where
- ▶ each leaf $\tau \in T_{leaf}$ is assigned a class $C(\tau) \in C$,
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Inference: Given an input \vec{x} , we traverse the tree from the root to a leaf, always choosing edges labeled with values of attributes from \vec{x} . The output is the class labeling the leaf.

$$T = \{O, H, W, z_1, z_2, z_3, z_4, z_5\}$$

$$T_{leaf} = \{z_1, z_2, z_3, z_4, z_5\}, T_{int} = \{O, H, W\}$$

$$E = \{(O, H), (O, W), (H, z_1), (H, z_2), (O, z_3), (W, z_4), (W, z_5)\}$$

$$C(z_1) = C(z_3) = N_O, C(z_2) = C(z_4) = Yes$$
Outlook
High Normal Strong We Yes

$$A(O) = Outlook$$
, $A(H) = Humidity$, $A(W) = Wind$
 $V(O, H) = Sunny$, $V(O, z_3) = Overcast$, $V(O, W) = Rain$
 $V(H, z_1) = High$, $V(H, z_2) = Normal$
 $V(W, z_4) = Strong$, $V(W, z_5) = Weak$

Inference: For (Rain, Hot, High, Strong) we reach z_4 , yielding No.

Training Dataset

Consider a training dataset

$$\mathcal{D} = \{ (\vec{x}_k, c_k) \mid k = 1, \dots, p \}$$

Here $\vec{x}_k \in V(A_1) \times \cdots \times V(A_k)$ and $c_k \in C$ for every k.

Technically $\mathcal D$ can be a multiset containing several occurrences of the same vector.

Index	Outlook	Temperature	Humidity	Wind	PlayTennis
1	Sunny	Hot	High	Weak	No
2	Sunny	Hot	High	Strong	No
3	Overcast	Hot	High	Weak	Yes
4	Rain	Mild	High	Weak	Yes
5	Rain	Cool	Normal	Weak	Yes
6	Rain	Cool	Normal	Strong	No
7	Overcast	Cool	Normal	Strong	Yes
8	Sunny	Mild	High	Weak	No
9	Sunny	Cool	Normal	Weak	Yes
10	Rain	Mild	Normal	Weak	Yes
11	Sunny	Mild	Normal	Strong	Yes
12	Overcast	Mild	High	Strong	Yes
13	Overcast	Hot	Normal	Weak	Yes
14	Rain	Mild	High	Strong	No

$$\mathcal{D} = \big\{ ((Sunny, Hot, High, Weak), No), \\ ((Sunny, Hot, High, Strong), No) \big\}$$

((Rain, Mild, High, Strong), No)}

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 - for every $v \in V(A)$ introduce an edge (τ, τ_v) assigned v.

```
1: function ID3(dataset \mathcal{D}, attribute set \mathcal{A})
         Create a root node \tau for the tree
 2:
         if \mathcal{D} = \emptyset then
 3:
             Return the single node \tau assigned with a default class.
 4:
 5:
         else if all examples in \mathcal{D} are of the same class c then
             Return the single-node tree, where \tau is assigned c
 6:
         else if set of attributes A is empty then
 7:
             Return the single-node tree where \tau is assigned
 8:
             the most common class in \mathcal{D}
 9:
         else
10:
             Choose attribute A \in \mathcal{A} best classifying examples in \mathcal{D}.
             Set the decision attribute for \tau to A
11:
12:
             for each value v \in D(A) do
                  Compute a decision tree ID3(\mathcal{D}_{v}, \mathcal{A} \setminus \{A\}) with root \tau_{v},
13:
                  add a new edge (\tau, \tau_v) assigned v.
14:
             end for
15:
         end if
16:
17: return 	au
18: end function
```

Best Classifying Attribute

We aim to choose an attribute that best informs us about the class.

As a result, we would possibly use as few attributes as possible and obtain a small tree containing only class-relevant decisions.

How to choose an attribute that best classifies examples in \mathcal{D} ?

There are several measures used in practice.

The most common are

- information gain
- Gini impurity decrease

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We need some notation:

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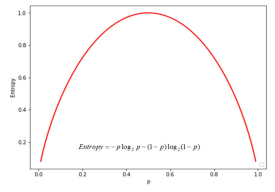
▶ The *information gain* of an attribute A is then defined by

$$\textit{Gain}(\mathcal{D}, \textit{A}) = \textit{Entropy}(\mathcal{D}) - \sum_{\textit{v} \in \textit{V}(\textit{A})} \frac{|\mathcal{D}_{\textit{v}}|}{|\mathcal{D}|} \textit{Entropy}(\mathcal{D}_{\textit{v}})$$

In every step of the ID3 algorithm, we choose an attribute maximizing the information gain for the current dataset \mathcal{D} .

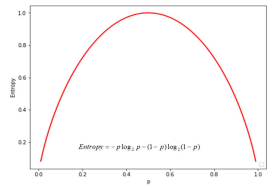
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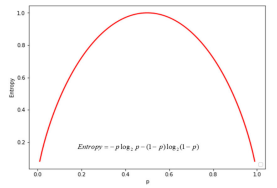
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 $\sum_{v \in V(A)} \frac{|\mathcal{D}_v|}{|\mathcal{D}|} Entropy(\mathcal{D}_v) \text{ is weighted uncertainty of classes}$ in each \mathcal{D}_v (weighted by the relative size of \mathcal{D}_v).

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- ▶ $Gain(\mathcal{D}, A)$ measures reduction in uncertainty of classes by splitting \mathcal{D} according to A.

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$$\mathit{Gini}(\mathcal{D}) = 1 - \sum_{c \in \mathcal{C}} p_c^2$$

► The *impurity decrease* of an attribute *A* is then defined similarly to the gain in the entropy case

$$\mathit{ImpDec}(\mathcal{D}, A) = \mathit{Gini}(\mathcal{D}) - \sum_{v \in V(A)} \frac{|\mathcal{D}_v|}{|\mathcal{D}|} \mathit{Gini}(\mathcal{D}_v)$$

In every step of the ID3 algorithm, we choose an attribute maximizing the impurity decrease for the current dataset \mathcal{D} .

73

What is the intuition behind $\textit{Gini}(\mathcal{D})$?

What is the intuition behind $Gini(\mathcal{D})$?

Assume we randomly independently choose objects from \mathcal{D} .

 $1 - \sum_{c \in C} p_c^2$ is the probability of choosing two objects of different classes in two consecutive independent trials.

Indeed, p_c is the probability of choosing an object of class c, p_c^2 the probability of choosing objects of the class c twice, and $\sum_{c \in C} p_c^2$ the probability of choosing two objects of the same class.

In what follows (and at the exam), we will work only with the Gini impurity as it is easier to compute.

Consider our tennis example (see the table).

- ▶ Consider the whole dataset \mathcal{D} .
 - $p_{Yes} = 9/14$
 - $p_{No} = 5/14$
 - ightharpoonup Gini(D) = 1 (9/14)² (5/14)² = 0.45918

Consider our tennis example (see the table).

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- ightharpoonup For A = Outlook we get
 - ightharpoonup Gini $(\mathcal{D}_{Sunny}) = 1 (2/5)^2 (3/5)^2 = 0.48$
 - $ightharpoonup Gini(\mathcal{D}_{Overcast}) = 1 1^2 0^2 = 0$
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Thus

$$ImpDec(\mathcal{D}, Outlook) = 0.459 - (5/14) \cdot 0.48 - (4/14) \cdot 0 - (5/14) \cdot 0.48$$
$$= 0.117$$

- ▶ $ImpDec(\mathcal{D}, Temperature) = 0.018$
- ▶ $ImpDec(\mathcal{D}, Humidity) = 0.091$
- \blacktriangleright ImpDec(\mathcal{D} , Wind) = 0.030

So the largest information gain is given by the Outlook.

Going further on, consider $\mathcal{D} = \mathcal{D}_{Sunny}$. We get

- ▶ ImpDec(D, Temperature) = 0.279
- ▶ $ImpDec(\mathcal{D}, Humidity) = 0.48$
- ▶ $ImpDec(\mathcal{D}, Wind) = 0.013$

The best choice attribude after *Sunny* in *Outlook* is *Humidity*.

Going further on, consider $\mathcal{D} = \mathcal{D}_{Sunny}$. We get

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The best choice attribude after Sunny in Outlook is Humidity.

Now consider $\mathcal{D} = \mathcal{D}_{Rain}$.

- ▶ ImpDec(D, Temperature) = 0.013
- ▶ $ImpDec(\mathcal{D}, Humidity) = 0.013$
- ► $ImpDec(\mathcal{D}, Wind) = 0.48$

The best choice attribude after Rain in Outlook is Wind.

Continuous-Valued Attributes

What if values of an attribute A come from a continuous variable?

 \boldsymbol{A} is a numerical attribute that can take any value in an interval, such as temperature, size, time, etc.

Continuous-Valued Attributes

What if values of an attribute A come from a continuous variable? A is a numerical attribute that can take any value in an interval, such as temperature, size, time, etc.

Consider an internal node $\tau \in T_{int}$ assigned such a continuous attribute A. Then

- ightharpoonup au is assigned a threshold value called a *cut point* $H \in \mathbb{R}$,
- ▶ there are two edges e_{true} , e_{false} from τ ,
- e_{true} labeled with True and e_{false} labeled with False.

During inference, when considering an example \vec{x} in the node τ ,

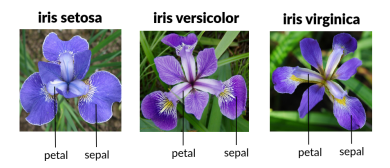
- ightharpoonup evaluate $A(\vec{x}) \leq H$,
- ▶ if $A(\vec{x}) \leq H$, then follow e_{true} ,
- else follow efalse.

True False
te values in the training s

In training, the cut point is chosen from the attribute values in the training set using information gain/impurity decrease similar to discrete attributes.

Temperature ≤ 15

Iris Example



Attributes

Sepal.Length, Sepal.Width, Petal.Length, Petal.Width

Classes (Variety) Setosa, Versicolor, Virginica

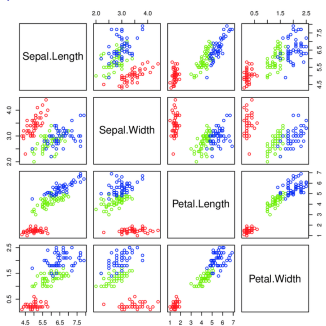
Iris Example

The dataset (150 examples):

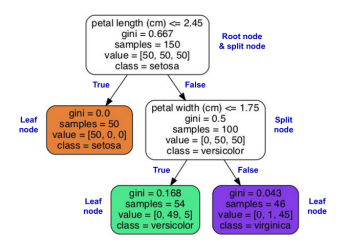
Sepal.Length	Sepal.Width	Petal.Length	Petal.Width	Variety
5.5	3.5	1.3	0.2	Setosa
6.8	2.8	4.8	1.4	Versicolor
6.7	3.1	4.7	1.5	Versicolor
6.9	3.1	5.1	2.3	Virginica
7.3	2.9	6.3	1.8	Virginica
5.4	3.7	1.5	0.2	Setosa
4.6	3.4	1.4	0.3	Setosa
6.2	2.8	4.8	1.8	Virginica
5.4	3.0	4.5	1.5	Versicolor
4.7	3.2	1.6	0.2	Setosa
6.7	3.3	5.7	2.1	Virginica
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4.4	2.9	1.4	0.2	Setosa
6.0	3.4	4.5	1.6	Versicolor
5.1	3.5	1.4	0.2	Setosa
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5.9	3.2	4.8	1.8	Versicolor
5.6	2.8	4.9	2.0	Virginica

79

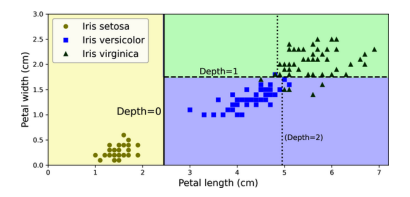
Iris Example



Iris Example - Decision Tree



Iris Example - Decision Tree Boudaries



If the leaves are split further, the Depth = 2 boundary would be added.

How important are attributes for the trained tree \mathcal{T} ? Depends on

- ightharpoonup how close they are to the root of \mathcal{T} ,
- ▶ how large information gain/decrease in impurity they give.

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Then define the importance as the average decrease in Gini impurity (i.e., average ImpDec) in the nodes of T[A]:

$$GiniImportance(A) = \sum_{ au \in T[A]} \frac{|\mathcal{D}[au]|}{|\mathcal{D}|} ImpDec(\mathcal{D}[au], A)$$

Decision Trees

Practical Issues

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- Data preprocessing
- Model tunning (overfitting and underfitting)
- Sensitivity to changes in data/hyperparameters
- ► Learning representation problems (the XOR)

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Imbalanced classes might cause problems because of small information gain/impurity decrease in splitting.

Imbalanced Classes

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- ▶ there are two classes, $C = \{0, 1\}$,
- ▶ 10⁶ examples have the class 1,
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- thus the Gini impurity $1 p_1^2 p_0^2 \approx 0$.

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Consider an attribute A with $V(A) = \{a, b\}$.

Splitting \mathcal{D} according to A gives to sets \mathcal{D}_a and \mathcal{D}_b .

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Splitting $\mathcal D$ according to A gives to sets $\mathcal D_a$ and $\mathcal D_b$.

What is the impurity decrease caused by the attribute?

$$ImpDec(\mathcal{D}, A) = Gini(\mathcal{D}) - \frac{|\mathcal{D}_a|}{|\mathcal{D}|}Gini(\mathcal{D}_a) - \frac{|\mathcal{D}_b|}{|\mathcal{D}|}Gini(\mathcal{D}_b)$$

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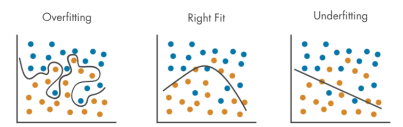
$$ImpDec(\mathcal{D}, A) = Gini(\mathcal{D}) - \frac{|\mathcal{D}_a|}{|\mathcal{D}|}Gini(\mathcal{D}_a) - \frac{|\mathcal{D}_b|}{|\mathcal{D}|}Gini(\mathcal{D}_b)$$

For small $|\mathcal{D}_{\mathsf{a}}|$ (say ≤ 1000) we have small $|\mathcal{D}_{\mathsf{a}}|/|\mathcal{D}|$

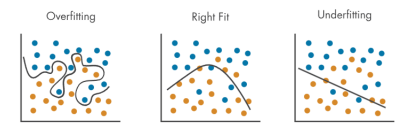
For not so small \mathcal{D}_a we have $Gini(\mathcal{D}_a) \approx 0$.

In both cases, the impurity decrease is very small.

The behavior of the model on the training set:

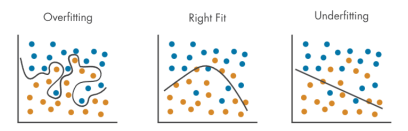


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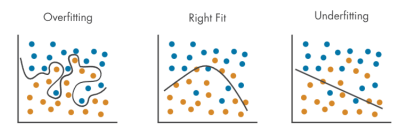
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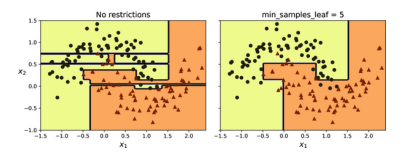
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The behavior of the model on the training set:



- ► The left one is strongly overfitting. It would possibly not work well on new data.
- ► The right one is strongly underfitting. It would probably give poor classification results.
- ► The middle one seems good (but still needs to be tested on fresh data).

Model Tuning - Overfitting in Decision Trees

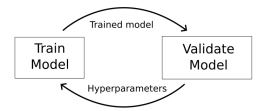


See the overfitting on the left and the "nice" model on the right.

Both overfitting and underfitting are best avoided. But how do we find out?

Model Tuning (In General)

Recall from the first lecture:

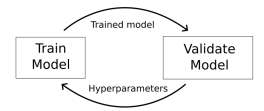


The validation should be done on a **validation set** separated from the training set.

We will discuss more sophisticated techniques later.

Model Tuning (In General)

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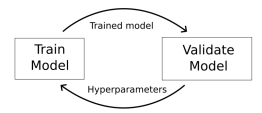
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We will discuss more sophisticated techniques later.

What hyperparameters to set? (see the next slide)

Model Tuning (In General)

Recall from the first lecture:



The validation should be done on a **validation set** separated from the training set.

We will discuss more sophisticated techniques later.

What hyperparameters to set? (see the next slide)

What to observe? In the case of decision trees, one should observe the difference between performance measures (e.g., classification accuracy) on the training and validation sets.

The too-large difference implies an improperly fitting model.

There are several approaches available for decision trees.

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The post-pruning approach has been more successful in practice than the pre-pruning because it is usually hard to say when to stop growing the tree.

We shall meet this controversy also in deep learning, where recent history shows a similar phenomenon.

The ensemble methods will be covered later when we discuss random forests.

Hyperparameters controlling the size of the tree:

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- Minimum number of examples required to be in a leaf Similar to the previous one. A higher number means we cannot have very specific branches concerned with particular combinations of values.
- ▶ Minimum information gain/impurity decrease

 A small impurity decrease means that the split does not contribute too much to the classification (their proportions after a split are similar to proportions before a split). However, keep in mind that it is weighted average impurity after the split.

Post-Pruning - Reduced Error Pruning

Train a large tree and then remove nodes that make classification worse on the validation set.

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Given a decision tree \mathcal{T} and its internal node $\tau \in \mathcal{T}_{int}$, we denote by $\mathcal{T}_{-\tau}$ the tree obtained from \mathcal{T} by removing the subtree rooted in τ , i.e., τ is a leaf of $\mathcal{T}_{-\tau}$.

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```
1: Train \mathcal{T} to maximum fit on the training dataset.
 2: while true do
 3:
            Err[\mathcal{T}] \leftarrow the error of \mathcal{T} on the validation set.
            for \tau \in T_{int} do
 4:
                  Err[\mathcal{T}_{-\tau}] \leftarrow the error of \mathcal{T}_{-\tau} on the validation set.
 5:
           end for
 6:
            if Err[\mathcal{T}] \leq \min\{Err[\mathcal{T}_{-\tau}] \mid \tau \in \mathcal{T}_{int}\}\ then return \mathcal{T}
 7:
 8:
           else
                 \mathcal{T} \leftarrow argmin\{Err[\mathcal{T}_{-\tau}] \mid \tau \in \mathcal{T}_{int}\}
 9:
            end if
10:
11: end while
```

The error $Err[\mathcal{T}]$ can be any measure of the "badness" of the decision tree \mathcal{T} . For example, 1-Accuracy.

Other Pruning Methods

There are more pruning methods.

- ► Rule Post-Pruning:
 - Transform the tree into a set of rules.
 Rules correspond to paths in the tree; they have a form of implication: Specific values of attributes imply a class.
 - Remove the attribute conditions from the premises of the implications.

This gives a more refined pruning: Instead of removing the whole subtree, each path of the subtree can be pruned individually.

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Using cost complexity measure: Evaluate trees not only based on the classification error but also based on their size.

Typically introduce regularization into the error functions: Given a decision tree \mathcal{T}

$$Err_{\alpha}(\mathcal{T}) = Err(\mathcal{T}) + \alpha |\mathcal{T}|$$

The original paper by Breiman et al. (1984) defined $Err(\mathcal{T})$ to be the misclassification rate on the training dataset, and $|\mathcal{T}|$ is the number of nodes of the tree \mathcal{T} .

Sensitivity to Small Changes and Randomness

- Decision trees are sensitive to small changes in data and hyperparameters.
 - Several attributes may provide (almost) identical information gain but divide the training dataset very differently.
- ➤ Some implementations choose attributes partially in random (sci-kit-learn). You may get completely different trees.

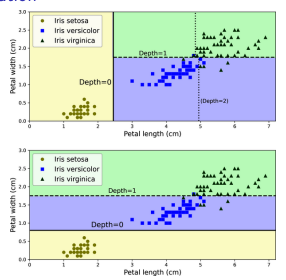
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A solution is to train an ensemble of many decision trees and then use majority voting for classification.

This is the fundamental idea behind random forests (see later lectures).

Iris - Illustration

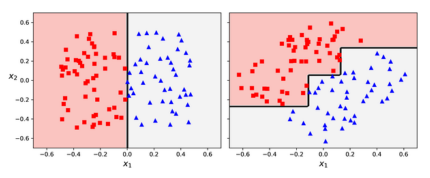


Decision trees trained on the Iris dataset.

Iris Setosa is perfectly separated by many choices for the first split.

Axis Sensitivity

The decision makes divisions along particular axes:

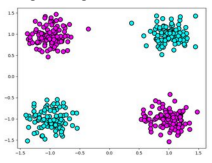


That is, rotated data may result in a completely different model.

That is why decision trees are often preceded by the *principal* component analysis (PCA) transformation, which aligns data along the axes of maximum data variance.

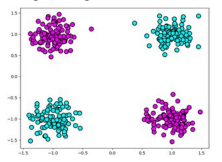
XOR Training Problem

Consider the following training dataset:

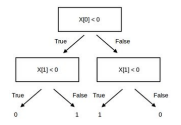


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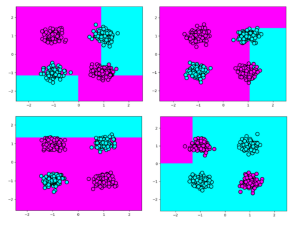


An ideal decision tree would look like this:



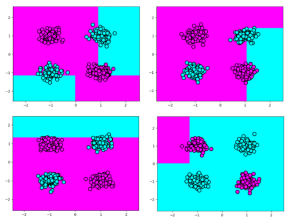
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Max depth = 2:



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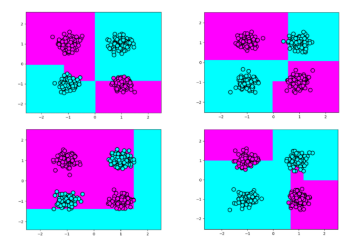


The problem: Both information gain and decrease in impurity consider only the relationship of a *single* attribute and the class.

However, there is no relationship between a single attribute and the class; both attributes need to be considered together!

More Attempts at Training on XOR

Max depth = 3:



It's better but still fails occasionally.

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- Little data preparation, unlike other techniques requiring normalization, dummy variables, or missing value removal.
- ► Handles numerical and categorical data.
- Not sensitive to outliers since the splitting is based on the proportion of examples within the split ranges and not on absolute values.
- ► The cost of using a well-balanced tree is logarithmic in the number of data points used to train it.

- Overfitting: Trees can be over-complex and not generalize well, needing pruning or limits on tree depth.
- Instability: Small data variations can result in very different trees. This is mitigated in ensemble methods.
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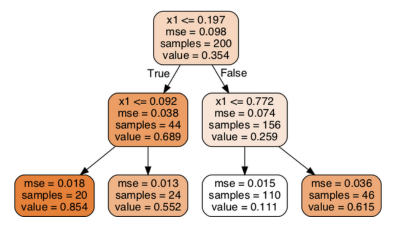
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- Difficulty expressing certain concepts, such as XOR, parity, or multiplexer problems (see the next slide).
- ▶ Bias in trees: Decision trees can create biased trees if some classes dominate. Balancing the dataset is recommended.
- ► Learning optimal trees is NP-complete: Heuristic algorithms like greedy algorithms are used, which do not guarantee globally optimal trees. Ensemble methods can help.

History of Decision Trees

- Hunt and colleagues use exhaustive search decision-tree methods (CLS) to model human concept learning in the 1960's.
- ▶ In the late 70's, Quinlan developed ID3 with the information gain heuristic to learn expert systems from examples.
- Simultaneously, Breiman, Friedman, and colleagues develop CART (Classification and Regression Trees), similar to ID3.
- In the 1980s, various improvements were introduced to handle noise, continuous features, missing features, and improved splitting criteria. Various expert-system development tools results.
- Quinlan's updated decision-tree package (C4.5) released in 1993.

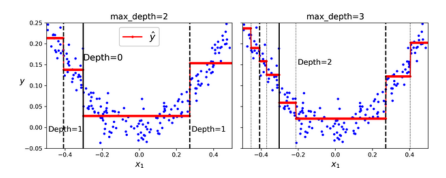
Comment on Regression Trees

Decision trees can also be used to approximate functions. Assign a function value to the leaves instead of classes.



Here, "mse" is the mean-squared-error.

Comment on Regression Trees



Intuitively, for every subinterval of x_1 , the value (the red line) is at the average y over the subinterval.

How are the subintervals being set?

A *regression tree* is a decision tree whose leaves are labeled by values from \mathbb{R} .

We follow the same procedure as in decision trees during inference on an input \vec{x} .

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The training procedure is the same as for the decision trees, except that the splits and cut points are selected differently.

Given a dataset $\mathcal{D} = \{(\vec{x}_1, d_1), \dots, (\vec{x}_p, d_p)\}$, we denote by $\bar{\mathcal{D}}$ the average *desired* value in \mathcal{D} , that is $\bar{\mathcal{D}} = \frac{1}{p} \sum_{k=1}^{p} d_k$.

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We are looking for a value H of the attribute A such that the split:

$$\mathcal{D}_{\leq H} = \{ (\vec{x}, d) \in \mathcal{D} \mid A(\vec{x}) \leq H \} \quad \mathcal{D}_{>H} = \{ (\vec{x}, d) \in \mathcal{D} \mid A(\vec{x}) > H \}$$

Minimizes the following split error:

$$\frac{1}{|\mathcal{D}_{\leq H}|} \sum_{(\vec{x}, d) \in \mathcal{D}_{\leq H}} \left(d - \bar{\mathcal{D}}_{\leq H}\right)^2 + \frac{1}{|\mathcal{D}_{> H}|} \sum_{(\vec{x}, d) \in \mathcal{D}_{> H}} \left(d - \bar{\mathcal{D}}_{> H}\right)^2$$

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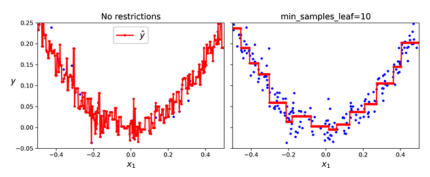
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If Δ is large enough, split on A and H that minimize the split error. Otherwise, stop splitting and label the leaf with \bar{D} .

107

Without any lower bound on the number of examples in the leaves, the algorithm will eventually overfit by splitting into (possibly) singleton leaves.



Probabilistic Classification

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The degree of belief (Bayesian), or the relative frequency (frequentists), is the *probability*.

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Experiment: Roll one dice once. Sample space: $\Omega = \{1, \dots, 6\}$

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▶ Basic laws: $P(\Omega) = 1$, $P(\emptyset) = 0$, given disjoint sets A, B we have $P(A \cup B) = P(A) + P(B)$, $P(\Omega \setminus A) = 1 - P(A)$.

Conditional Probability and Independence

▶ $P(A \mid B)$ is the probability of A given B (assume P(B) > 0) defined by

$$P(A \mid B) = P(A \cap B)/P(B)$$

(We assume that B is all and only information known.)

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▶ A and B are independent if $P(A \cap B) = P(A) \cdot P(B)$. It is easy to show that if P(B) > 0, then
A, B are independent iff $P(A \mid B) = P(A)$.

Random Variables and Random Vectors

- A random variable X is a function $X : \Omega \to \mathbb{R}$. A dice: $X : \{1, ..., 6\} \to \{0, 1\}$ such that $X(n) = n \mod 2$.
- ▶ A *random vector* is a function $X : \Omega \to \mathbb{R}^d$.

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- ▶ A random vector is a function $X : \Omega \to \mathbb{R}^d$. We use $X = (X_1, \dots, X_d)$ where X_i is a random variable returning the *i*-th component of X.
- Consider random variables X_1, X_2 and Y. The variables X_1, X_2 are *conditionally independent given* Y if for all x_1, x_2 and y we have that

$$P(X_1 = x_1, X_2 = x_2 \mid Y = y) =$$

 $P(X_1 = x_1 \mid Y = y) \cdot P(X_2 = x_2 \mid Y = y)$

Random Vectors – Example

Let Ω be a space of colored geometric shapes that are divided into two categories (1 and 0).

Assume a random vector $X = (X_{color}, X_{shape}, X_{cat})$ where

- $ightharpoonup X_{color}: \Omega \rightarrow \{red, blue\},$
- ► $X_{shape}: \Omega \rightarrow \{circle, square\},\$
- $X_{cat} : \Omega \to \{\mathbf{1}, \mathbf{0}\}.$

The following tables give probability distribution of values:

category 1:

	circle	square
red	0.2	0.02
blue	0.02	0.01

category **0**:

	circle	square
red	0.05	0.3
blue	0.2	0.2

Random Vectors – Example

Example:

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$$P(red, circle) = P(X_{color} = red, X_{shape} = circle)$$

= $P(red, circle, \mathbf{1}) + P(red, circle, \mathbf{0})$
= $0.2 + 0.05 = 0.25$

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Thus also, all conditional probabilities can be computed:

$$P(\mathbf{1} \mid red, circle) = \frac{P(red, circle, \mathbf{1})}{P(red, circle)} = \frac{0.2}{0.25} = 0.8$$

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 - ▶ Denote by $\vec{x} \in \mathbb{R}^n$ values of X,
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Bayes classifier: Given a vector of feature values \vec{x} ,

$$C^{Bayes}(\vec{x}) := egin{cases} \mathbf{1} & ext{if } P(Y = \mathbf{1} \mid X = \vec{x}) \geq P(Y = \mathbf{0} \mid X = \vec{x}) \\ \mathbf{0} & ext{otherwise}. \end{cases}$$

Intuitively, C^{Bayes} assigns to \vec{x} the most probable category it might be in.

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A machine is supposed to correctly distinguish apples from apricots based on their weight and diameter.

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We are given a fruit of a diameter of 5cm that weighs 40g.

The Bayes classifier compares $P(Y = 1 \mid X = (40g, 5cm))$ with $P(Y = 0 \mid X = (40g, 5cm))$ and selects the more probable category given the features.

Crucial question: Is such a classifier good?

There are other classifiers, e.g., one which compares the weight divided by 10 with the diameter and decides based on the answer, or maybe a classifier that sums the weight and the diameter and compares the result with a constant, etc.

Bayes Classifier

Let C be an arbitrary *classifier*, that is a function that to every feature vector $\vec{x} \in \mathbb{R}^n$ assigns a class from $\{0,1\}$.

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Define the error of the classifier C by

$$E_C = P(Y \neq C)$$

(Here we slightly abuse notation and apply C to samples, technically we apply the composition $C \circ X$ of C and X which first determines the features using X and then classifies according to C).

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Theorem

The Bayes classifier C^{Bayes} minimizes E_C , that is

$$E_{C^{Bayes}} = \min_{C \ is \ a \ classifier} E_{C}$$

Practical Use of Bayes Classifier

The crucial problem: The probability P is not known! In particular, where to get $P(Y=\mathbf{1}\mid X=\vec{x})$? Note that $P(Y=\mathbf{0}\mid X=\vec{x})=1-P(Y=\mathbf{1}\mid X=\vec{x})$

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Given no other assumptions, this requires a table showing the probability of the category $\mathbf{1}$ for each possible feature vector \vec{x} .

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Where do you get these probabilities?

In some cases, the probabilities might come from the knowledge of the solved problem (e.g., applications in physics might be supported by a theory giving the probabilities).

In most cases, however, P is estimated from sampled data by

$$\bar{P}(Y = 1 \mid X = \vec{x}) = \frac{\text{number of samples with } Y = 1 \text{ and } X = \vec{x}}{\text{number of samples with } X = \vec{x}}$$

(We use \bar{P} to denote an estimate of P from data.)

Estimating P

Consider a problem with $X = (X_1, X_2, X_3)$ where each X_i returns either 0 or 1. What might the data look like?

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Part of the data table:

Y	<i>X</i> ₁	X_2	<i>X</i> ₃
1	1	0	1
1	0	1	1
0	1	0	1
0	0	0	1
1	0	0	0
0	1	1	1
•••			

All data with $X_1 = 1$, $X_2 = 0$, $X_3 = 1$:

Y	X_1	<i>X</i> ₂ 0	<i>X</i> ₃
1	1	0	1
1	1	0	1
0	1	0	1
0	1	0	1
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The probability table and the necessary data are typically too large!

Concretely, if all $X_1, ..., X_n$ are binary, there are 2^n probabilities $P(Y = 1 \mid X = \vec{x})$, one for each possible $\vec{x} \in \{0, 1\}^n$.

Let's Look at It the Other Way Round

Theorem (Bayes, 1764)

$$P(A \mid B) = \frac{P(B \mid A) \cdot P(A)}{P(B)}$$

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Proof.

$$P(A \mid B) = \frac{P(A \cap B)}{P(B)} = \frac{\frac{P(A \cap B)}{P(A)} \cdot P(A)}{P(B)} = \frac{P(B \mid A) \cdot P(A)}{P(B)}$$

121

Determine the category for \vec{x} by computing

$$P(Y = y \mid X = \vec{x}) = \frac{P(Y = y) \cdot P(X = \vec{x} \mid Y = y)}{P(X = \vec{x})}$$

for both $y \in \{0,1\}$ and deciding whether or not the following holds:

$$P(Y = 1 \mid X = \vec{x}) \ge P(Y = 0 \mid X = \vec{x})$$

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So, to make the classifier, we need to compute the following:

- ▶ The prior P(Y = 1) (then P(Y = 0) = 1 P(Y = 1))
- ► The conditionals $P(X = \vec{x} \mid Y = y)$ for $y \in \{0,1\}$ and for every \vec{x}

Estimating the Prior and Conditionals

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If the dimension of features is small, $P(X = \vec{x} \mid Y = y)$ can be estimated from data similarly as $P(Y = 1 \mid X = \vec{x})$ by

$$\bar{P}(X = \vec{x} \mid Y = y) = \frac{\text{number of samples with } Y = y \text{ and } X = \vec{x}}{\text{number of samples with } Y = y}$$

Unfortunately, for higher dimensional data too many samples are needed to estimate all $P(X = \vec{x} \mid Y = y)$ (there are too many \vec{x} 's).

So where is the advantage of using the Bayes thm.??

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So where is the advantage of using the Bayes thm.??

We introduce independence assumptions about the features!

Naive Bayes

We assume that features are (conditionally) independent *given* the category. That is for all $\vec{x} = (x_1, \dots, x_n)$ and $y \in \{0, 1\}$ we assume:

$$P(X = x \mid Y = y) = P(X_1 = x_1, \dots, X_n = x_n \mid Y)$$

$$= \prod_{i=1}^{n} P(X_i = x_i \mid Y = y)$$

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► Therefore, we only need to specify $P(X_i = x_i \mid Y = y)$ for each possible pair of a feature-value x_i and $y \in \{0, 1\}$.

Note that if all X_i are binary (values in $\{0,1\}$), this requires specifying only 2n parameters:

$$P(X_i = 1 \mid Y = \mathbf{1})$$
 and $P(X_i = 1 \mid Y = \mathbf{0})$ for each X_i as $P(X_i = 0 \mid Y = y) = 1 - P(X_i = 1 \mid Y = y)$ for $y \in \{\mathbf{0}, \mathbf{1}\}$.

Compared to specifying 2^n parameters without any independence assumption.

Estimating the marginal probabilities

Estimate the probabilities $P(X_i = x_i \mid Y = y)$ by

$$\bar{P}(X_i = x_i \mid Y = y) = \frac{\text{number of samples with } X_i = x_i \text{ and } Y = y}{\text{number of samples with } Y = y}$$

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Example: Consider a problem with $X = (X_1, X_2, X_3)$ where each X_i returns either 0 or 1. The data is

Y	X_1	X_2	<i>X</i> ₃
1	1	0	1
1	0	1	1
0	1	0	1
0	0	0	1
1	0	0	0
0	1	1	1

$$ar{P}(X_1 = 1 \mid Y = 1) = 1/3$$
 $ar{P}(X_1 = 1 \mid Y = 0) = 2/3$
 $ar{P}(X_2 = 1 \mid Y = 1) = 1/3$ $ar{P}(X_2 = 1 \mid Y = 0) = 1/3$
 $ar{P}(X_3 = 1 \mid Y = 1) = 2/3$ $ar{P}(X_3 = 1 \mid Y = 0) = 1$

Naive Bayes - Example

Consider classification of geometric shapes:

 $X_1 \in \{small, medium, large\}$

 $X_2 \in \{\textit{red}, \textit{blue}, \textit{green}\}$

 $X_3 \in \{ square, triangle, circle \}$

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 $X_2 \in \{red, blue, green\}$

 $X_3 \in \{square, triangle, circle\}$

Assume that we have already estimated the following probabilities:

	-
Y = 1	Y = 0
0.5	0.5
0.4	0.4
0.1	0.2
0.5	0.4
0.9	0.3
0.05	0.3
0.05	0.4
0.05	0.4
0.05	0.3
0.9	0.3
	0.5 0.4 0.1 0.5 0.9 0.05 0.05 0.05 0.05

Does (medium, red, circle) belong to the category 1?

	Y = 1	Y = 0
$\bar{P}(Y)$	0.5	0.5
$\bar{P}(medium \mid Y)$	0.1	0.2
$\bar{P}(red \mid Y)$	0.9	0.3
P(circle Y)	0.9	0.3

	Y = 1	<i>Y</i> = 0
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$\bar{P}(red \mid Y)$	0.9	0.3
P̄(circle Y)	0.9	0.3

$$P(Y = 1 \mid X = \vec{x}) = = P(1) \cdot P(medium \mid 1) \cdot P(red \mid 1) \cdot P(circle \mid 1) / P(X = \vec{x}) = 0.5 \cdot 0.1 \cdot 0.9 \cdot 0.9 / P(X = \vec{x}) = 0.0405 / P(X = \vec{x})$$

	Y = 1	Y = 0
$\bar{P}(Y)$	0.5	0.5
$\bar{P}(medium \mid Y)$	0.1	0.2
P(red Y)	0.9	0.3
P̄(circle Y)	0.9	0.3

$$P(Y = 1 \mid X = \vec{x}) = = P(1) \cdot P(medium \mid 1) \cdot P(red \mid 1) \cdot P(circle \mid 1) / P(X = \vec{x}) = 0.5 \cdot 0.1 \cdot 0.9 \cdot 0.9 / P(X = \vec{x}) = 0.0405 / P(X = \vec{x})$$

$$P(Y = \mathbf{0} \mid X = \vec{x}) =$$

= $P(\mathbf{0}) \cdot P(medium \mid \mathbf{0}) \cdot P(red \mid \mathbf{0}) \cdot P(circle \mid \mathbf{0}) / P(X = \vec{x})$
= $0.5 \cdot 0.2 \cdot 0.3 \cdot 0.3 / P(X = \vec{x}) = 0.009 / P(X = \vec{x})$

(Note that we used the estimates \bar{P} of P to finish the computation above.)

Y = 1	<i>Y</i> = 0
0.5	0.5
0.1	0.2
0.9	0.3
0.9	0.3
	0.5 0.1 0.9

$$P(Y = \mathbf{1} \mid X = \vec{x}) =$$

$$= P(\mathbf{1}) \cdot P(medium \mid \mathbf{1}) \cdot P(red \mid \mathbf{1}) \cdot P(circle \mid \mathbf{1}) / P(X = \vec{x})$$

$$\doteq 0.5 \cdot 0.1 \cdot 0.9 \cdot 0.9 / P(X = \vec{x}) = 0.0405 / P(X = \vec{x})$$

$$P(Y = \mathbf{0} \mid X = \vec{x}) = \\ = P(\mathbf{0}) \cdot P(medium \mid \mathbf{0}) \cdot P(red \mid \mathbf{0}) \cdot P(circle \mid \mathbf{0}) / P(X = \vec{x}) \\ \doteq 0.5 \cdot 0.2 \cdot 0.3 \cdot 0.3 / P(X = \vec{x}) = 0.009 / P(X = \vec{x})$$

(Note that we used the estimates \bar{P} of P to finish the computation above.) Apparently,

$$P(Y = 1 \mid X = \vec{x}) \doteq 0.0405/P(X = \vec{x}) > 0.009/P(X = \vec{x}) \doteq P(0 \mid X = \vec{x})$$

So we classify \vec{x} to the category **1**.

Estimating Probabilities in Practice

We already know that $P(X_i = x_i \mid Y = y)$ can be estimated by

$$\bar{P}(X_i = x_i \mid Y = y) = \ell_{y,x_i} / \ell_y$$

where

- $ightharpoonup \ell_y = \text{number of samples with } Y = y$

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We already know that $P(X_i = x_i \mid Y = y)$ can be estimated by

$$\bar{P}(X_i = x_i \mid Y = y) = \ell_{y,x_i} / \ell_y$$

where

- \blacktriangleright ℓ_{y,x_i} = number of samples with Y=y and $X_i=x_i$
- ho ℓ_y = number of samples with Y = y

Problem: If, by chance, a rare value x_i of a feature X_i never occurs in the training data, we get

$$\bar{P}(X_i = x_i \mid Y = y) = 0$$
 for both $y \in \{0, 1\}$

But then $\bar{P}(X = x) = 0$ for x containing the value x_i for X_i , and thus $\bar{P}(Y = y \mid X = x)$ is not well defined.

Moreover, $\bar{P}(Y = y) \cdot \bar{P}(X = x \mid Y = y) = 0$ (for $y \in \{0, 1\}$) so even this cannot be used for classification.

Probability Estimation Example

Training data:

Size	Color	Shape	Class	
small	red	circle	1	
large	red	circle	1	
small	red	triangle	0	
large	blue	circle	0	

Estimated probabilities:

	Y = 1	Y = 0
$\bar{P}(Y)$	0.5	0.5
$\bar{P}(small \mid Y)$	0.5	0.5
$\bar{P}(medium \mid Y)$	0	0
$\bar{P}(large \mid Y)$	0.5	0.5
$\bar{P}(red \mid Y)$	1	0.5
$\bar{P}(blue \mid Y)$	0	0.5
$\bar{P}(green \mid Y)$	0	0
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(0)		
$\bar{P}(square \mid Y)$	0	0
$\bar{P}(triangle \mid Y)$	0	0.5
$ar{P}(\mathit{circle} \mid Y)$	1	0.5

Note that $\bar{P}(medium \mid \mathbf{1}) = P(medium \mid \mathbf{0}) = 0$ and thus also $\bar{P}(medium, red, circle) = 0$.

So what is $\bar{P}(1 \mid medium, red, circle)$?

Smoothing

► To account for estimation from small samples, probability estimates are adjusted or *smoothed*.

Smoothing

- ► To account for estimation from small samples, probability estimates are adjusted or *smoothed*.
- Laplace smoothing adds one to every count of feature values

$$\tilde{P}(X_i = x_i \mid Y = y) = \frac{\ell_{y,x_i} + 1}{\ell_y + v_i}$$

where

- ho ℓ_v = number of training samples with Y = y,
- $\ell_{y,x_i} = \text{number of training samples with } Y = y \text{ and } X_i = x_i,$
- \triangleright v_i is the number of all distinct values of the variable X_i .

To understand note that

$$\ell_y = \sum_{x_i \text{ is a value of } X_i} \ell_{y,x_i}$$

and thus

$$\begin{split} \bar{P}(X_i = x_i \mid Y = y) &= \ell_{y, x_i} / \sum_{x_i \text{ is a value of } X_i} \ell_{y, x_i} \\ \tilde{P}(X_i = x_i \mid Y = y) &= (\ell_{y, x_i} + 1) / \sum_{x_i \text{ is a value of } X_i} (\ell_{y, x_i} + 1) \end{split}$$

Laplace Smoothing Example

- ► Assume training set contains 10 samples of category 1:
 - ► 4 small
 - 0 medium
 - ► 6 large

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- Assume training set contains 10 samples of category 1:
 - ▶ 4 small
 - 0 medium
 - 6 large
- Estimate parameters as follows
 - $\tilde{P}(small \mid \mathbf{1}) = (4+1)/(10+3) = 0.384$
 - $\tilde{P}(medium \mid \mathbf{1}) = (0+1)/(10+3) = 0.0769$
 - $\tilde{P}(large \mid \mathbf{1}) = (6+1)/(10+3) = 0.538$

Continuous Features

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Usually, $P(X_i | Y = y)$ is used to denote the *density* of X_i conditioned on Y = y.

- ► The densities $P(X_i | Y = y)$ are usually estimated using Gaussian densities as follows:
 - Estimate the mean μ_{iy} and the standard deviation σ_{iy} based on training data.
 - Then put

$$\bar{P}(X_i \mid Y = y) = \frac{1}{\sigma_{iy}\sqrt{2\pi}} \exp\left(\frac{-(X_i - \mu_{iy})^2}{2\sigma_{iy}^2}\right)$$

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 - Even if the probabilities are not accurately estimated, it often picks the correct maximum probability category.
- Directly constructs a model from parameter estimates that are calculated from the training data.
- Typically handles outliers and noise well.
- Missing values are easy to deal with (simply average overall missing values in feature vectors).

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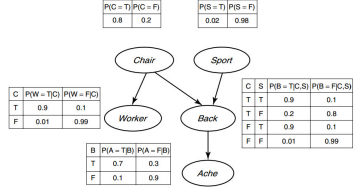
E.g. Variables "rain" and "grass wet" are (usually) strongly dependent.

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(But now in a well-defined sense.)

Bayesian networks are a graphical model that uses a directed acyclic graph to specify dependencies among variables.

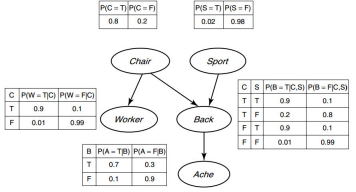
Bayasian Natworks - Frampla



Now, e.g.,

$$P(C, S, W, B, A) = P(C) \cdot P(S) \cdot P(W \mid C) \cdot P(B \mid C, S) \cdot P(A \mid B)$$

Bayasian Natworks - Framnla

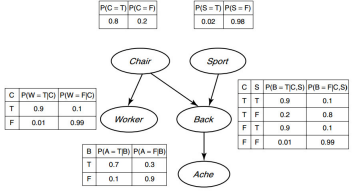


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Now, we may, e.g., infer the probability $P(C = T \mid A = T)$ that we sit in the wrong chair, assuming that our back aches.

We have to store only 10 numbers as opposed to $2^5 - 1$ possible probabilities for all vectors of values of C, S, W, B, A.

Bayesian Networks - Learning & Naive Bayes

Many algorithms have been developed for learning:

- the structure of the graph of the network,
- the conditional probability tables.

The methods are based on maximum-likelihood estimation, gradient descent, etc.

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Can you express the naive Bayes for $Y, X_1, ..., X_n$ using a Bayesian network?

Classifier Evaluation

Assume binary classification into two classes $\{0,1\}$.

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Consider a classification dataset:

$$\{(\vec{x}_k, c_k) \mid k = 1, \dots, p\}$$

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There are many possible metrics ...

I will call the class 1 *positive* and the class 0 *negative*.

Note that the class 0 is not negative in the numerical sense but in the absence of something (e.g., predicted illness).

Confusion Matrix for Binary Classifier

		Predicted	
		1	0
Actual	1	TP	FN
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Example

Given a sample of 12 individuals, eight have cancer, and four are cancer-free.

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Assume that we have trained a classifier with the following results:

Index	1	2	3	4	5	6	7	8	9	10	11	12
Actual	1	1	1	1	1	1	1	1	0	0	0	0
Predicted	0	0	1	1	1	1	1	1	1	0	0	0
Result	FN	FN	TP	TP	TP	TP	TP	TP	FP	TN	TN	TN

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Actual condition	Predicted condition				
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Terminology

- ► TP aka hit
- ► TN aka correct rejection
- ► FP aka type I error, false alarm, overestimation
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In what follows, we also use

- ightharpoonup P = TP + FN of all cases with the *actual* class 1
- ightharpoonup N = TN + FP of all cases with the *actual* class 0
- ▶ PP = TP + FP of all cases with the *predicted* class 1
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There is a large number of derived metrics. We consider some of the most used in practice.

Accuracy

$$\mbox{Accuracy} = \frac{\mbox{TP} + \mbox{TN}}{\mbox{P} + \mbox{N}}$$

Intuitively, Accuracy is the proportion of correctly classified cases w.r.t. all cases.

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Example: Consider our cancer predictor with the confusion matrix

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The Accuracy is

$$ACC = \frac{TP + TN}{P + N} = \frac{6+3}{12} = \frac{3}{4}$$

Accuracy can be misleading when the classes are imbalanced:

- Consider 100 cases, 90 in the class 0 and 10 in the class 1,
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The Accuracy is 91/100 > 0.9. Pretty good, right?

However, the classifier is pretty bad in the positive cases.

In the case of cancer prediction, such a classifier would be a disaster.

Precision & Recall

To mitigate the defect of the Accuracy, we may compute the following metrics:

$$Precision = \frac{TP}{PP} \quad (= \text{how often is predicted positive actually positive})$$

Precision is also known as positive predictive value (PPV)

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$$Recall = \frac{TP}{P} \quad (= \text{how often is actually positive predicted positive})$$

Recall is also known as true positive rate, sensitivity, hit rate, and power.

Precision & Recall - Example

Example: In our cancer example:

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- Precision measures how often is the patient predicted to be ill truly ill (in our case, 6/7)
- Recall measures how often is an ill patient found to be ill (in our case, 6/8)

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$$\begin{aligned} &\mathsf{Precision} = 1 \\ &\mathsf{Recall} = \frac{1}{10} \end{aligned}$$

You can see that the predictor is very precise (on the class 1) but useless due to the weak Recall.

Let us get back to our cancer example:

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Consider *Precision* and *Recall*.

By now, you should remember what they measure.

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Which of the two is more important in medicine?

Which of the two is more important for plagiarism detectors?

Can we get a single number summarizing both Precision and Recall?

For example, to compare two classifiers.

F_1 Score

 F_1 score is the harmonic mean of Recall and Precision:

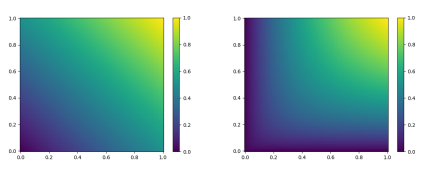
$$\mathsf{F_1} = \frac{2}{\mathsf{Recall}^{-1} + \mathsf{Precision}^{-1}} = \frac{2\mathsf{TP}}{2\mathsf{TP} + \mathsf{FP} + \mathsf{FN}}$$

F_1 Score

 F_1 score is the harmonic mean of Recall and Precision:

$$\mathsf{F_1} = \frac{2}{\mathsf{Recall}^{-1} + \mathsf{Precision}^{-1}} = \frac{2\mathsf{TP}}{2\mathsf{TP} + \mathsf{FP} + \mathsf{FN}}$$

Compare the arithmetic (left) and harmonic (right) mean:



The harmonic mean prefers the two values closer to each other. For example, the harmonic mean of 2/3 and 1/3 is (approx) 0.44444.

F_1 Score - Examples

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Here
$$F_1 = \frac{2\text{TP}}{2\text{TP}+\text{FP}+\text{FN}} = (2 \cdot 6)/((2 \cdot 6) + 1 + 2) = 0.8$$
.

F_1 Score - Examples

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.

Our imbalanced example:

Actual	Predicted	
	Pos Neg	
Pos	1	9
Neg	0	90
Total	90 +	10 = 100

Here
$$F_1 = \frac{2\text{TP}}{2\text{TP}+\text{FP}+\text{FN}} = (2 \cdot 1)/((2 \cdot 1) + 0 + 9) = 0.18$$
.
Note that the average of Precision and Recall is 0.55, which would give us a

much less severe warning that the classifier is bad.

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	Pos	Neg
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Precision =
$$90/99$$
 Recall = $90/90$
$$F_1 = \frac{2TP}{2TP + FP + FN} = (2 \cdot 90)/(2 \cdot 90 + 9 + 0) = 0.95$$

Imbalanced Classes Once More

Note that the standard definitions of Precision and Recall for binary classifiers reveal only part of the truth.

In particular, true negatives are not used in the definition of F_1 .

Consider

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Precision =
$$90/99$$
 Recall = $90/90$
 $F_1 = \frac{2TP}{2TP + FP + FN} = (2 \cdot 90)/(2 \cdot 90 + 9 + 0) = 0.95$

All great, except that the classifier sucks on the negative cases. If you are concerned with the negative cases, swap the classes and compute another set of metrics.

F_1 Score

 $ightharpoonup F_1$ is often used as a summary score for binary classifiers instead of Accuracy.

Works better with imbalanced classes.

*F*₁ Score

F₁ is often used as a summary score for binary classifiers instead of Accuracy.

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- Criticised for giving Precision and Recall the same importance.
- ▶ Is not symmetric, ignores true negatives, i.e., is misleading for some cases of imbalanced classes.

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► F₁ is often used as a summary score for binary classifiers instead of Accuracy.

Works better with imbalanced classes.

- Criticised for giving Precision and Recall the same importance.
- Is not symmetric, ignores true negatives, i.e., is misleading for some cases of imbalanced classes.
- Fowlkes-Mallows index is a geometric mean of Precision and Recall (used in clustering).

The geometric mean is between the arithmetic and harmonic mean. For example, the geometric mean of 2/3 and 1/3 is (approx) 0.4714.

More Derived Metrics

Positive predictive value (PPV),	False omission
precision	rate (FOR)
$= \frac{\mathrm{TP}}{\mathrm{PP}} = 1 - \mathrm{FDR}$	$= \frac{FN}{PN} = 1 - NPV$
False discovery rate (FDR) $= \frac{FP}{PP} = 1 - PPV$	Negative predictive value (NPV) $= \frac{TN}{PN} = 1 - FOR$

You can see that the negative predictive value becomes the Precision when we swap the classes (and vice versa).

More Derived Metrics

True positive rate (TPR), recall, sensitivity (SEN), probability of detection, hit rate, power
$$= \frac{TP}{P} = 1 - FNR$$
 False positive rate (FPR), probability of false alarm, fall-out
$$= \frac{FP}{N} = 1 - TNR$$
 True negative rate (TNR), specificity (SPC), selectivity
$$= \frac{TN}{N} = 1 - FPR$$

Note that *specificity* becomes Recall when we swap the classes (and vice versa).

For example, medical doctors communicate in terms of *sensitivity* and *specificity*.

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$$TPR = Sensitivity = Recall = TP/P = 6/8$$

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$$TPR = Sensitivity = Recall = TP/P = 6/8$$

$$TNR = Specificity = TN/N = 3/4$$

How often is negative predicted negative?

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$$TPR = Sensitivity = Recall = TP/P = 6/8$$

$$TNR = Specificity = TN/N = 3/4$$

How often is negative predicted negative?

$$FPR = Prob.$$
 of false $alarm = FP/N = 1/4$

How often is negative predicted positive?

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Total	8 + 4 = 12		

$$TPR = Sensitivity = Recall = TP/P = 6/8$$

$$TNR = Specificity = TN/N = 3/4$$

How often is negative predicted negative?

$$FPR = Prob.$$
 of false $alarm = FP/N = 1/4$

How often is negative predicted positive?

$$FNR = Miss rate = FN/P = 2/8$$

How often is positive predicted negative?

Evaluating Multi-class Classifiers

Assume classification into classes from a finite set C.

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Consider a classification dataset:

$$\{(\vec{x}_k,c_k)\mid k=1,\ldots,p\}$$

Here \vec{x}_k is a vector of attributes/features and $c_k \in C$ for all k.

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How good are the predictions h_1, \ldots, h_p w.r.t. c_1, \ldots, c_p ?

There are many possible metrics ...

Consider an arbitrary (finite) number of classes in C.

Confusion Matrix

Assume that $C = \{1, \dots, m\}$.

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Formally,

$$M_{ij} = |\{k \mid c_k = i \land h_k = j\}|$$

Actual		Р	redict	ed	
	1		j	• • •	m
1	M_{11}		M_{1j}		M_{1m}
:	:		÷		:
i	M_{i1}	• • •	M_{ij}	• • •	M_{im}
:	:		:		:
m	M_{m1}		M_{mj}		M_{mm}

Example

Actual	Predicted
big	big
big	big
small	big
medium	medium
big	small
big	big
small	small
small	small
medium	medium
medium	small
small	small
big	big
medium	small
small	medium
big	big

Example

Actual	Predicted
big	big
big	big
small	big
medium	medium
big	small
big	big
small	small
small	small
medium	medium
medium	small
small	small
big	big
medium	small
small	medium
big	big

Actual	Predicted		
	big medium small		
big	5	0	1
medium	0	2	2
small	1	1	3

Note that the diagonal counts the correctly classified samples.

The off-diagonal elements correspond to misclassified samples.

We can easily generalize Accuracy, Precision, Recall, and F_1 -score from the binary classification to multiple classes.

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- $\begin{array}{l} \blacktriangleright \ M_{i\bullet} = \sum_{j=1}^m M_{ij} \\ \blacktriangleright \ M_{\bullet j} = \sum_{j=1}^m M_{ij} \end{array}$

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- $M_{\bullet \bullet} = \sum_{i=1}^{m} \sum_{j=1}^{m} M_{ij}$

Now, the metrics:

$$\mathsf{Accuracy} = \frac{\sum_{k=1}^{m} M_{kk}}{M_{\bullet \bullet}}$$

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$$Accuracy = \frac{\sum_{k=1}^{m} M_{kk}}{M_{\bullet \bullet}}$$

For a given class $i \in C$:

$$Precision[i] = \frac{M_{ii}}{M_{\bullet i}} \qquad Recall[i] = \frac{M_{ii}}{M_{i\bullet}}$$

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For a given class $i \in C$:

$$Precision[i] = \frac{M_{ii}}{M_{\bullet i}} \qquad Recall[i] = \frac{M_{ii}}{M_{i\bullet}}$$

$$F_1[i] = \frac{2 * \text{Precision}[i] * \text{Recall}[i]}{\text{Precision}[i] + \text{Recall}[i]}$$

Note that Precision, Recall, and F_1 can be defined only for a given class!

Example

Actual	Predicted		
	big medium small		
big	5	0	1
medium	0	2	2
small	1	1	3

Compute the metrics.

Example

Accuracy =
$$(5+2+3)/15 = 0.66$$

Precision[big] = 5/6

 $\mathsf{Precision}[\mathsf{medium}] = 2/3$

Precision[small] = 3/6

Recall[big] = 5/6

Recall[medium] = 2/4

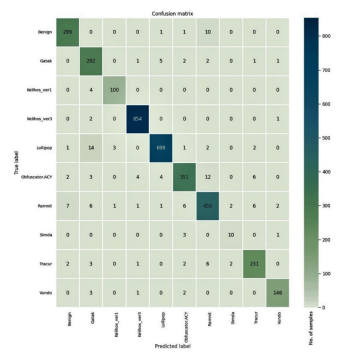
Recall[small] = 3/5

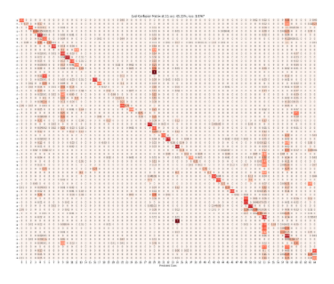
$$F_1[big] = \frac{2*(5/6)*(5/6)}{(5/6)+(5/6)} = 5/6 = 0.83$$

$$F_1[medium] = 0.57$$

$$F_1[medium] = 0.54$$

How do you get a single number out of these? Average Precision, Recall, and F_1 are usually computed, but one needs to be careful about the variance.





Machine learning/data mining is needed to understand the matrix.

Probabilistic Classifier Evaluation

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Assume binary classification into two classes $\{0,1\}$.

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How to interpret the predictions h_1, \ldots, h_p ?

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Let us fix predictions h_1, \ldots, h_p .

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Given a threshold $T \in [0,1]$ we define

$$h_k^T = \begin{cases} 1 & \text{if } h_k \ge T \\ 0 & \text{if } h_k < T \end{cases}$$

For every ${\cal T}$ we can compute all the metrics (Precision, Recall, etc.)

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Given a metric MET and a threshold T, we denote by MET[T] the metric MET evaluated on h_1^T, \ldots, h_p^T .

We obtain

$$\mathsf{TP}[T] = |\{k \mid h_k^T = 1 \land c_k = 1\}|$$

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and

 $\mathsf{TN}[T], \mathsf{FP}[T], \mathsf{FN}[T], \mathsf{Accuracy}[T], \mathsf{Precision}[T], \mathsf{Recall}[T], F_1[T], \dots$

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We obtain

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and

$$\mathsf{TN}[T], \mathsf{FP}[T], \mathsf{FN}[T], \mathsf{Accuracy}[T], \mathsf{Precision}[T], \mathsf{Recall}[T], F_1[T], \dots$$

However, all metrics are now functions of the threshold T.

Index	1	2	3	4	5	6	7	8	9	10	11	12
Actual	1	1	1	1	1	0	0	1	1	0	0	0
Predicted	.98	.95	.9	.86	.66	.48	.43	.42	.36	.15	.1	.05
T=0.5	TP	TP	TP	TP	TP	TN	TN	FN	FN	TN	TN	TN
T=0.42	TP	TP	TP	TP	TP	FP	FP	TP	FN	TN	TN	TN
T=0.1	TP	TP	TP	TP	TP	FP	FP	TP	TP	FP	FP	TN

Index	1	2	3	4	5	6	7	8	9	10	11	12
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Predicted	.98	.95	.9	.86	.66	.48	.43	.42	.36	.15	.1	.05
T=0.5	TP	TP	TP	TP	TP	TN	TN	FN	FN	TN	TN	TN
T=0.42	TP	TP	TP	TP	TP	FP	FP	TP	FN	TN	TN	TN
T=0.1	TP	TP	TP	TP	TP	FP	FP	TP	TP	FP	FP	TN

For example, consider T=0.42, then

Index	1	2	3	4	5	6	7	8	9	10	11	12
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Predicted	.98	.95	.9	.86	.66	.48	.43	.42	.36	.15	.1	.05
T=0.5	TP	TP	TP	TP	TP	TN	TN	FN	FN	TN	TN	TN
T=0.42	TP	TP	TP	TP	TP	FP	FP	TP	FN	TN	TN	TN
T=0.1	TP	TP	TP	TP	TP	FP	FP	TP	TP	FP	FP	TN

For example, consider T=0.42, then

$$\mathsf{TP}[T] = 6 \quad \mathsf{FP}[T] = 2 \quad \mathsf{FN}[T] = 1 \quad \mathsf{TN}[T] = 3$$

Index	1	2	3	4	5	6	7	8	9	10	11	12
Actual	1	1	1	1	1	0	0	1	1	0	0	0
Predicted	.98	.95	.9	.86	.66	.48	.43	.42	.36	.15	.1	.05
T=0.5	TP	TP	TP	TP	TP	TN	TN	FN	FN	TN	TN	TN
T=0.42	TP	TP	TP	TP	TP	FP	FP	TP	FN	TN	TN	TN
T=0.1	TP	TP	TP	TP	TP	FP	FP	TP	TP	FP	FP	TN

For example, consider T = 0.42, then

$$TP[T] = 6$$
 $FP[T] = 2$ $FN[T] = 1$ $TN[T] = 3$

$$\mathsf{Accuracy}[T] = \frac{3+6}{12} \quad \mathsf{Precision}[T] = \frac{6}{6+2} \quad \mathsf{Recall}[T] = \frac{6}{6+1}$$

Index	1	2	3	4	5	6	7	8	9	10	11	12
Actual	1	1	1	1	1	0	0	1	1	0	0	0
Predicted	.98	.95	.9	.86	.66	.48	.43	.42	.36	.15	.1	.05
T=0.5	TP	TP	TP	TP	TP	TN	TN	FN	FN	TN	TN	TN
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Accuracy[
$$T$$
] = $\frac{3+6}{12}$ Precision[T] = $\frac{6}{6+2}$ Recall[T] = $\frac{6}{6+1}$

$$F_1[T] = \frac{2 \cdot 6/8 \cdot 6/7}{6/8 + 6/7} = 0.8$$

Consider two metrics for a given T:

$$TPR[T] = \frac{TP[T]}{P[T]}$$
 (True Positive Rate)

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ROC curve is then a function ROC : $[0,1] \rightarrow [0,1]^2$ defined by

$$ROC(T) = (TPR[T], FPR[T])$$

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Observe that

$$ROC(0) = (1, 1)$$

Because the classifier with T=0 simply classifies everything as positive, i.e., into the class 1.

Both TPR[T] and FPR[T] are non-increasing in T.

Index	1	2	3	4	5	6	7	8	9	10	11	12
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▶ $0.00 \le T \le 0.05$: TPR = 1 and FPR = 1

Index	1	2	3	4	5	6	7	8	9	10	11	12
Actual	1	1	1	1	1	0	0	1	1	0	0	0
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- ▶ $0.00 \le T \le 0.05$: TPR = 1 and FPR = 1
- ▶ $0.05 < T \le 0.10$: TPR = 1 and FPR = 4/5

Index	1	2	3	4	5	6	7	8	9	10	11	12
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- ▶ $0.15 < T \le 0.36$: TPR = 1 and FPR = 2/5

Index	1	2	3	4	5	6	7	8	9	10	11	12
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- ▶ $0.10 < T \le 0.15$: TPR = 1 and FPR = 3/5
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Index	1	2	3	4	5	6	7	8	9	10	11	12
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- ▶ $0.42 < T \le 0.43$: TPR = 5/7 and FPR = 2/5
- ▶ $0.43 < T \le 0.48$: TPR = 5/7 and FPR = 1/5

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- ▶ $0.48 < T \le 0.66$: TPR = 5/7 and FPR = 0

Index	1	2	3	4	5	6	7	8	9	10	11	12
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- ▶ $0.48 < T \le 0.66$: TPR = 5/7 and FPR = 0
- ▶ $0.66 < T \le 0.86$: TPR = 4/7 and FPR = 0

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- ▶ $0.43 < T \le 0.48$: TPR = 5/7 and FPR = 1/5
- ▶ $0.48 < T \le 0.66$: TPR = 5/7 and FPR = 0
- ▶ $0.66 < T \le 0.86$: TPR = 4/7 and FPR = 0
- ▶ $0.86 < T \le 0.90$: TPR = 3/7 and FPR = 0

Index	1	2	3	4	5	6	7	8	9	10	11	12
Actual	1	1	1	1	1	0	0	1	1	0	0	0
Predicted	.98	.95	.9	.86	.66	.48	.43	.42	.36	.15	.1	.05

- ▶ $0.00 \le T \le 0.05$: TPR = 1 and FPR = 1
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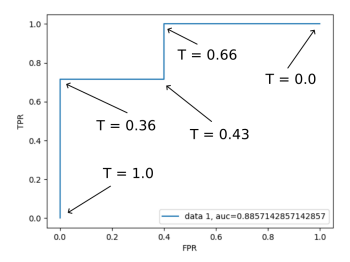
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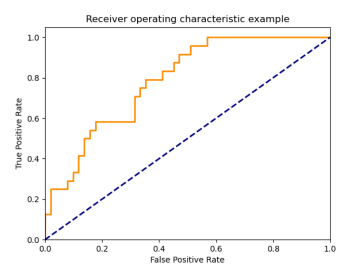
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ROC

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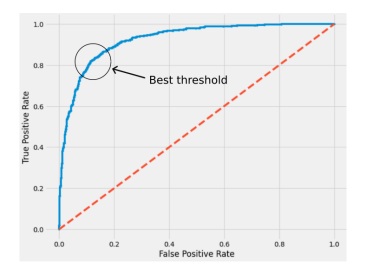


Iris Dataset - A Classifier



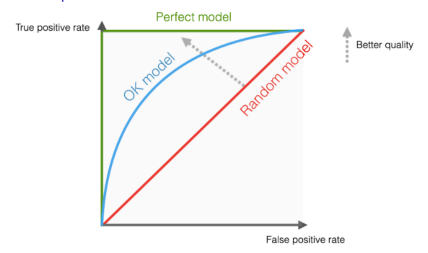
Example from the scikit-learn manual - SVM classifier trained in Iris

Using ROC and Threshold



Search for the best threshold at the elbow of the ROC curve.

ROC - Explanation

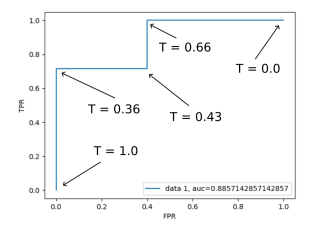


The larger the area under the ROC curve (ROC-AUC), the better.

ROC-AUC ranges from 0 to 1. ROC-AUC \approx 0.5 indicates random guessing.

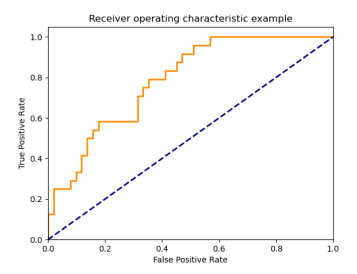
ROC-AUC

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ROC-AUC = 0.8857

Iris - ROC-AUC



ROC-AUC = 0.79

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Consider our cancer detection example:

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The ROC-AUC is the probability of succeeding in the $h_i > h_j$ test.

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 Accuracy, Precision, Recall, F₁
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There are still several questions unanswered:

- When to use the metrics.
- How to estimate the influence of sampling the dataset.

Use of Evaluation Metrics

In our case, the following scenarios are typical:

▶ **Final test**: Evaluate the model on the test set (separated at the beginning of training) and then compute the metrics. May inform the user about the quality of the model.

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There are (at least) two scenarios in which this happens:

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- Hyperparameter fine-tuning.
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Keep in mind that the metrics are artificial, and the results of the model are roughly summarized.

It would be best if you always strived to test the proper functionality of your model in as natural conditions as possible.

For example, a model for medical diagnosis should be evaluated by medical doctors who may observe many features of its behavior that are difficult to express quantitatively.

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We will consider these issues in some later lecture. Concretely,

- ► Bias-variance tradeoff
- Statistical tests for testing
 - significance of the metrics values,
 - paired t-tests for comparing models.

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Thresholding

- ▶ Introduce a threshold $0 \le t \le 1$
- ▶ Demand, one of the two metrics (typically the Recall), to be at least t. That is

$$Recall_1 \ge t$$
 $Recall_2 \ge t$

Compare the values of the other metric numerically. In our case, decide whether

$$Precision_1 \ge Precision_2$$

(Still need to be concerned about the statistical significance.)

Example

Actual condition	Predicted condition					
	Canc.	Non-canc.				
Cancer	6	2				
Non-canc.	1	3				
Total	8 + 4 = 12					

Actual condition	Predicted condition					
	Canc.	Non-canc.				
Cancer	5	3				
Non-canc.	0	4				
Total	8 + 4 = 12					

$$\begin{aligned} &\mathsf{Precision}_1 = \frac{6}{7} & \mathsf{Recall}_1 = \frac{6}{8} \\ &\mathsf{Precision}_2 = \frac{5}{5} = 1 & \mathsf{Recall}_2 = \frac{5}{8} \end{aligned}$$

Consider a threshold t on the Recall.

The second classifier is better if the threshold t is 5/8, then the second classifier is better.

If the threshold t is 6/8, then the second classifier is unacceptable.

Numerical features

Throughout this lecture we assume that all features are numerical, i.e., feature vectors belong to \mathbb{R}^n .

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- Throughout this lecture we assume that all features are numerical, i.e., feature vectors belong to \mathbb{R}^n .
- Most non-numerical features can be conveniently transformed to numerical ones.

For example:

Colors {blue, red, yellow} can be represented by

$$\{(1,0,0),(0,1,0),(0,0,1)\}$$

(one-hot encoding)

- Words can be embedded into vector spaces by various means (word2vec etc.)
- ▶ A black-and-white picture of *x* × *y* pixels can be encoded as a vector of *xy* numbers that capture the shades of gray of the pixels.

(Even though this is not the best way of representing images.)

Basic Problems

We consider two basic problems:

▶ (Binary) classification

Our goal: Classify inputs into two categories.



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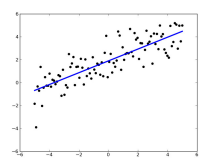
▶ (Binary) classification

Our goal: Classify inputs into two categories.

Regression

Our goal: Find a (hypothesized) functional dependency in data.





Linear Models Binary Classification

Binary classification in \mathbb{R}^n

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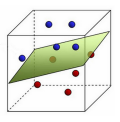
Comments:

- ▶ In practice, we often do not strictly demand $h(\vec{x}) = c$ for all training examples $(\vec{x}, c) \in D$ (often it is impossible)
- ▶ We are more interested in good generalization, that is how well h classifies new instances that do not belong to D. (Recall that we usually evaluate accuracy of the resulting hypothesized function h on a test set.)

Models

We consider two kinds of hypothesis spaces:

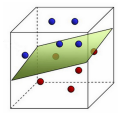
Linear (affine) classifiers (this lecture)



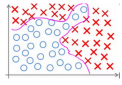
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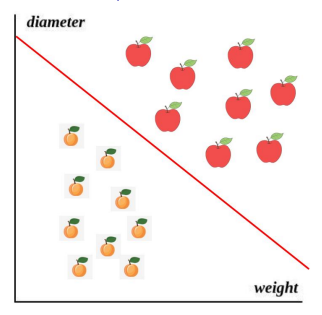
► Linear (affine) classifiers (this lecture)



Non-linear classifiers (kernel SVM, neural networks) (later lectures)



Linear Classifier – Example



Length and Scalar Product of Vectors

▶ We consider vectors $\vec{x} = (x_1, ..., x_n) \in \mathbb{R}^m$.

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- ► Euclidean metric on vectors: $||\vec{x}|| = \sqrt{\sum_{i=1}^{n} x_i^2}$ The distance between two vectors (points) \vec{x}, \vec{y} is $||\vec{x} - \vec{y}||$.

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- Scalar product $\vec{x} \cdot \vec{y}$ of vectors $\vec{x} = (x_1, \dots, x_n)$ and $\vec{y} = (y_1, \dots, y_n)$ defined by

$$\vec{x} \cdot \vec{y} = \sum_{i=1}^{n} x_i y_i$$

- Recall that $\vec{x} \cdot \vec{y} = ||\vec{x}|| \, ||\vec{y}|| \cos \theta$ where θ is the angle between \vec{x} and \vec{y} . That is $\vec{x} \cdot \vec{y}$ is the length of the projection of \vec{y} on \vec{x} multiplied by $||\vec{x}||$.
- Note that $\vec{x} \cdot \vec{x} = ||\vec{x}||^2$

Linear Classifier

A *linear classifier* $h[\vec{w}]$ is determined by a vector of *weights* $\vec{w} = (w_0, w_1, \dots, w_n) \in \mathbb{R}^{n+1}$ as follows:

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Given
$$\vec{x} = (x_1, \dots, x_n) \in \mathbb{R}^n$$
,

$$h[\vec{w}](\vec{x}) := \begin{cases} 1 & w_0 + \sum_{i=1}^n w_i \cdot x_i \ge 0 \\ 0 & w_0 + \sum_{i=1}^n w_i \cdot x_i < 0 \end{cases}$$

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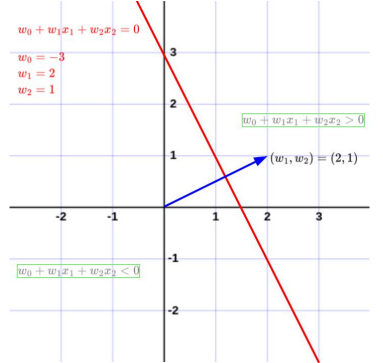
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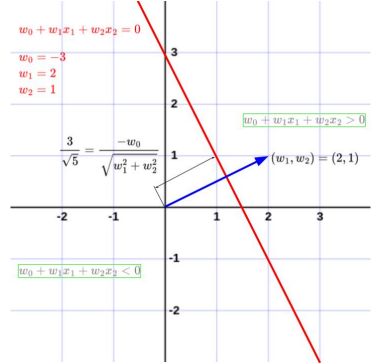
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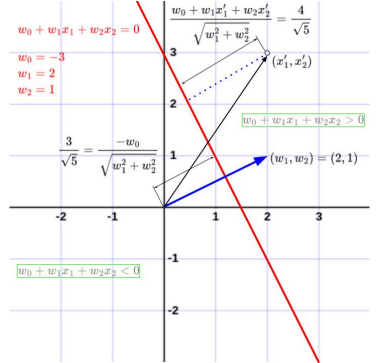
More succinctly:

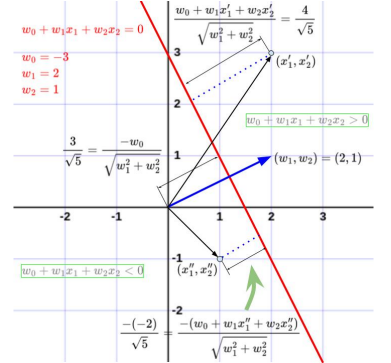
$$h(\vec{x}) = sgn\left(w_0 + \sum_{i=1}^n w_i \cdot x_i\right)$$
 where $sgn(y) = \begin{cases} 1 & y \ge 0 \\ 0 & y < 0 \end{cases}$

We define separating hyperplane determined by \vec{w} as the set of all $\vec{x} \in \mathbb{R}^n$ satisfying $w_0 + \sum_{i=1}^n w_i \cdot x_i = 0$.

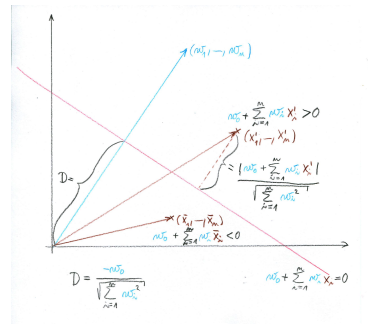








Linear Classifier - Geometry



Linear Classifier – Notation

Given
$$\vec{x} = (x_1, \dots, x_n) \in \mathbb{R}^n$$
 we define an *augmented feature vector*

$$\mathbf{\tilde{x}} = (x_0, x_1, \dots, x_n)$$
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Linear Classifier - Notation

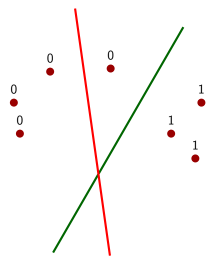
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This makes the notation for the linear classifier more succinct:

$$h[\vec{w}](\vec{x}) = sgn(\vec{w} \cdot \tilde{\mathbf{x}})$$

Linear Classifier – Learning



- classification in the plane using a linear classifier
- ▶ if a point is incorrectly classified, the learning algorithm turns the line (hyperplane) to improve the classification

► Given a training set

$$D = \{ \left(\vec{x}_1, c_1\right), \left(\vec{x}_2, c_2\right)\right), \dots, \left(\vec{x}_p, c_p\right)) \}$$
 Here $\vec{x}_k = (x_{k1} \dots, x_{kn}) \in \mathbb{R}^n$ and $c_k \in \{0, 1\}$.

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Recall that
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 Here $\vec{x}_k = \left(x_{k1}, \dots, x_{kn}\right) \in \mathbb{R}^n$ and $c_k \in \{0, 1\}$. Recall that $\tilde{\mathbf{x}}_k = \left(x_{k0}, x_{k1}, \dots, x_{kn}\right)$ where $x_{k0} = 1$.

▶ A weight vector $\vec{\mathbf{w}} \in \mathbb{R}^{n+1}$ is **consistent with** D if

$$h[\vec{w}](\vec{x}_k) = sgn(\vec{w} \cdot \tilde{\mathbf{x}}_k) = c_k$$
 for all $k = 1, ..., p$

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▶ Our goal is to find a consistent \vec{w} assuming that D is linearly separable.

Online learning algorithm:

Idea: Cyclically go through the training examples in *D* and adapt weights. Whenever an example is incorrectly classified, turn the hyperplane so that the example becomes closer to its correct half-space.

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Here $k = (t \mod p) + 1$, i.e., the examples are considered cyclically, and $0 < \varepsilon \le 1$ is a **learning rate**.

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Theorem (Rosenblatt)

If D is linearly separable, then there is t^* such that $\vec{w}^{(t^*)}$ is consistent with D.

Example

Training set:

$$D = \{((2,-1),1),((2,1),1),((1,3),0)\}$$

That is

$$\vec{x}_1 = (2,-1)$$
 $\vec{x}_1 = (1,2,-1)$ $\vec{x}_2 = (2,1)$ $\vec{x}_3 = (1,3)$ $\vec{x}_3 = (1,1,3)$

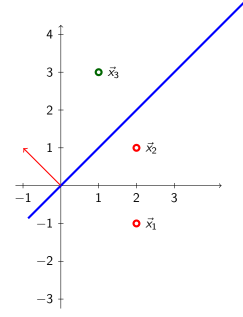
$$c_1 = 1$$

$$c_2 = 1$$

$$c_3 = 0$$

Assume that the initial vector $\vec{w}^{(0)}$ is $\vec{w}^{(0)} = (0, -1, 1)$. Consider $\varepsilon = 1$.

Example: Separating by $\vec{w}^{(0)}$



Denoting $\vec{w}^{(0)} = (w_0, w_1, w_2) = (0, -1, 1)$ the blue separating line is given by $w_0 + w_1x_1 + w_2x_2 = 0$.

The red vector normal to the blue line is (w_1, w_2) .

The points on the side of (w_1, w_2) are assigned 1 by the classifier, the others zero. (In this case \vec{x}_3 is assigned one and \vec{x}_1, \vec{x}_2 are assigned zero, all of this is inconsistent with $c_1=1, c_2=1, c_3=0$.)

Example: Computing $\vec{w}^{(1)}$

We have

$$\vec{\mathbf{w}}^{(0)} \cdot \tilde{\mathbf{x}}_1 = (0, -1, 1) \cdot (1, 2, -1) = 0 - 2 - 1 = -3$$

thus

$$sgn\left(ec{w}^{\left(0
ight) }\cdot \mathbf{ ilde{x}}_{1}
ight) =0$$

and thus

$$sgn\left(\vec{w}^{(0)}\cdot\mathbf{\tilde{x}}_1\right)-c_1=0-1=-1$$

(I.e., $\vec{x_1}$ is not correctly classified, and $\vec{w}^{(0)}$ is not consistent with D.) Hence.

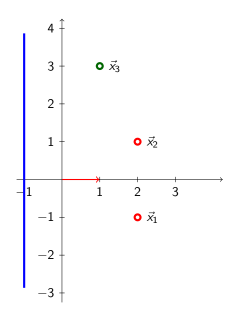
$$\vec{w}^{(1)} = \vec{w}^{(0)} - \left(sgn\left(\vec{w}^{(0)} \cdot \tilde{\mathbf{x}}_1\right) - c_1\right) \cdot \tilde{\mathbf{x}}_1$$

$$= \vec{w}^{(0)} + \tilde{\mathbf{x}}_1$$

$$= (0, -1, 1) + (1, 2, -1)$$

$$= (1, 1, 0)$$

Example: Separating by $\vec{w}^{(1)}$



Example: Computing $\vec{w}^{(2)}$

We have

$$\vec{w}^{(1)} \cdot \tilde{\mathbf{x}}_2 = (1, 1, 0) \cdot (1, 2, 1) = 1 + 2 = 3$$

thus

$$sgn\left(ec{w}^{(1)}\cdot\mathbf{ ilde{x}}_{2}
ight)=1$$

and thus

$$sgn\left(\vec{w}^{(1)}\cdot\tilde{\mathbf{x}}_{2}\right)-c_{2}=1-1=0$$

(I.e., $\vec{x_2}$ is currently correctly classified by $\vec{w}^{(1)}$. However, as we will see, $\vec{x_3}$ is not well classified.)

Hence,

$$\vec{w}^{(2)} = \vec{w}^{(1)} = (1, 1, 0)$$

Example: Computing $\vec{w}^{(3)}$

We have

$$\vec{\mathbf{w}}^{(2)} \cdot \tilde{\mathbf{x}}_3 = (1, 1, 0) \cdot (1, 1, 3) = 1 + 1 = 2$$

thus

$$sgn\left(ec{w}^{(2)}\cdot\mathbf{ ilde{x}}_{3}
ight)=1$$

and thus

$$sgn\left(\vec{w}^{(2)}\cdot\tilde{\mathbf{x}}_3\right)-c_3=1-0=1$$

(This means that \vec{x}_3 is not well classified, and $\vec{w}^{(2)}$ is not consistent with D.) Hence,

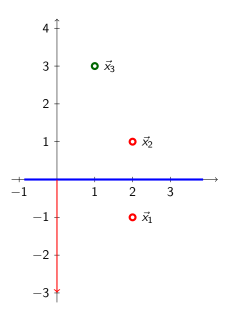
$$\vec{w}^{(3)} = \vec{w}^{(2)} - \left(sgn\left(\vec{w}^{(2)} \cdot \tilde{\mathbf{x}}_3\right) - c_3\right) \cdot \tilde{\mathbf{x}}_3$$

$$= \vec{w}^{(2)} - \tilde{\mathbf{x}}_3$$

$$= (1, 1, 0) - (1, 1, 3)$$

$$= (0, 0, -3)$$

Example: Separating by $\vec{w}^{(3)}$



Example: Computing $\vec{w}^{(4)}$

We have

$$\vec{w}^{(3)} \cdot \tilde{\mathbf{x}}_1 = (0,0,-3) \cdot (1,2,-1) = 3$$

thus

$$sgn\left(ec{w}^{(3)}\cdot\mathbf{\tilde{x}}_{1}
ight)=1$$

and thus

$$sgn\left(\vec{w}^{(3)}\cdot\mathbf{\tilde{x}}_1\right)-c_1=1-1=0$$

(I.e., $\vec{x_1}$ is currently correctly classified by $\vec{w}^{(3)}$. However, we shall see that $\vec{x_2}$ is not.)

Hence,

$$\vec{w}^{(4)} = \vec{w}^{(3)} = (0, 0, -3)$$

Example: Computing $\vec{w}^{(5)}$

We have

$$\vec{\mathbf{w}}^{(4)} \cdot \tilde{\mathbf{x}}_2 = (0, 0, -3) \cdot (1, 2, 1) = -3$$

thus

$$sgn\left(\vec{w}^{(4)}\cdot\tilde{\mathbf{x}}_{2}\right)=0$$

and thus

$$sgn\left(\vec{w}^{(4)}\cdot\mathbf{\tilde{x}}_{2}\right)-c_{2}=0-1=-1$$

(I.e., $\vec{x_2}$ is not correctly classified, and $\vec{w}^{(4)}$ is not consistent with D.) Hence.

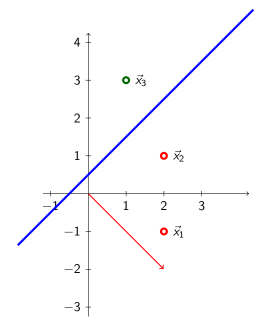
$$\vec{w}^{(5)} = \vec{w}^{(4)} - \left(sgn\left(\vec{w}^{(4)} \cdot \tilde{\mathbf{x}}_2\right) - c_2\right) \cdot \tilde{\mathbf{x}}_2$$

$$= \vec{w}^{(4)} + \tilde{\mathbf{x}}_2$$

$$= (0, 0, -3) + (1, 2, 1)$$

$$= (1, 2, -2)$$

Example: Separating by $\vec{w}^{(5)}$



Example: The result

The vector $\vec{w}^{(5)}$ is consistent with D:

$$\begin{split} sgn\left(\vec{w}^{(5)} \cdot \tilde{\mathbf{x}}_1\right) &= sgn\left((1,2,-2) \cdot (1,2,-1)\right) = sgn(7) = 1 = c_1 \\ sgn\left(\vec{w}^{(5)} \cdot \tilde{\mathbf{x}}_2\right) &= sgn\left((1,2,-2) \cdot (1,2,1)\right) = sgn(3) = 1 = c_2 \\ sgn\left(\vec{w}^{(5)} \cdot \tilde{\mathbf{x}}_3\right) &= sgn\left((1,2,-2) \cdot (1,1,3)\right) = sgn(-3) = 0 = c_3 \end{split}$$

Batch learning algorithm:

Compute a sequence of weight vectors $\vec{w}^{(0)}, \vec{w}^{(1)}, \vec{w}^{(2)}, \ldots$

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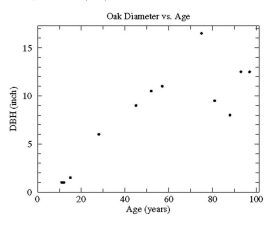
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Linear Regression - Oaks in Wisconsin

This example is from How to Lie with Statistics by Darrell Huff (1954)

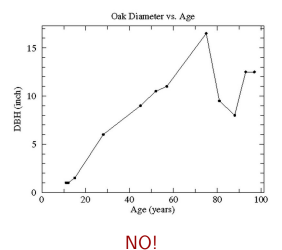
Age	DBH
(years)	(inch)
97	12.5
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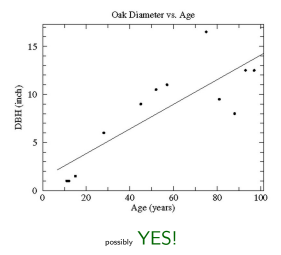
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In what follows we use the squared error defined by

$$E = \frac{1}{2} \sum_{(\vec{x}, f) \in D} (h(\vec{x}) - f)^2$$

Our goal is to minimize E.

The main reason is that this function has nice mathematical properties (as opposed, e.g., to $\sum_{(\vec{x},f)\in D}|h(\vec{x})-f|$).

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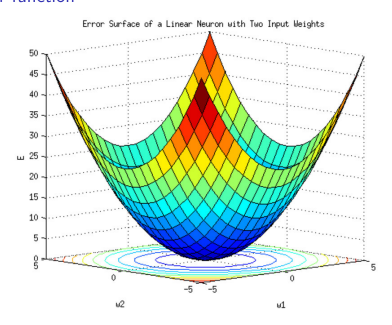
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Squared Error Function:

$$E(\vec{w}) = \frac{1}{2} \sum_{k=1}^{p} (\vec{w} \cdot \tilde{\mathbf{x}}_{k} - f_{k})^{2} = \frac{1}{2} \sum_{k=1}^{p} \left(\sum_{i=0}^{n} w_{i} x_{ki} - f_{k} \right)^{2}$$

Error function



Consider the **gradient** of the error function:

$$\nabla E(\vec{w}) = \left(\frac{\partial E}{\partial w_0}(\vec{w}), \dots, \frac{\partial E}{\partial w_n}(\vec{w})\right) = \sum_{k=1}^p (\vec{w} \cdot \tilde{\mathbf{x}}_k - f_k) \cdot \tilde{\mathbf{x}}_k$$

What is the gradient $\nabla E(\vec{w})$? It is a vector in \mathbb{R}^{n+1} which points in the direction of the steepest *ascent* of E (its length corresponds to the steepness). Note that here the vectors $\tilde{\mathbf{x}}_k$ are *fixed* parameters of E!

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Fact:

If
$$\nabla E(\vec{w}) = \vec{0} = (0, \dots, 0)$$
, then \vec{w} is a global minimum of E .

This follows from the fact that E is a convex paraboloid that has a unique extreme, which is a minimum.



Consider n = 1, which means that $\vec{w} = (w_0, w_1)$ and we write x instead of \vec{x} since $\vec{x} \in \mathbb{R}^n = \mathbb{R}^1 = \mathbb{R}$.

Then the model is $h[\vec{w}](x) = w_0 + w_1 \cdot x$.

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= \{(x₁, f₁), (x₂, f₂), (x₃, f₃)\}

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$$\frac{\partial E}{\partial w_0} = (w_0 + w_1 \cdot 2 - 1) \cdot 1 + (w_0 + w_1 \cdot 3 - 2) \cdot 1 + (w_0 + w_1 \cdot 4 - 5) \cdot 1$$

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$$\frac{\partial E}{\partial w_1} = (w_0 + w_1 \cdot 2 - 1) \cdot 2 + (w_0 + w_1 \cdot 3 - 2) \cdot 3 + (w_0 + w_1 \cdot 4 - 5) \cdot 4$$

Consider n=1, which means that $\vec{w}=(w_0,w_1)$ and we write x instead of \vec{x} since $\vec{x} \in \mathbb{R}^n = \mathbb{R}^1 = \mathbb{R}$.

Then the model is $h[\vec{w}](x) = w_0 + w_1 \cdot x$.

Consider a concrete training set:

$$\mathcal{T} = \{(2,1), (3,2), (4,5)\}$$

= \{(x₁, f₁), (x₂, f₂), (x₃, f₃)\}

$$E(w_0, w_1) = \frac{1}{2}[(w_0 + w_1 \cdot 2 - 1)^2 + (w_0 + w_1 \cdot 3 - 2)^2 + (w_0 + w_1 \cdot 4 - 5)^2]$$

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$$\nabla E(\vec{w}) = (\frac{\partial E}{\partial w_0}, \frac{\partial E}{\partial w_1}) = (w_0 + w_1 \cdot 2 - 1) \cdot (1, 2) + (w_0 + w_1 \cdot 3 - 2) \cdot (1, 3) + (w_0 + w_1 \cdot 4 - 5) \cdot (1, 4)$$

Gradient Descent:

▶ Weights $\vec{w}^{(0)}$ are initialized randomly close to $\vec{0}$.

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Here $0 < \varepsilon \le 1$ is a learning rate.

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Proposition

For sufficiently small $\varepsilon > 0$ the sequence $\vec{w}^{(0)}, \vec{w}^{(1)}, \vec{w}^{(2)}, \dots$ converges (component-wisely) to the global minimum of E.

Training set:

$$D = \{(x_1, f_1), (x_2, f_2), (x_3, f_3)\} = \{(0, 0), (2, 1), (2, 2)\}$$

Note that input vectors are one dimensional, so we write them as numbers. That is

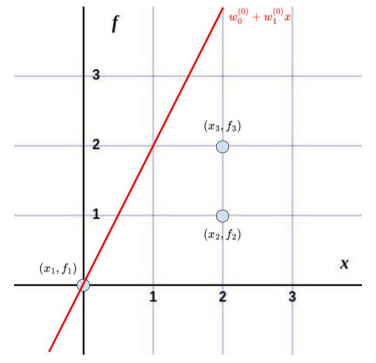
$$x_1 = 0$$

 $x_2 = 2$
 $x_3 = 2$
 $\mathbf{\tilde{x}}_1 = (1,0)$
 $\mathbf{\tilde{x}}_2 = (1,2)$
 $\mathbf{\tilde{x}}_3 = (1,2)$

$$f_1 = 0$$

 $f_2 = 1$
 $f_3 = 2$

Assume that the initial vector $\vec{w}^{(0)}$ is $\vec{w}^{(0)} = (w_0^{(0)}, w_1^{(0)}) = (0, 2)$. Consider $\varepsilon = \frac{1}{10}$.



Training set:

 $D = \{(x_1, f_1), (x_2, f_2), (x_3, f_3)\} = \{(0, 0), (2, 1), (2, 2)\}$ Augmented input vectors: $\tilde{\mathbf{x}}_1 = (1, 0), \ \tilde{\mathbf{x}}_2 = (1, 2), \ \tilde{\mathbf{x}}_1 = (1, 2)$

$$\nabla E(\vec{w}) = \left(\frac{\partial E}{\partial w_0}(\vec{w}), \frac{\partial E}{\partial w_1}(\vec{w})\right) = (w_0 + w_1 \cdot x_1 - f_1) \cdot \tilde{\mathbf{x}}_1 + (w_0 + w_1 \cdot x_2 - f_2) \cdot \tilde{\mathbf{x}}_2 + (w_0 + w_1 \cdot x_3 - f_3) \cdot \tilde{\mathbf{x}}_3$$

For $\vec{w}^{(0)} = (0,2)$ we have

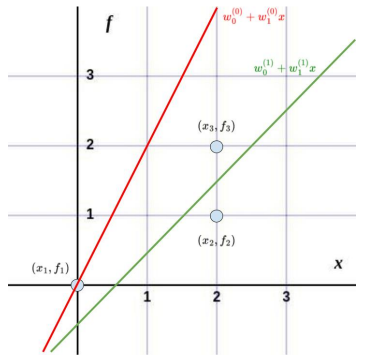
$$\nabla E(\vec{w}^{(0)}) = (0 + 2 \cdot 0 - 0) \cdot (1, 0)$$

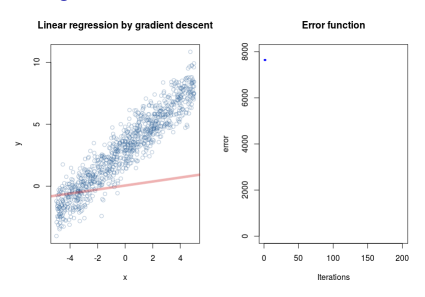
$$+ (0 + 2 \cdot 2 - 1) \cdot (1, 2)$$

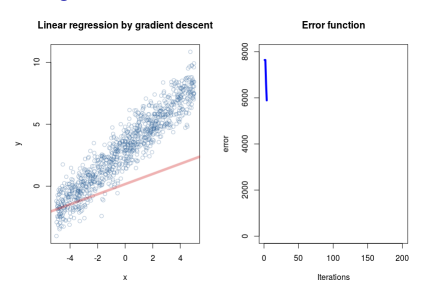
$$+ (0 + 2 \cdot 2 - 2) \cdot (1, 2) = (3, 6) + (2, 4) = (5, 10)$$

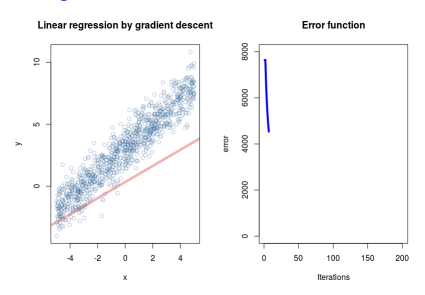
Finally, $\vec{w}^{(1)}$ is computed by

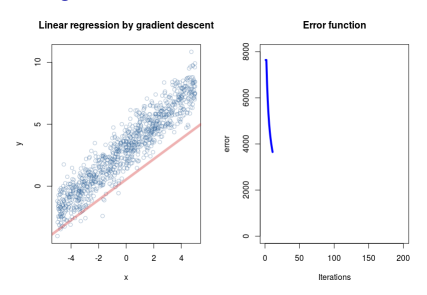
$$ec{w}^{(1)} = ec{w}^{(0)} - arepsilon \cdot
abla E(ec{w}^{(0)}) = (0,2) - rac{1}{10} \cdot (5,10) = (-1/2,1)$$

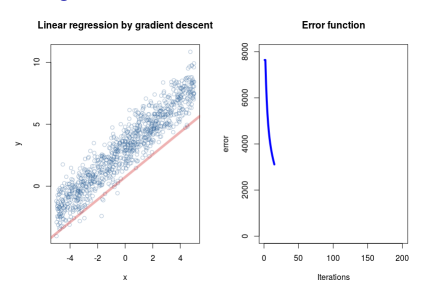


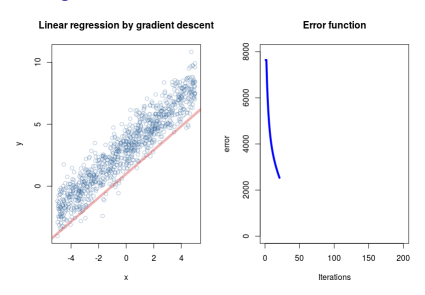


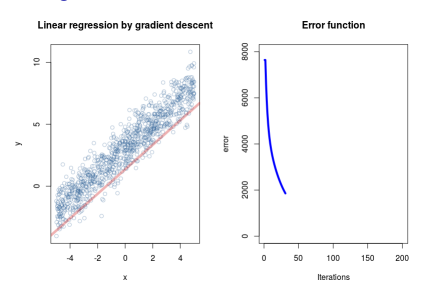


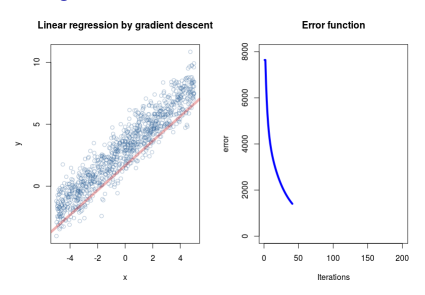


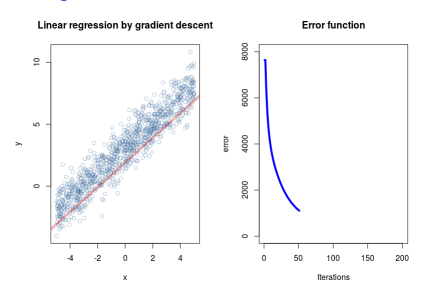


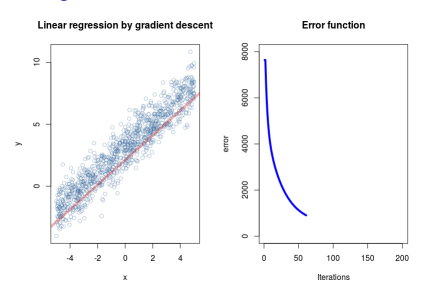


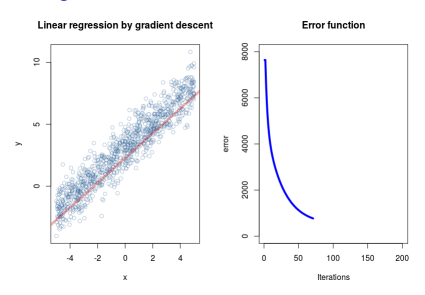


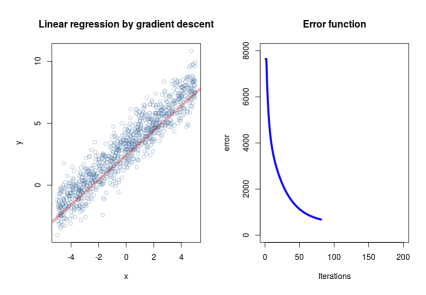


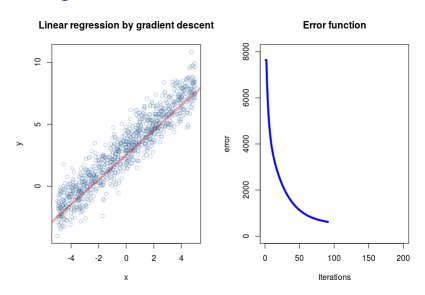


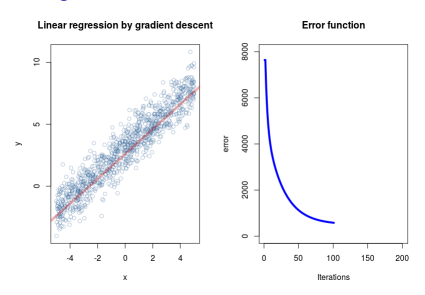


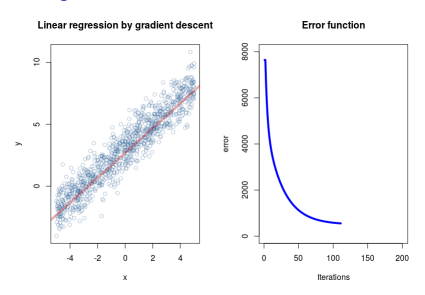


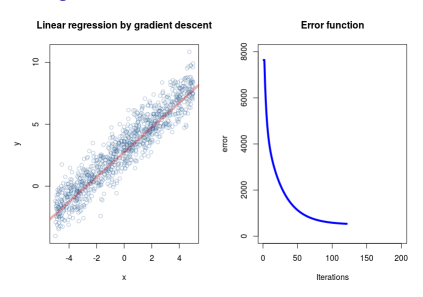


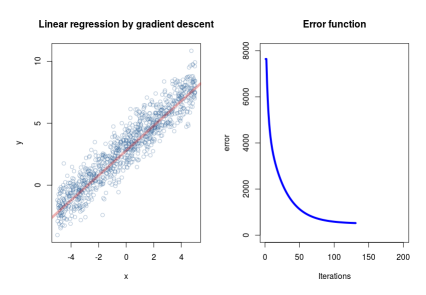


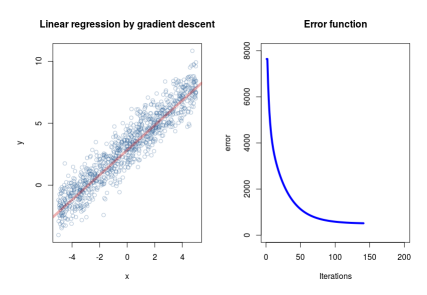


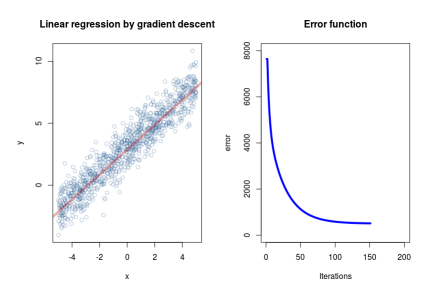


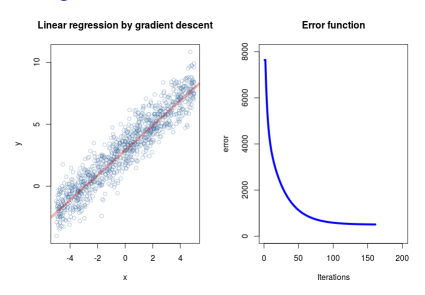


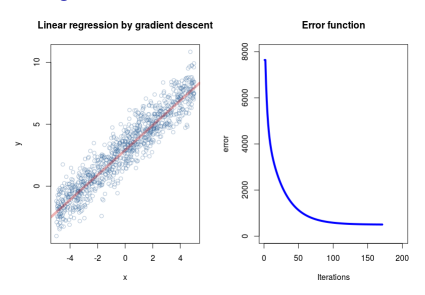


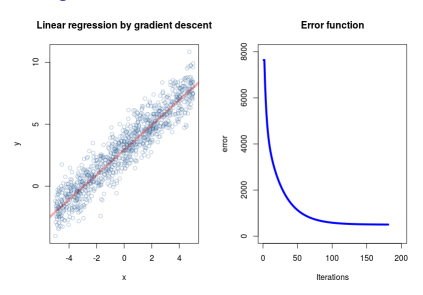


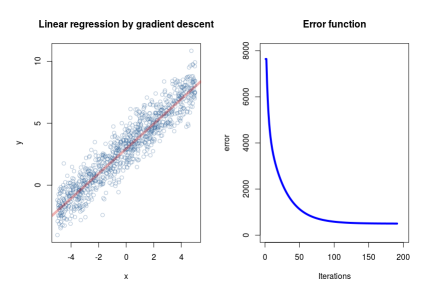


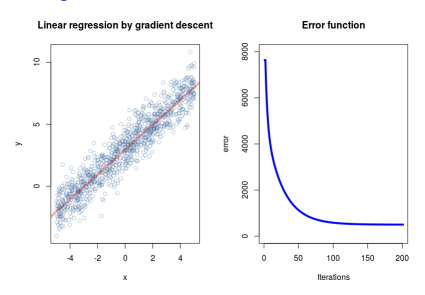












Finding the Minimum in Dimension One

Assume n = 1. Then, the error function E is

$$E(w_0, w_1) = \frac{1}{2} \sum_{k=1}^{p} (w_0 + w_1 x_k - f_k)^2$$

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Minimize E w.r.t. w_0 a w_1 :

$$\frac{\partial E}{\partial w_0} = 0 \quad \Leftrightarrow \quad w_0 = \bar{f} - w_1 \bar{x} \quad \Leftrightarrow \quad \bar{f} = w_0 + w_1 \bar{x}$$

where
$$\bar{x} = \frac{1}{p} \sum_{k=1}^{p} x_k$$
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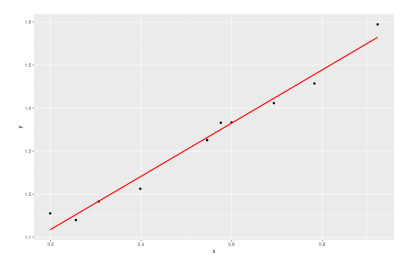
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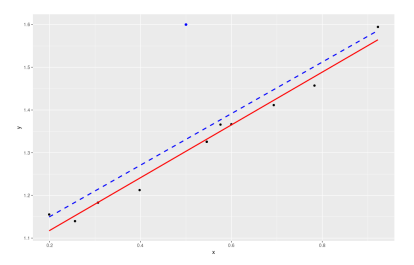
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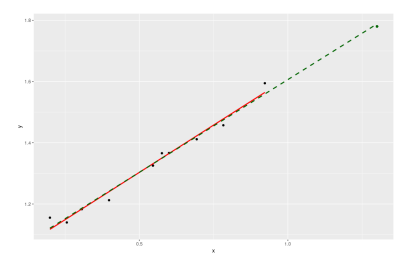
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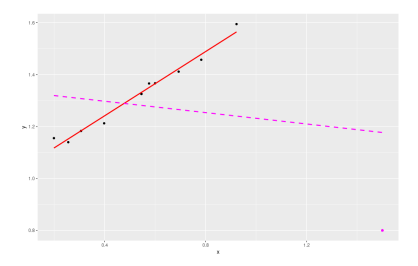
$$\frac{\partial E}{\partial w_1} = 0 \quad \Leftrightarrow \quad w_1 = \frac{\frac{1}{p} \sum_{k=1}^p (f_k - f)(x_k - \bar{x})}{\frac{1}{p} \sum_{k=1}^p (x_k - \bar{x})^2}$$

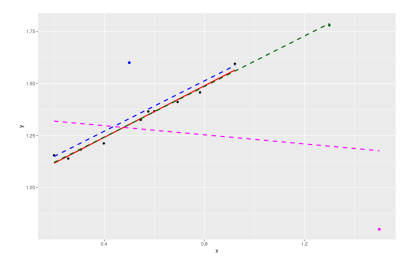
i.e.
$$w_1 = cov(f, x)/var(x)$$









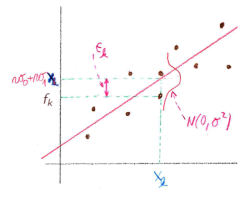


Maximum Likelihood vs Least Squares (Dim 1)

Fix a training set $D = \{(x_1, f_1), (x_2, f_2), \dots, (x_p, f_p)\}$ Assume that each f_k has been generated randomly by

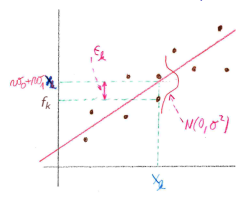
$$f_k = (\mathbf{w_0} + \mathbf{w_1} \cdot \mathbf{x_k}) + \epsilon_k$$

where w_0 , w_1 are **unknown weights**, and ϵ_k are independent, normally distributed noise values with mean 0 and some variance σ^2



How "probable" is it to generate the correct f_1, \ldots, f_p ?

Maximum Likelihood vs Least Squares (Dim 1)



How "probable" is it to generate the correct f_1, \ldots, f_p ?

The following conditions are equivalent:

- \triangleright w_0 , w_1 minimize the squared error E
- ▶ w_0 , w_1 maximize the likelihood (i.e., the "probability") of generating the correct values f_1, \ldots, f_p using $f_k = (w_0 + w_1 \cdot x_k) + \epsilon_k$

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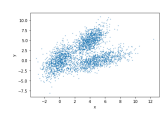
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- ► Linear models are less likely to overfit (low variance) the training data but sometimes tend to underfit (high bias).
- Linear models are prone to outliers.

Unsupervised Learning

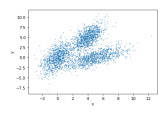
Clustering

Often data form clusters based on some notion of similarity.



Clustering

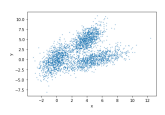
Often data form clusters based on some notion of similarity.



This means that the data distribution is *multimodal*, i.e., contains several regions of higher probability mass.

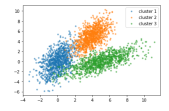
Clustering

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We aim to group data into clusters of "similar" examples without using any additional information. (no supervision).

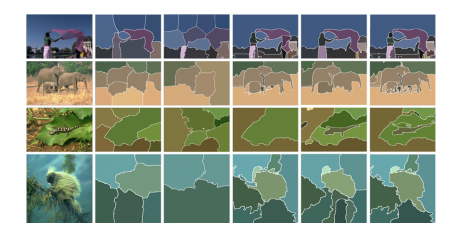


Motivation

Clustering is useful, e.g., in

- Customer segmentation based on their purchases.
- Data exploration identify patterns in data
- Semi-supervised learning cluster labeled examples with the unlabeled ones
- Search engines searching for images similar to a given image
- Image segmentation
- **.**..

Segmentation



Consider a dataset

$$D = \{\vec{x}_1, \dots, \vec{x}_p\}$$

Note that no target class/value is provided.

Clustering is a partition $\mathcal{U} = \{U_1, \dots, U_K\}$ of D into K clusters.

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For concreteness:

- We stick with numerical features, which means that the dataset $D = \{\vec{x}_1, \dots, \vec{x}_p\}$ contains vectors $\vec{x}_i \in \mathbb{R}^n$.
- Assume the Euclidean distance d.

Note that clustering may be based on completely different similarity/dissimilarity measures and non-numerical data.

K-Means Clustering

The K-means clustering model consists of

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- An assignment $q_{ij} \in \{0,1\}$ for $i=1,\ldots,p$ and $j=1,\ldots,K$ of inputs $\vec{x_i}$ to clusters U_j so that

$$\sum_j q_{ij} = 1$$
 for $i = 1, \dots, p$

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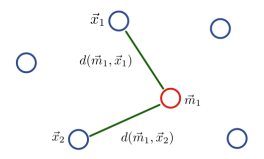
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How good is a given model?

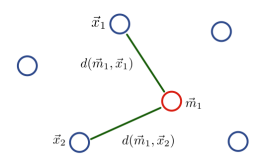
Error Function

Measure the distance of inputs $\vec{x_i}$ to their cluster prototypes $\vec{m_j}$



Error Function

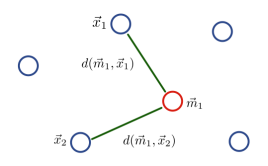
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$$E(\{q_{ij}\}, \{\vec{m}_j\}) = \sum_{i=1}^{p} \sum_{j=1}^{K} q_{ij} d(\vec{x}_i, \vec{m}_j)^2$$

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We aim to *minimize* this error, i.e., to find proper positions of cluster prototypes and their assignment to minimize the total squared distance of examples to their prototypes.

The Problem: Minimize

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Note that

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- ▶ If we fix $\{q_{ij}\}$, we can minimize $E(\{q_{ij}\}, \{\vec{m}_j\})$ by letting each \vec{m}_j to minimize the total squared distance to its prototypes:

$$\sum_{i} q_{ij} d(\vec{x}_i, \vec{m}_j)^2$$

The Problem: Minimize

$$E(\{q_{ij}\}, \{\vec{m}_j\}) = \sum_{i=1}^{p} \sum_{j=1}^{K} q_{ij} d(\vec{x}_i, \vec{m}_j)^2 \text{ w.r.t. } \{q_{ij}\}, \{\vec{m}_j\}$$

Note that

- If we fix $\{\vec{m}_j\}$, we can minimize $E(\{q_{ij}\}, \{\vec{m}_j\})$ by setting $q_{ij} = 1$ iff \vec{m}_j is the *closest prototype* to \vec{x}_i .
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$$\sum_{i} q_{ij} d(\vec{x}_i, \vec{m}_j)^2$$

This is achieved by putting each prototype \vec{m}_j into the centroid of all inputs it represents:

$$\vec{m}_j = \frac{1}{\sum_{i=1}^p q_{ij}} \sum_{i=1}^p q_{ij} \vec{x}_i$$

Note that $\sum_{i=1}^{p} q_{ij}$ is the size of the cluster represented by \vec{m}_{i} .

Algorithm 1 K-means clustering

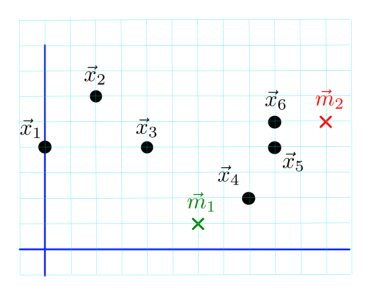
- 1: Initialize K cluster centers $\vec{m}_1, \vec{m}_2, \dots, \vec{m}_K$ randomly
- 2: repeat
- 3: **for** each data point $\vec{x_i}$ **do**
- 4: Assign $\vec{x_i}$ to the nearest centroid, i.e., set $q_{ij} = 1$ for

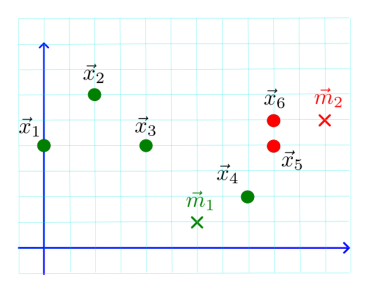
$$j = \arg\min_{j} d(\vec{x}_i, \vec{m}_j)^2$$

- 5: end for
- 6: **for** each cluster prototype \vec{m}_i **do**
- 7: Update \vec{m}_j to be the centroid of all points assigned to it

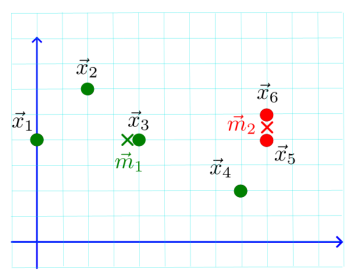
$$\vec{m}_{j} = \frac{1}{\sum_{i=1}^{p} q_{ij}} \sum_{i=1}^{p} q_{ij} \vec{x}_{i}$$

- 8: end for
- 9: until convergence

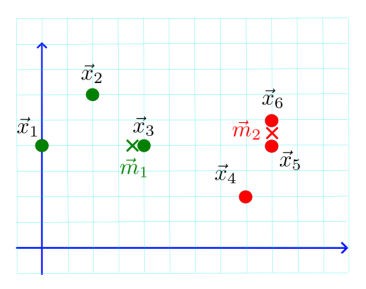




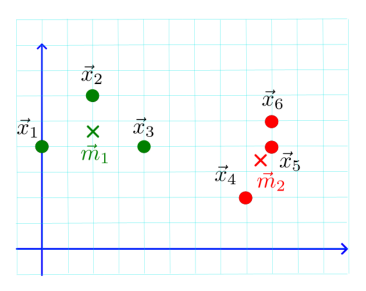
Lines 3-5: Assign examples to the prototypes.



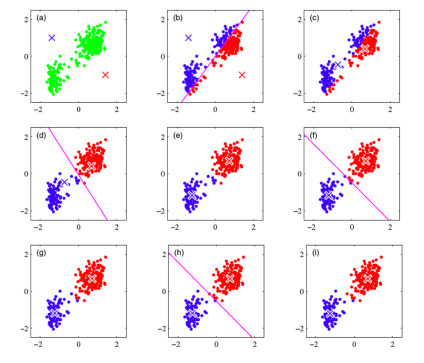
Lines 6-8: Move the prototypes to the centroids of their examples.



Lines 3-5: Assign examples to the prototypes.



Lines 6-8: Move the prototypes to the centroids of their examples.



Convergence of K-means Clustering

Every step of K-means reduces the error $E(\{q_{ij}\}, \{\vec{m}_j\})$:

- ▶ We always assign an input vector to the closest prototype.
- ► We always move the prototype to be "closest" to the input vectors it represents.

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Convergence can be tested by computing the error and checking whether it has not changed in the last step.

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There are only finitely many possible assignments to q_{ij} , and we always minimize the distance of inputs to their assigned centers.

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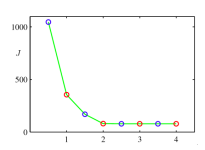
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Convergence can be tested by computing the error and checking whether it has not changed in the last step.

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There are only finitely many possible assignments to q_{ij} , and we always minimize the distance of inputs to their assigned centers.

Example error development during training. Blue circles mean reassignment, and red circles mean moving prototypes.



Setting K - the Elbow Method

K-means clustering minimizes the *inertia* measure:

$$E(\{q_{ij}\},\{\vec{m}_j\}) = \sum_{i=1}^{p} \sum_{j=1}^{k} q_{ij} d(\vec{x}_i,\vec{m}_j)^2$$

That is the sum of squared distances of all examples of D to the cluster prototypes.

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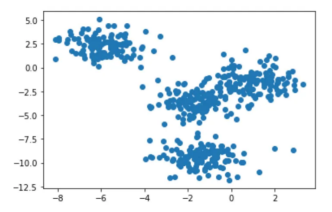
Note that the error does not consider the distance between the centers of the clusters.

Still, it is a valid measure that can be used to select the number of clusters.

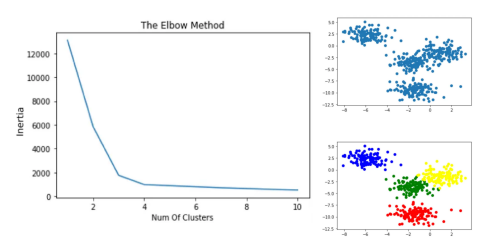
Elbow Method

The following method for setting up the hyperparameters can be used in general. Let us illustrate the elbow method on K-means clustering with the inertia measure.

Consider the following data:



Elbow Method



We could choose four clusters because adding more leads only to small decrements in the inertia.

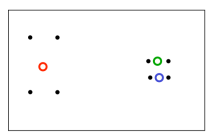
Bad Behavior

Minimizing $E(\{q_{ij}\}, \{\vec{m_j}\})$ starting from random positions of prototypes does not always produce "nice" results.

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Minimizing $E(\{q_{ij}\}, \{\vec{m_j}\})$ starting from random positions of prototypes does not always produce "nice" results.

Some runs correspond to apparently bad solutions to the clustering problem even though a better solution exists.



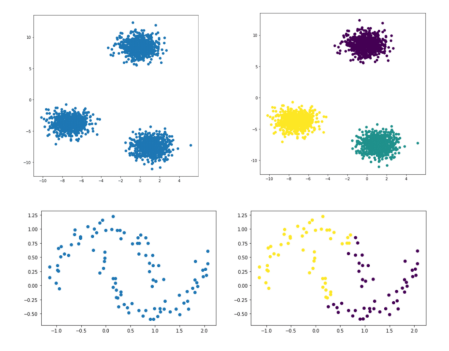
Possible solution: Start the algorithm several times with random initialization of the prototypes.

- Prototype initialization is a big issue in K-means. There are various strategies. For example:
 - Start with all centers in a single corner.
 - Include randomness in the setting of centers throughout the algorithm.
 - Initialize sequentially, always fit prototypes, and then choose a new one as far away from the others as possible.
 - ► Use hierarchical clustering (next slides) to find clusters and initialize *K*-means with their centroids.

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- As the squared error is behind the basic method, outliers may strongly affect its behavior (as in the linear regression case).
- ▶ Other problematic properties of data include
 - non-convex clusters
 - clusters of different sizes
 - non-linearly separable clusters
 - overlapping clusters



Consider a dataset

$$D = \{\vec{x}_1, \dots, \vec{x}_p\}$$

Here $\vec{x_i} \in \mathbb{R}^n$ for all i = 1, ..., p. Assume a distance d (e.g., Euclidean).

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Idea:

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More concretely:

Maintain a set of clusters

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Idea:

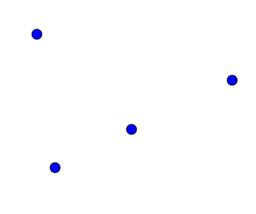
- Start by merging the closest examples (w.r.t. d)
- ▶ Incrementally build *larger clusters* by merging smaller clusters.

More concretely:

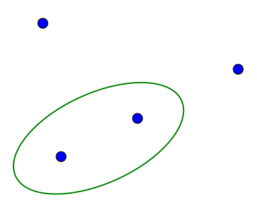
- Maintain a set of clusters
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How do we determine the closest clusters?

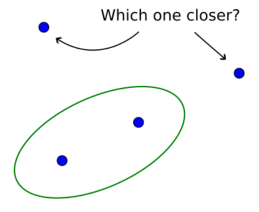
Closest Clusters

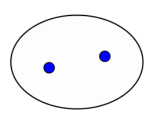


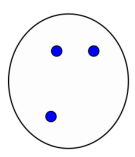
Closest Clusters

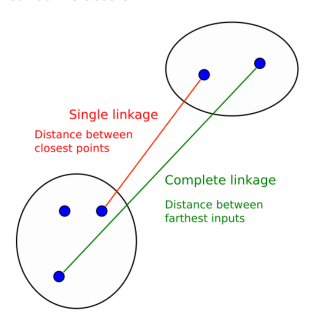


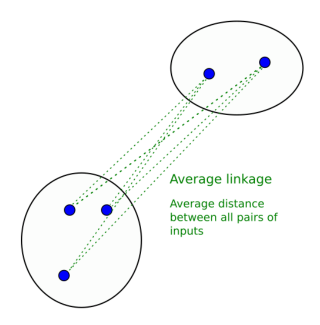
Closest Clusters



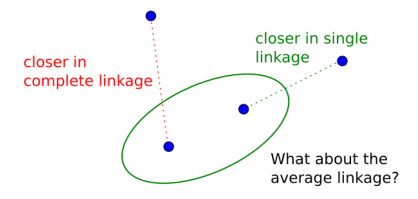








Which One is Closer?



Consider two clusters $U_j, U_k \subseteq D$.

single_linkage
$$(U_j, U_k)$$

= min $\{d(\vec{x}, \vec{z}) \mid \vec{x} \in U_j, \vec{z} \in U_k\}$

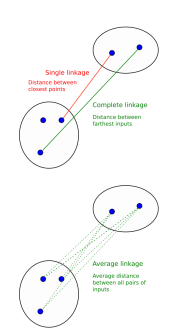
complete_linkage
$$(U_j, U_k)$$

= max $\{d(\vec{x}, \vec{z}) \mid \vec{x} \in U_j, \vec{z} \in U_k\}$

average_linkage(
$$U_j, U_k$$
)

$$= \frac{1}{|U_j||U_k|} \sum_{\vec{x} \in U_j} \sum_{\vec{z} \in U_k} d(\vec{x}, \vec{z})$$

Each linkage can result in a different clustering.



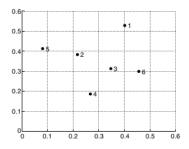
Agglomerative Hierarchical Clustering Algorithm

Maintain a set of clusters
Initially, each $\vec{x_i}$ in its own cluster
repeat
Pick two closest clusters

Using the distance measure *d* and single, average, or complete linkage. Merge them into a new cluster

until only one cluster is left

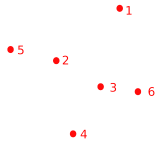
Example



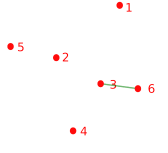
Point	x Coordinate	y Coordinate
p1	0.40	0.53
p2	0.22	0.38
р3	0.35	0.32
p4	0.26	0.19
p5	0.08	0.41
p6	0.45	0.30

	p1	p2	р3	p4	p5	p6
p1	0.00	0.24	0.22	0.37	0.34	0.23
p2	0.24	0.00	0.15	0.20	0.14	0.25
р3	0.22	0.15	0.00	0.16	0.28	0.11
p4	0.37	0.20	0.16	0.00	0.29	0.22
p5	0.34	0.14	0.28	0.29	0.00	0.39
p6	0.23	0.25	0.11	0.22	0.39	0.00

	p1	p2	р3	p4	p5	p6
p1	0.00	0.24	0.22	0.37	0.34	0.23
p2	0.24	0.00	0.15	0.20	0.14	0.25
p3	0.22	0.15	0.00	0.16	0.28	0.11
p4	0.37	0.20	0.16	0.00	0.29	0.22
p5	0.34	0.14	0.28	0.29	0.00	0.39
p6	0.23	0.25	0.11	0.22	0.39	0.00



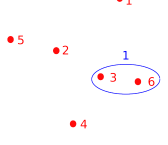
	p1	p2	p3	p4	p_5	p6
p1	0.00	0.24	0.22	0.37	0.34	0.23
p2	0.24	0.00	0.15	0.20	0.14	0.25
р3	0.22	0.15	0.00	0.16	0.28	0.11
p4	0.37	0.20	0.16	0.00	0.29	0.22
p5	0.34	0.14	0.28	0.29	0.00	0.39
p6	0.23	0.25	0.11	0.22	0.39	0.00



$$d(3,6) = 0.11$$

which is the minimum distance between points.

	p1	p2	р3	p4	p5	p6
p1	0.00	0.24	0.22	0.37	0.34	0.23
p2	0.24	0.00	0.15	0.20	0.14	0.25
p3	0.22	0.15	0.00	0.16	0.28	0.11
p4	0.37	0.20	0.16	0.00	0.29	0.22
p5	0.34	0.14	0.28	0.29	0.00	0.39
p6	0.23	0.25	0.11	0.22	0.39	0.00



	p1	p2	р3	p4	p5	p6
p1	0.00	0.24	0.22	0.37	0.34	0.23
p2	0.24	0.00	0.15	0.20	0.14	0.25
р3	0.22	0.15	0.00	0.16	0.28	0.11
p4	0.37	0.20	0.16	0.00	0.29	0.22
p5	0.34	0.14	0.28	0.29	0.00	0.39
р6	0.23	0.25	0.11	0.22	0.39	0.00

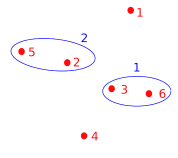
• 5 2 1 0 3 • 6 0 0 4

• 1

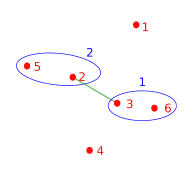
$$d(2,5) = 0.14$$

which is the second smallest distance.

	p1	p2	р3	p4	p5	p6
p1	0.00	0.24	0.22	0.37	0.34	0.23
p2	0.24	0.00	0.15	0.20	0.14	0.25
p3	0.22	0.15	0.00	0.16	0.28	0.11
p4	0.37	0.20	0.16	0.00	0.29	0.22
p5	0.34	0.14	0.28	0.29	0.00	0.39
p6	0.23	0.25	0.11	0.22	0.39	0.00



	p1	p2	р3	p4	p5	p6
p1	0.00	0.24	0.22	0.37	0.34	0.23
p2	0.24	0.00	0.15	0.20	0.14	0.25
р3	0.22	0.15	0.00	0.16	0.28	0.11
p4	0.37	0.20	0.16	0.00	0.29	0.22
p5	0.34	0.14	0.28	0.29	0.00	0.39
р6	0.23	0.25	0.11	0.22	0.39	0.00

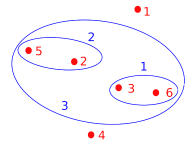


$$d(2,3) = 0.15 =$$

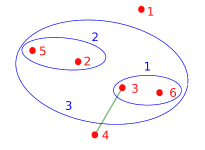
 $min\{d(2,3), d(2,6), d(5,3), d(5,6)\}$

which is smaller than d(1,2)=0.24, d(1,3)=0.22, d(4,2)=0.2, d(4,3)=0.16, d(4,1)=0.37 the min. distances of points in all other pairs of clusters.

	p1	p2	р3	p4	p5	p6
p1	0.00	0.24	0.22	0.37	0.34	0.23
p2	0.24	0.00	0.15	0.20	0.14	0.25
р3	0.22	0.15	0.00	0.16	0.28	0.11
p4	0.37	0.20	0.16	0.00	0.29	0.22
p5	0.34	0.14	0.28	0.29	0.00	0.39
p6	0.23	0.25	0.11	0.22	0.39	0.00



	p1	p2	р3	p4	p5	p6
p1	0.00	0.24	0.22	0.37	0.34	0.23
p2	0.24	0.00	0.15	0.20	0.14	0.25
р3	0.22	0.15	0.00	0.16	0.28	0.11
p4	0.37	0.20	0.16	0.00	0.29	0.22
p5	0.34	0.14	0.28	0.29	0.00	0.39
p6	0.23	0.25	0.11	0.22	0.39	0.00

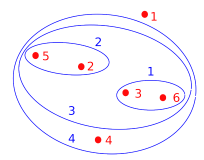


$$d(4,3) = 0.15$$

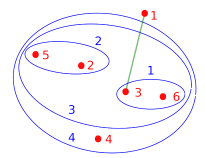
= min{d(4,3), d(4,5), d(4,2), d(4,6)}

which is smaller than d(1,3) = 0.22, the distance of 1 to the cluster 3.

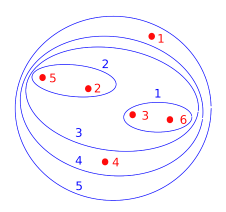
	p1	p2	р3	p4	p5	p6
p1	0.00	0.24	0.22	0.37	0.34	0.23
p2	0.24	0.00	0.15	0.20	0.14	0.25
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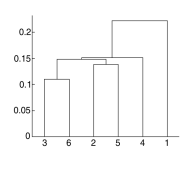


	p1	p2	р3	p4	p5	p6
p1	0.00	0.24	0.22	0.37	0.34	0.23
p2	0.24	0.00	0.15	0.20	0.14	0.25
р3	0.22	0.15	0.00	0.16	0.28	0.11
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p5	0.34	0.14	0.28	0.29	0.00	0.39
p6	0.23	0.25	0.11	0.22	0.39	0.00



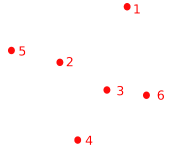
	p1	p2	р3	p4	p_5	p6
p1	0.00	0.24	0.22	0.37	0.34	0.23
p2	0.24	0.00	0.15	0.20	0.14	0.25
p3	0.22	0.15	0.00	0.16	0.28	0.11
p4	0.37	0.20	0.16	0.00	0.29	0.22
p5	0.34	0.14	0.28	0.29	0.00	0.39
p6	0.23	0.25	0.11	0.22	0.39	0.00





Example - Average Linkage

	p1	p2	р3	p4	p5	p6
p1	0.00	0.24	0.22	0.37	0.34	0.23
p2	0.24	0.00	0.15	0.20	0.14	0.25
р3	0.22	0.15	0.00	0.16	0.28	0.11
p4	0.37	0.20	0.16	0.00	0.29	0.22
p5	0.34	0.14	0.28	0.29	0.00	0.39
p6	0.23	0.25	0.11	0.22	0.39	0.00



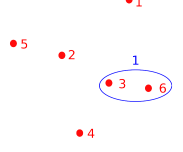
Example - Average Linkage

	p1	p2	р3	p4	p5	p6
p1	0.00	0.24	0.22	0.37	0.34	0.23
p2	0.24	0.00	0.15	0.20	0.14	0.25
р3	0.22	0.15	0.00	0.16	0.28	0.11
p4	0.37	0.20	0.16	0.00	0.29	0.22
p5	0.34	0.14	0.28	0.29	0.00	0.39
p6	0.23	0.25	0.11	0.22	0.39	0.00

• 1 • 5 • 2 • -3 • 6

Example - Average Linkage

	p1	p2	р3	p4	p5	p6
p1	0.00	0.24	0.22	0.37	0.34	0.23
p2	0.24	0.00	0.15	0.20	0.14	0.25
р3	0.22	0.15	0.00	0.16	0.28	0.11
p4	0.37	0.20	0.16	0.00	0.29	0.22
p5	0.34	0.14	0.28	0.29	0.00	0.39
p6	0.23	0.25	0.11	0.22	0.39	0.00

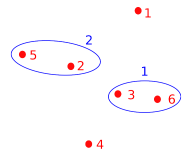


	p1	p2	р3	p4	p_5	p6
p1	0.00	0.24	0.22	0.37	0.34	0.23
p2	0.24	0.00	0.15	0.20	0.14	0.25
р3	0.22	0.15	0.00	0.16	0.28	0.11
p4	0.37	0.20	0.16	0.00	0.29	0.22
p5	0.34	0.14	0.28	0.29	0.00	0.39
p6	0.23	0.25	0.11	0.22	0.39	0.00

$$d(2,5) = 0.14$$

which is second smallest distance.

	p1	p2	р3	p4	p5	p6
p1	0.00	0.24	0.22	0.37	0.34	0.23
p2	0.24	0.00	0.15	0.20	0.14	0.25
p3	0.22	0.15	0.00	0.16	0.28	0.11
p4	0.37	0.20	0.16	0.00	0.29	0.22
p5	0.34	0.14	0.28	0.29	0.00	0.39
p6	0.23	0.25	0.11	0.22	0.39	0.00



	p1	p2	р3	p4	p5	p6
p1	0.00	0.24	0.22	0.37	0.34	0.23
p2	0.24	0.00	0.15	0.20	0.14	0.25
р3	0.22	0.15	0.00	0.16	0.28	0.11
p4	0.37	0.20	0.16	0.00	0.29	0.22
p5	0.34	0.14	0.28	0.29	0.00	0.39
p6	0.23	0.25	0.11	0.22	0.39	0.00

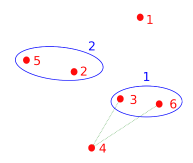
The average distance between 4 and both points of $\{3,6\}$ is

$$\frac{1}{2}(d(4,3)+d(4,6))=0.19$$

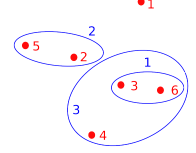
which is smaller than the average distance between all points of clusters 1, 2:

$$\frac{d(5,2)+d(5,3)+d(2,3)+d(2,6)}{4}$$

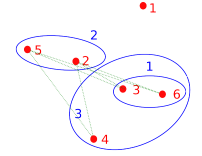
(equal to 0.205), and the average distance of 1 to any cluster.



	p1	p2	р3	p4	p5	p6
p1	0.00	0.24	0.22	0.37	0.34	0.23
p2	0.24	0.00	0.15	0.20	0.14	0.25
р3	0.22	0.15	0.00	0.16	0.28	0.11
p4	0.37	0.20	0.16	0.00	0.29	0.22
p5	0.34	0.14	0.28	0.29	0.00	0.39
p6	0.23	0.25	0.11	0.22	0.39	0.00

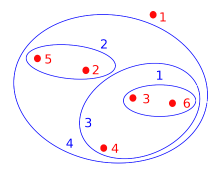


	p1	p2	р3	p4	p5	p6
p1	0.00	0.24	0.22	0.37	0.34	0.23
p2	0.24	0.00	0.15	0.20	0.14	0.25
p3	0.22	0.15	0.00	0.16	0.28	0.11
p4	0.37	0.20	0.16	0.00	0.29	0.22
p5	0.34	0.14	0.28	0.29	0.00	0.39
p6	0.23	0.25	0.11	0.22	0.39	0.00

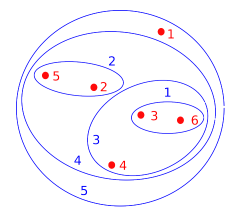


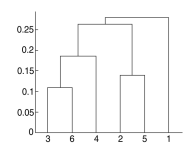
The average distance between clusters 2, 3 is 0.26 which is smaller than the average distance of 1 to any of the two clusters 1, 2 (the average distances are 0.273 and 0.29).

	p1	p2	р3	p4	p5	p6
p1	0.00	0.24	0.22	0.37	0.34	0.23
p2	0.24	0.00	0.15	0.20	0.14	0.25
р3	0.22	0.15	0.00	0.16	0.28	0.11
p4	0.37	0.20	0.16	0.00	0.29	0.22
p5	0.34	0.14	0.28	0.29	0.00	0.39
p6	0.23	0.25	0.11	0.22	0.39	0.00



	p1	p2	р3	p4	p5	p6
p1	0.00	0.24	0.22	0.37	0.34	0.23
p2	0.24	0.00	0.15	0.20	0.14	0.25
р3	0.22	0.15	0.00	0.16	0.28	0.11
p4	0.37	0.20	0.16	0.00	0.29	0.22
p5	0.34	0.14	0.28	0.29	0.00	0.39
p6	0.23	0.25	0.11	0.22	0.39	0.00





	p1	p2	р3	p4	p5	p6
p1	0.00	0.24	0.22	0.37	0.34	0.23
p2	0.24	0.00	0.15	0.20	0.14	0.25
р3	0.22	0.15	0.00	0.16	0.28	0.11
p4	0.37	0.20	0.16	0.00	0.29	0.22
p5	0.34	0.14	0.28	0.29	0.00	0.39
p6	0.23	0.25	0.11	0.22	0.39	0.00

• 1

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• 3 • 6

• 4

	p1	p2	р3	p4	p5	p6
p1	0.00	0.24	0.22	0.37	0.34	0.23
p2	0.24	0.00	0.15	0.20	0.14	0.25
р3	0.22	0.15	0.00	0.16	0.28	0.11
p4	0.37	0.20	0.16	0.00	0.29	0.22
p5	0.34	0.14	0.28	0.29	0.00	0.39
p6	0.23	0.25	0.11	0.22	0.39	0.00

•5 •2 •3•6

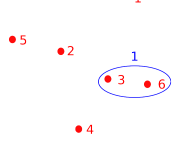
• 4

• 1

$$d(3,6) = 0.11$$

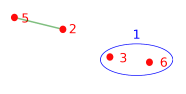
which is the minimum distance between points.

	p1	p2	р3	p4	p_5	p6
p1	0.00	0.24	0.22	0.37	0.34	0.23
p2	0.24	0.00	0.15	0.20	0.14	0.25
р3	0.22	0.15	0.00	0.16	0.28	0.11
p4	0.37	0.20	0.16	0.00	0.29	0.22
p5	0.34	0.14	0.28	0.29	0.00	0.39
p6	0.23	0.25	0.11	0.22	0.39	0.00



	p1	p2	р3	p4	p5	p6
p1	0.00	0.24	0.22	0.37	0.34	0.23
p2	0.24	0.00	0.15	0.20	0.14	0.25
р3	0.22	0.15	0.00	0.16	0.28	0.11
p4	0.37	0.20	0.16	0.00	0.29	0.22
p5	0.34	0.14	0.28	0.29	0.00	0.39
p6	0.23	0.25	0.11	0.22	0.39	0.00

• 1

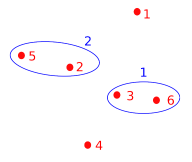


$$d(2,5) = 0.14$$

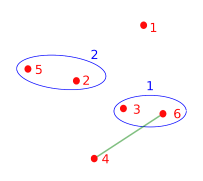
which is the second smallest distance.

• 4

	p1	p2	р3	p4	p5	p6
p1	0.00	0.24	0.22	0.37	0.34	0.23
p2	0.24	0.00	0.15	0.20	0.14	0.25
р3	0.22	0.15	0.00	0.16	0.28	0.11
p4	0.37	0.20	0.16	0.00	0.29	0.22
p5	0.34	0.14	0.28	0.29	0.00	0.39
p6	0.23	0.25	0.11	0.22	0.39	0.00



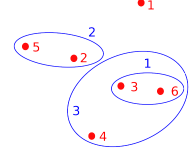
	p1	p2	p3	p4	p5	p6
p1	0.00	0.24	0.22	0.37	0.34	0.23
p2	0.24	0.00	0.15	0.20	0.14	0.25
р3	0.22	0.15	0.00	0.16	0.28	0.11
p4	0.37	0.20	0.16	0.00	0.29	0.22
p5	0.34	0.14	0.28	0.29	0.00	0.39
p6	0.23	0.25	0.11	0.22	0.39	0.00



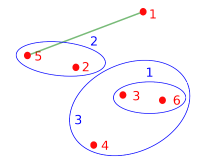
$$d(4,6) = 0.22 =$$
 $\max\{d(4,3), d(4,6)\}$

which is smaller than d(4,5) = 0.29, d(1,5) = 0.34, d(1,6) = 0.23, d(5,6) = 0.39, d(4,1) = 0.37 the max distances of points in all other pairs of clusters.

	p1	p2	р3	p4	p5	p6
p1	0.00	0.24	0.22	0.37	0.34	0.23
p2	0.24	0.00	0.15	0.20	0.14	0.25
р3	0.22	0.15	0.00	0.16	0.28	0.11
p4	0.37	0.20	0.16	0.00	0.29	0.22
p5	0.34	0.14	0.28	0.29	0.00	0.39
p6	0.23	0.25	0.11	0.22	0.39	0.00



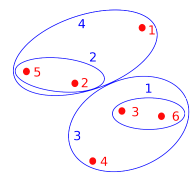
	p1	p2	р3	p4	p5	p6
p1	0.00	0.24	0.22	0.37	0.34	0.23
p2	0.24	0.00	0.15	0.20	0.14	0.25
p3	0.22	0.15	0.00	0.16	0.28	0.11
p4	0.37	0.20	0.16	0.00	0.29	0.22
p5	0.34	0.14	0.28	0.29	0.00	0.39
p6	0.23	0.25	0.11	0.22	0.39	0.00



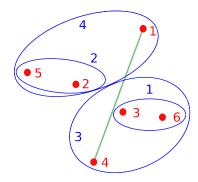
$$d(1,5) = 0.34$$

which is smaller than d(1,4) = 0.37, d(5,6) = 0.39, which are the maximum distances of points in all other pairs of clusters.

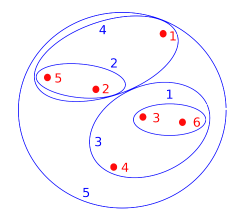
	p1	p2	р3	p4	p5	p6
p1	0.00	0.24	0.22	0.37	0.34	0.23
p2	0.24	0.00	0.15	0.20	0.14	0.25
р3	0.22	0.15	0.00	0.16	0.28	0.11
p4	0.37	0.20	0.16	0.00	0.29	0.22
p5	0.34	0.14	0.28	0.29	0.00	0.39
р6	0.23	0.25	0.11	0.22	0.39	0.00

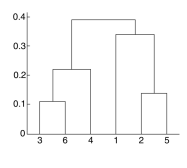


	p1	p2	р3	p4	p5	p6
p1	0.00	0.24	0.22	0.37	0.34	0.23
p2	0.24	0.00	0.15	0.20	0.14	0.25
р3	0.22	0.15	0.00	0.16	0.28	0.11
p4	0.37	0.20	0.16	0.00	0.29	0.22
p5	0.34	0.14	0.28	0.29	0.00	0.39
р6	0.23	0.25	0.11	0.22	0.39	0.00



	p1	p2	р3	p4	p5	p6
p1	0.00	0.24	0.22	0.37	0.34	0.23
p2	0.24	0.00	0.15	0.20	0.14	0.25
р3	0.22	0.15	0.00	0.16	0.28	0.11
p4	0.37	0.20	0.16	0.00	0.29	0.22
p5	0.34	0.14	0.28	0.29	0.00	0.39
p6	0.23	0.25	0.11	0.22	0.39	0.00



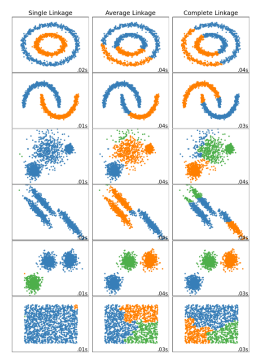


Properties of Agglomerative Hierarchical Clustering

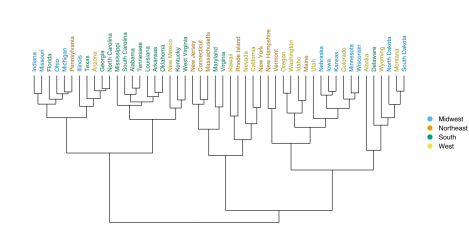
- Provides hierarchy of clusters different cut levels provide different levels of coarseness of clusters
- ► Compared with *k*-means, it does not depend on the initialization and may provide better clusters than *k*-means.

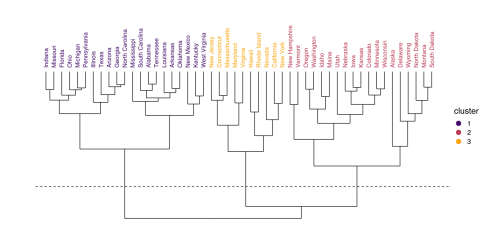
Properties of Agglomerative Hierarchical Clustering

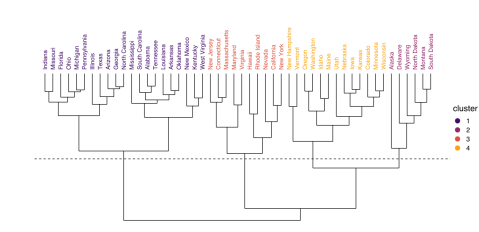
- Provides hierarchy of clusters different cut levels provide different levels of coarseness of clusters
- Compared with k-means, it does not depend on the initialization and may provide better clusters than k-means.
- Lack of global objective function
 - ► The agglomerative hierarchical clustering uses local criteria to decide which clusters to merge.
- Agglomerative clustering has a "rich get richer" behavior that leads to uneven cluster sizes
- Merging decision cannot be undone bad for noisy data
- Computationally expensive.

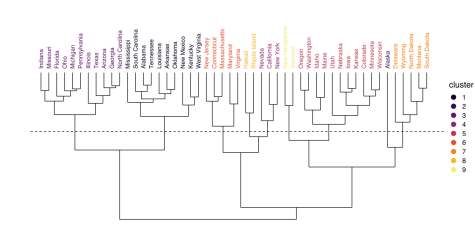


```
# A tibble: 50 × 20
   state
            homeo...¹ multi...² income med_i...³ poverty fed_s...⁴ smoke murder robbery
   <chr>
              <fdb>>
                      <dbl>
                             <dbl>
                                     <dbl>
                                            <dbl>
                                                    <dbl> <dbl> <dbl>
                                                                         <fdb>>
1 Alabama
               71.1
                      15.5
                             22984
                                     42081
                                            17.1
                                                    11.7
                                                           24.8
                                                                   8.2
                                                                        141.
2 Alaska
               64.7
                       24.6
                             30726
                                     66521
                                            9.5
                                                    16.8
                                                           25
                                                                   4.8
                                                                         80.9
3 Arizona
               67.4
                       20.7
                             25680
                                     50448
                                            15.3
                                                    9.85
                                                           20.4
                                                                   7.5
                                                                        144.
4 Arkansas
              67.7 15.2
                             21274
                                     39267
                                             18
                                                     9.61
                                                           23.5
                                                                   6.7
                                                                        91.1
 5 Californ...
               57.4
                       30.7
                             29188
                                     60883
                                             13.7
                                                     8.89
                                                          15.2
                                                                   6.9
                                                                        176.
6 Colorado
              67.6
                       25.6
                             30151
                                     56456
                                             12.2
                                                    9.15
                                                           19.9
                                                                   3.7
                                                                         84.6
7 Connecti...
               69.2
                      34.6
                             36775
                                     67740
                                             9.2
                                                    14.8
                                                           16.5
                                                                   2.9
                                                                        113
 8 Delaware
               73.6
                      17.7
                             29007
                                     57599
                                             11
                                                     8.89
                                                           20.7
                                                                   4.4
                                                                         155.
 9 Florida
               69.7
                       30
                             26551
                                     47661
                                             13.8 9.62
                                                           21.6
                                                                   5
                                                                         169.
10 Georgia
               67.2
                      20.5 25134
                                    49347
                                             15.7 8.88 22.2
                                                                   6.2
                                                                         155.
 ... with 40 more rows, 10 more variables: agg_assault <dbl>, larceny <dbl>,
   motor theft <dbl>. soc sec <dbl>. nuclear <dbl>. coal <dbl>.
#
   tr deaths <dbl>. tr deaths no alc <dbl>. unempl <dbl>. popdens2010 <dbl>.
    and abbreviated variable names 'homeownership, 'multiunit, 'med_income,
#
#
    4fed_spend
```









Cluster Validation

Cluster Validity

For supervised classification (= we have class labels) we have a variety of measures to evaluate how good our model is: Accuracy, Precision, Recall, F_1 , etc.

Cluster Validity

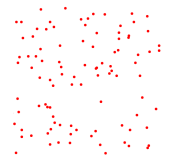
For supervised classification (= we have class labels) we have a variety of measures to evaluate how good our model is: Accuracy, Precision, Recall, F_1 , etc.

For cluster analysis (=unsupervised learning), the analogous question is:

How to evaluate the "goodness" of the resulting clusters?

Keep in mind that the dataset can be large and high-dimensional. Visualization might be difficult.

Random points:



Random points: Hierarchical K-means

 Determining the clustering tendency of a set of data, i.e., distinguishing whether non-random structure exists in the data (e.g., to avoid overfitting).

- 1. Determining the **clustering tendency** of a set of data, i.e., distinguishing whether non-random structure exists in the data (e.g., to avoid overfitting).
- 2. Internal Validation: Evaluating how well the cluster analysis results fit the data without reference to external information.

- 1. Determining the **clustering tendency** of a set of data, i.e., distinguishing whether non-random structure exists in the data (e.g., to avoid overfitting).
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- 4. Compare clusterings to determine which is better.

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- 2. Internal Validation: Evaluating how well the cluster analysis results fit the data without reference to external information.
- 3. External Validation: Compare the cluster analysis results to externally known class labels (class labels).
- 4. Compare clusterings to determine which is better.
- 5. Determining the 'correct' number of clusters.

For 2, 3, and 4, we can further distinguish whether we want to evaluate the entire clustering or just individual clusters.

Measures of Cluster Validity

Numerical measures applied to judge various aspects of cluster validity are classified into the following three types.

- ► Internal Index: Used to measure the goodness of a clustering structure without respect to external information.
- External Index: Used to measure the extent to which cluster labels match externally supplied class labels.
- Relative Index: Used to compare two different clusterings or clusters.

Internal Index

Consider a dataset

$$D = \{\vec{x}_1, \dots, \vec{x}_p\}$$

Assume that a clustering algorithm produced a partition $\mathcal{U} = \{U_1, \dots, U_K\}$ of D into K clusters.

No other information has been provided.

We aim to measure the clustering's "niceness" (??)

Internal Index

Consider a dataset

$$D = \{\vec{x}_1, \dots, \vec{x}_p\}$$

Assume that a clustering algorithm produced a partition $\mathcal{U} = \{U_1, \dots, U_K\}$ of D into K clusters.

No other information has been provided.

We aim to measure the clustering's "niceness" (??)

Assume that we have a distance measure d measuring how far apart the objects being clustered.

For concreteness:

- We stick with numerical features, which means that the dataset $D = \{\vec{x}_1, \dots, \vec{x}_p\}$ contains vectors $\vec{x}_i \in \mathbb{R}^n$.
- Assume the Euclidean distance d.

Note that the validity measures may be based on completely different similarity/dissimilarity measures and non-numerical data.

Consider a dataset
$$D = \{\vec{x}_1, \dots, \vec{x}_p\}$$
 and its clustering $\mathcal{U} = \{U_1, \dots, U_K\}$.

Let us utilize the concept of distance to cluster prototypes and consider the distance between prototypes.

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The proximity might be, e.g.,

- ▶ the distance $d(\vec{x}, \vec{z})$,
- ▶ the square of the distance, that is $d(\vec{x}, \vec{z})^2$,
- any other notion of dissimilarity based on the application.

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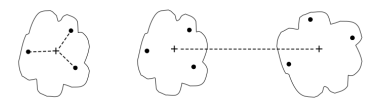
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We consider the notions of *cohesion* (proximity of examples within clusters) and *separation* (proximity of clusters).

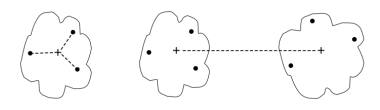


Prototype-based cohesion = the similarity of examples within a given cluster to a prototype of the cluster (e.g., centroid).

Given a cluster $U_j \in \mathcal{U}$ and its prototype $\vec{m}_j \in \mathbb{R}^n$,

$$cohesion(U_j) = \sum_{\vec{x} \in U_i} proximity(\vec{x}, \vec{m}_j)$$

Note that the prototype **does not** have to be an element of U_j . Intuitively, cohesion is the proximity of cluster's examples and a point somewhere "between" all examples of the cluster.



Prototype-based separation = dissimilarity of prototypes of different clusters.

Given a cluster $U_j \in \mathcal{U}$, its prototype $\vec{m}_j \in \mathbb{R}^n$, and a prototype of all examples $\vec{m} \in \mathbb{R}^n$ (e.g. the centroid of all examples)

$$separation(U_j) = proximity(\vec{m}_j, \vec{m})$$

Intuitively, separation is the proximity of the cluster's examples to the dataset's center.

Summarize the prototype-based cohesion and separation as follows:

$$\mathsf{cohesion}(\mathcal{U}) = \sum_{j=1}^{K} \mathsf{cohesion}(U_j)$$

$$= \sum_{j=1}^{K} \sum_{ec{x} \in U_j} \mathsf{proximity}(ec{x}, ec{m}_j)$$

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$$\begin{split} \text{separation}(\mathcal{U}) &= \sum_{j=1}^{K} |U_j| separation(U_j) \\ &= \sum_{i=1}^{K} |U_j| \textit{proximity}(\vec{m}_j, \vec{m}) \end{split}$$

If $proximity(\vec{x}, \vec{z})$ is defined as $d(\vec{x}, \vec{z})^2$ then the cohesion is the inertia.

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If $proximity(\vec{x}, \vec{z})$ is defined as $d(\vec{x}, \vec{z})^2$ then the cohesion is the inertia.

There is an interesting relationship between the above measures and the squared distances to the prototype of the whole dataset \vec{m} .

Consider a dataset $D = \{\vec{x}_1, \dots, \vec{x}_p\}$ and its clustering $\mathcal{U} = \{U_1, \dots, U_K\}$ of D.

Consider $proximity(\vec{x}, \vec{z}) = d(\vec{x}, \vec{z})^2$ and all prototypes to be centroids.

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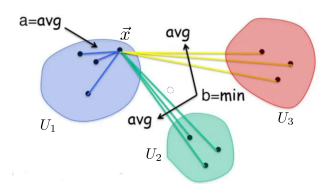
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The following holds:

$$\mathsf{TSS} = \mathsf{cohesion}(\mathcal{U}) + \mathsf{separation}(\mathcal{U})$$

Note that TSS is determined by D.

Silhouette score can be used to measure both qualities of clustering from the point of view of individual examples and from the point of view of the overall clustering.



$$\mathsf{silhouette}(\vec{x}) = \frac{b-a}{\mathsf{max}\{a,b\}}$$

Consider a clustering $\mathcal{U} = \{U_1, \dots, U_k\}$ and $\vec{x} \in U_j$.

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 and $\vec{x}\in U_j.$ If $|U_j|>1$ we define
$$a(\vec{x})=\frac{1}{|U_j|-1}\sum_{\vec{z}\in U_i\smallsetminus\{\vec{x}\}}d(\vec{x},\vec{z})$$

Consider a clustering $\mathcal{U} = \{U_1, \dots, U_k\}$ and $\vec{x} \in U_j$. If $|U_j| > 1$ we define

$$a(\vec{x}) = \frac{1}{|U_j| - 1} \sum_{\vec{z} \in U_i \setminus \{\vec{x}\}} d(\vec{x}, \vec{z})$$

$$b(\vec{x}) = \min_{k \neq j} \frac{1}{|U_k|} \sum_{\vec{z} \in U_k} d(\vec{x}, \vec{z})$$

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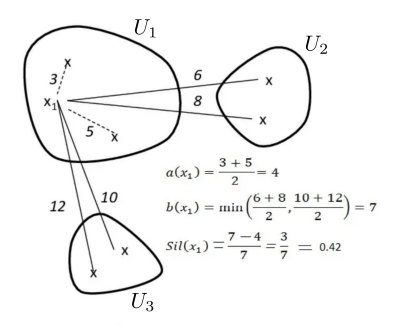
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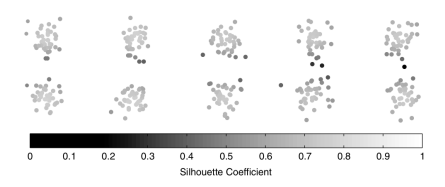
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If $|U_j| > 1$ we define

$$silhouette(\vec{x}) = \frac{b(\vec{x}) - a(\vec{x})}{\max\{a(\vec{x}), b(\vec{x})\}}$$

Else, we define $silhouette(\vec{x}) = 0$.





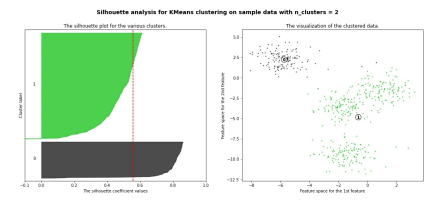
Silhouette for Clusters and Clusterings

We have defined the silhouette for a single $\vec{x} \in D$.

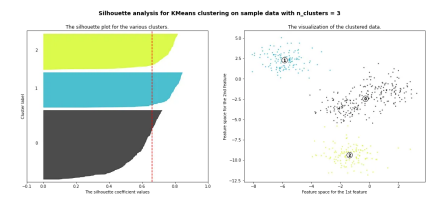
To obtain the silhouette score for a whole cluster U_j or for D we summarize using simple averaging:

$$silhouette(U_j) = \frac{1}{|U_j|} \sum_{\vec{x} \in U_j} silhouette(\vec{x})$$

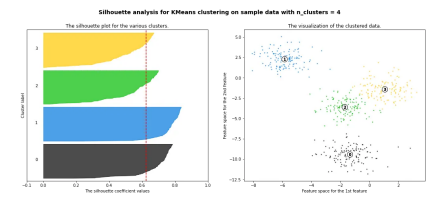
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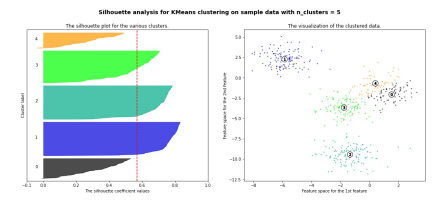
The colored graphs on the left are silhouette scores of the individual elements of clusters.



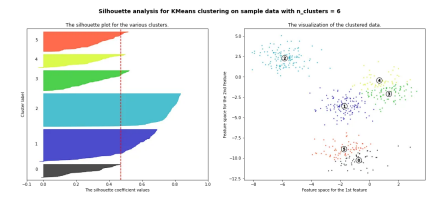
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External Index

Consider a *supervised learning* dataset

$$D = \{(\vec{x}_1, c_1), \dots, (\vec{x}_p, c_p)\}\$$

Here $c_i \in C$ is a class of $\vec{x_i}$.

Assume that a clustering algorithm produced a partition $\mathcal{U} = \{U_1, \dots, U_K\}$ of D into K clusters.

We measure how the clustering conforms with the given classes.

Purity

Consider the clustering to be a classification model.

Define a classifier $h:D\to C$ such that given $\vec{x_i}\in U_i\in\mathcal{U}$

 $h(\vec{x_i}) = \text{the most frequent class in } U_j$

Now we can measure the Accuracy of h.

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Accuracy of *h* is called *purity*.

Intuitively, it is the proportion of non-majority class elements in clusters.

Is it a good measure?

Probably not; many clusters lead to high purity (each element in its own cluster means purity = 1).

Classifier Point of View

Given $\vec{x_i}$, denote by $\mathcal{U}(\vec{x_i})$ the cluster $U_j \in \mathcal{U}$ containing $\vec{x_i}$.

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▶ TP = number of examples of the same class and the same cluster

$$\mathsf{TP} = |\{(i,j) \mid \mathcal{U}(\vec{x_i}) = \mathcal{U}(\vec{x_j}) \land c_i = c_j)\}$$

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- ▶ TN = number of examples of different classes and different clusters

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- ► FP = number of examples of the same class and different clusters $\mathsf{FP} = |\{(i,j) \mid \mathcal{U}(\vec{x_i}) = \mathcal{U}(\vec{x_j}) \land c_i \neq c_j)\}$

Classifier Point of View

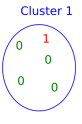
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Now, we may apply all the measures from the supervised model.

Example



$$TP = {4 \choose 2} + {5 \choose 2} + {2 \choose 2}$$

$$= 6 + 10 + 1 = 17$$

$$TN = 4 * 5 + 1 * 2 = 22$$

$$FP = 1 * 4 + 5 * 2 = 14$$

$$FN = 1 * 5 + 4 * 2 = 13$$

		Cluster	
		same	diff
Class	same	TP=17	FN=13
	diff	FP=14	TN=22

Rand Index

Accuracy (in this area known as Rand index) is

$$\mathsf{RandInd} = \mathsf{Accuracy} = \frac{\mathsf{TP} + \mathsf{TN}}{\mathsf{TP} + \mathsf{TN} + \mathsf{FP} + \mathsf{FN}}$$

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In our example,

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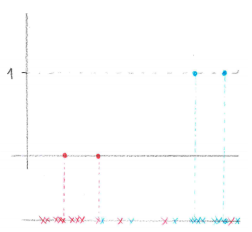
Here, note that the Rand index considers the purity and the number of clusters.

Note that the Rand index can be used to compare two clusterings: Simply consider class labels to be indicators of clusters.

Similarly, we may compute the other measures such as Precision, Recall, and F_1 with all their benefits and limitations.

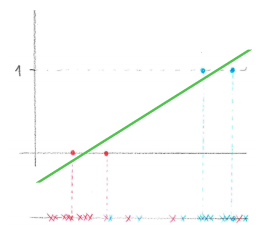
What about classification using regression?

Binary classification: Desired outputs 0 and 1 ... we want to capture the probability distribution of the classes



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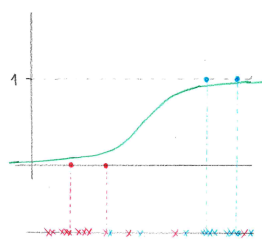
Binary classification: Desired outputs 0 and 1 ... we want to capture the probability distribution of the classes



... does not capture the probability well (it is not probability at all)

What about classification using regression?

Binary classification: Desired outputs 0 and 1 ... we want to capture the probability distribution of the classes



... logistic sigmoid $\frac{1}{1+e^{-(\vec{w}\cdot\vec{\mathbf{x}})}}$ is much better!

Logistic regression model $h[\vec{w}]$ is determined by a vector of weights $\vec{w} = (w_0, w_1, \dots, w_n) \in \mathbb{R}^{n+1}$ as follows:

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Given
$$\vec{x} = (x_1, \dots, x_n) \in \mathbb{R}^n$$
,

$$h[\vec{w}](\vec{x}) := \frac{1}{1 + e^{-\left(w_0 + \sum_{k=1}^n w_k x_k\right)}} = \frac{1}{1 + e^{-\vec{w} \cdot \tilde{\mathbf{x}}}}$$

Here

$$\tilde{\mathbf{x}} = (x_0, x_1, \dots, x_n)$$
 where $x_0 = 1$

is the augmented feature vector.

The model gives probability $h[\vec{w}](\vec{x})$ of the class 1 given an input \vec{x} . But why do we model such probability using $1/(1+e^{-\vec{w}\cdot\tilde{\mathbf{x}}})$??

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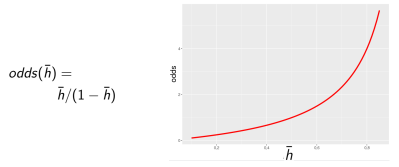
Denote by \bar{h} the probability $P(Y = 1 \mid X = \vec{x})$, i.e., the "true" probability of the class 1 given features \vec{x} .

The probability \bar{h} cannot be easily modeled using a linear function (the probabilities are between 0 and 1).

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What about odds of the class 1?

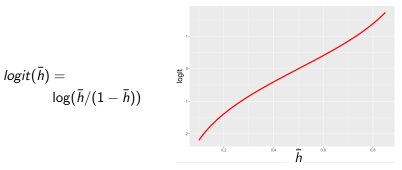


Better, at least it is unbounded on one side ...

The model gives probability $h[\vec{w}](\vec{x})$ of the class 1 given an input \vec{x} . But why do we model such probability using $1/(1+e^{-\vec{w}\cdot\tilde{\mathbf{x}}})$??

Denote by \bar{h} the probability $P(Y=1 \mid X=\vec{x})$, i.e., the "true" probability of the class 1 given features \vec{x} .

What about log odds (aka logit) of the class 1?



Looks almost linear, at least for probabilities not too close to 0 or 1 ...

Assume that \bar{h} is the actual probability of the class 1 for an "object" with features $\vec{x} \in \mathbb{R}^n$. Put

$$\log(ar{h}/(1-ar{h})) = ec{w} \cdot \mathbf{\tilde{x}}$$

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and

$$(1-\bar{h})/\bar{h}=e^{-\vec{w}\cdot\tilde{\mathbf{x}}}$$

and

$$\bar{h} = \frac{1}{1 + e^{-\vec{w} \cdot \tilde{\mathbf{x}}}} = h[\vec{w}](\vec{x})$$

If we model log odds using a linear function, the probability is obtained by applying the logistic sigmoid on the result of the linear function.

► Given a set *D* of training samples:

$$D = \{ (\vec{x}_1, c_1), (\vec{x}_2, c_2), \dots, (\vec{x}_p, c_p) \}$$

Here
$$\vec{x}_k = (x_{k1} \dots, x_{kn}) \in \mathbb{R}^n$$
 and $c_k \in \{0, 1\}$.

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Recall that
$$h[\vec{w}](\vec{x}_k) = 1/(1+e^{-\vec{w}\cdot\tilde{\mathbf{x}}_k})$$
 where $\tilde{\mathbf{x}}_k = (x_{k0}, x_{k1}\dots, x_{kn})$, here $x_{k0} = 1$

Our goal: Find \vec{w} such that for every $k=1,\ldots,p$ we have that $h[\vec{w}](\vec{x}_k)\approx c_k$

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Our goal: Find \vec{w} such that for every k = 1, ..., p we have that $h[\vec{w}](\vec{x}_k) \approx c_k$

Binary Cross-entropy:

$$E(\vec{w}) = -\sum_{k=1}^{p} c_k \log(h[\vec{w}](\vec{x}_k)) + (1-c_k) \log(1-h[\vec{w}](\vec{x}_k))$$

Gradient of the Error Function

Consider the **gradient** of the error function:

$$\nabla E(\vec{w}) = \left(\frac{\partial E}{\partial w_0}(\vec{w}), \dots, \frac{\partial E}{\partial w_n}(\vec{w})\right) = \sum_{k=1}^p \left(h[\vec{w}](\vec{x}_k) - c_k\right) \cdot \tilde{\mathbf{x}}_k$$

Fact 1

If $\nabla E(\vec{w}) = \vec{0} = (0, \dots, 0)$, then \vec{w} is a global minimum of E.

This follows from the fact that E is convex.

Using the squared error with the logistic sigmoid would lead to a non-convex error with several minima!

Logistic Regression – Learning

Gradient Descent:

• Weights $\vec{w}^{(0)}$ are initialized randomly close to $\vec{0}$.

Logistic Regression - Learning

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Note that the algorithm is almost similar to the batch perceptron algorithm!

Proposition

For sufficiently small $\varepsilon > 0$, the sequence $\vec{w}^{(0)}, \vec{w}^{(1)}, \vec{w}^{(2)}, \ldots$ converges (in a component-wise manner) to the global minimum of the error function E.

Logistic Regression - Using the Trained Model

We have already trained our logistic regression model, i.e., we have a vector of weights $\vec{w} = (w_0, w_1, \dots, w_n)$.

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The model is the function $h[\vec{w}]$ which for a given feature vector $\vec{x} = (x_1, \dots, x_n)$ returns the probability

$$h[\vec{w}](\vec{x}) = \frac{1}{1 + e^{-(w_0 + \sum_{k=1}^{n} w_k x_k)}}$$

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To decide whether a given \vec{x} belongs to the class 1 we use $h[\vec{w}]$ as a Bayes classifier: Assign \vec{x} to the class 1 iff $h[\vec{w}](\vec{x}) \ge 1/2$.

Other thresholds can also be used depending on the application and properties of the model. In such a case, given a threshold $\xi \in [0,1]$, assign \vec{x} to the class 1 iff $h[\vec{w}](\vec{x}) \geq \xi$.

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$$h[w_0, w_1](x_k) = \frac{1}{1 + e^{-(w_0 + w_1 \cdot x_k)}}$$

and 0 otherwise.

Here w_0 , w_1 are unknown weights.

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How "probable" is it to generate the correct classes c_1, \ldots, c_p ?

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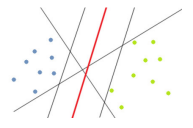
The following conditions are equivalent:

- \triangleright w_0 , w_1 minimize the binary cross-entropy E
- ▶ w_0 , w_1 maximize the likelihood (i.e., the "probability") of generating the correct values c_1, \ldots, c_p using the above described Bernoulli trials (i.e., that $c_k' = c_k$ for all $k = 1, \ldots, p$)

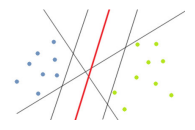
Note that the above equivalence is a property of the cross-entropy and is not dependent on the "implementation" of $h[w_0, w_1](x_k)$ using the logistic sigmoid.

Support Vector Machines (SVM)

SVM Idea – Which Linear Classifier is the Best?



SVM Idea – Which Linear Classifier is the Best?



Benefits of maximum margin:

- Intuitively, the maximum margin is good w.r.t. generalization.
- ▶ Only the *support vectors* (those on the margin) matter; others can, in principle, be ignored.

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Consider a linear classifier:

$$h[\vec{w}](\vec{x}) := \begin{cases} 1 & w_0 + \sum_{i=1}^n w_i \cdot x_i = \vec{w} \cdot \tilde{\mathbf{x}} \ge 0 \\ -1 & w_0 + \sum_{i=1}^n w_i \cdot x_i = \vec{w} \cdot \tilde{\mathbf{x}} < 0 \end{cases}$$

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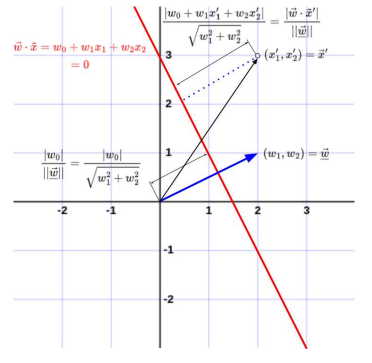
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The *distance* of \vec{x} from the separating hyperplane determined by \vec{w} is

$$d[\vec{w}](\vec{x}) = \frac{|\vec{w} \cdot \tilde{\mathbf{x}}|}{\|\vec{w}\|}$$

Recall that $\vec{w} \cdot \tilde{\mathbf{x}}$ is positive for \vec{x} on the side to which $\underline{\vec{w}}$ points and negative on the opposite side.



Margin

Given a training set

$$D = \{ (\vec{x}_1, y_1), (\vec{x}_2, y_2), \dots, (\vec{x}_p, y_p) \}$$
Here $\vec{x}_k = (x_{k1}, \dots, x_{kn}) \in X \subseteq \mathbb{R}^n$ and $y_k \in \{-1, 1\}$.

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Assume that D is linearly separable, let \vec{w} be consistent with D.

Margin of \vec{w} is twice the minimum distance between feature vectors \vec{x}_k and the separating hyperplane determined by \vec{w} , i.e.,

$$2\min_{k} d[\vec{w}](\vec{x}_{k}) = 2\min_{k} \frac{|\vec{w} \cdot \tilde{\mathbf{x}}_{k}|}{\|\underline{\vec{w}}\|}$$

▶ Our goal is to find \vec{w} consistent with D that maximizes the margin. Note that to maximize the margin it suffices to maximize $\min_k \frac{|\vec{w} \cdot \vec{x}_k|}{||\vec{w}||}$ over \vec{w} consistent with D.

Finding the Maximum Margin Classifier

We want to maximize the minimum distance of the feature vectors \vec{x}_k from the separating hyperplane determined by \vec{w} .

Finding the Maximum Margin Classifier

We want to maximize the minimum distance of the feature vectors \vec{x}_k from the separating hyperplane determined by \vec{w} .

Formally, we use the following:

To maximize the margin, find \vec{w} maximizing

$$\min_{k} \frac{|\vec{w} \cdot \tilde{\mathbf{x}}_{k}|}{||\vec{w}||}$$
 (= the distance of closest $\vec{x_k}$'s to the sep. hyperplane)

over the following constraints

$$\vec{w} \cdot \tilde{\mathbf{x}}_k > 0$$
 for all k satisfying $y_k = 1$

$$ec{w} \cdot \mathbf{\tilde{x}}_k < 0$$
 for all k satisfying $y_k = -1$

(the contraints make sure that \vec{w} is consistent with the training set D)

$$\min_{k} \frac{|\vec{w} \cdot \tilde{\mathbf{x}}_{k}|}{||\underline{\vec{w}}||}$$

over the following constraints

$$\vec{w} \cdot \tilde{\mathbf{x}}_k > 0$$
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Can be made more succinct:

To maximize the margin, find \vec{w} maximizing

$$\min_{k} \frac{y_k \cdot \vec{w} \cdot \tilde{\mathbf{x}}_k}{\|\vec{w}\|}$$
 over $\min_{k} (y_k \cdot \vec{w} \cdot \tilde{\mathbf{x}}_k) > 0$

The reason is that \vec{w} is consistent with D. That is, $\vec{w} \cdot \tilde{x}_k > 0$ for $y_k = 1$, and $\vec{w} \cdot \tilde{x}_k < 0$ for $y_k = -1$.

$$\min_{k} \frac{y_k \cdot \vec{w} \cdot \tilde{\mathbf{x}}_k}{\|\vec{w}\|} \quad \text{over} \quad \min_{k} (y_k \cdot \vec{w} \cdot \tilde{\mathbf{x}}_k) > 0$$

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Observation: For every \vec{w} satisfying $\min_k (y_k \cdot \vec{w} \cdot \tilde{\mathbf{x}}_k) > 0$ there is \vec{w}' satisfying $\min_k (y_k \cdot \vec{w}' \cdot \tilde{\mathbf{x}}_k) = 1$ such that

$$\min_{k} \frac{y_k \cdot \vec{w} \cdot \tilde{\mathbf{x}}_k}{\|\vec{w}\|} = \min_{k} \frac{y_k \cdot \vec{w}' \cdot \tilde{\mathbf{x}}_k}{\|\vec{w}'\|}$$

Proof: Just consider $\vec{w}' = \vec{w}/\xi$ where $\xi = \min_k (y_k \cdot \vec{w} \cdot \tilde{\mathbf{x}}_k)$.

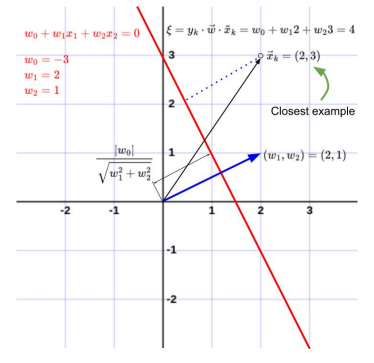
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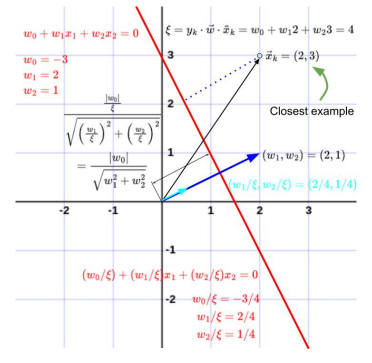
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$$\min_{k} \frac{y_k \cdot \vec{w} \cdot \mathbf{\tilde{x}}_k}{\|\vec{w}\|}$$
 over $\min_{k} (y_k \cdot \vec{w} \cdot \mathbf{\tilde{x}}_k) = 1$

$$\min_{k} \frac{y_k \cdot \vec{w} \cdot \tilde{\mathbf{x}}_k}{\|\vec{w}\|}$$
 over $\min_{k} (y_k \cdot \vec{w} \cdot \tilde{\mathbf{x}}_k) = 1$

can be further simplified to

$$rac{1}{\|ec{w}\|}$$
 over $\min_k (y_k \cdot ec{w} \cdot \widetilde{\mathbf{x}}_k) = 1$

$$rac{1}{|ec{w}||}$$
 over $\min_k (y_k \cdot ec{w} \cdot \mathbf{ ilde{x}}_k) = 1$

$$\frac{1}{\|\vec{w}\|}$$
 over $\min_{k} (y_k \cdot \vec{w} \cdot \tilde{\mathbf{x}}_k) = 1$

Can be adjusted by loosening the constraints:

To maximize the margin, find \vec{w} maximizing

$$rac{1}{\|ec{w}\|}$$
 over $\min_{k}(y_k \cdot ec{w} \cdot \widetilde{\mathbf{x}}_k) \geq 1$

If the latter is solved by \vec{w}' with $\min_k (y_k \cdot \vec{w}' \cdot \tilde{\mathbf{x}}_k) > 1$, then

$$\min_{k} \frac{y_k \cdot \vec{w}' \cdot \tilde{\mathbf{x}}_k}{\left|\left|\underline{\vec{w}}'\right|\right|} > \frac{1}{\left|\left|\underline{\vec{w}}'\right|\right|} \ge \frac{1}{\left|\left|\underline{\vec{w}}\right|\right|} = \frac{\min_{k} y_k \cdot \vec{w} \cdot \tilde{\mathbf{x}}_k}{\left|\left|\underline{\vec{w}}\right|\right|}$$

For all \vec{w} satisfying $\min_k (y_k \cdot \vec{w} \cdot \vec{x}_k) = 1$, which contradicts the fact that the maximum margin is attained by such a \vec{w} .

 $rac{1}{|ec{w}||}$ over $\min_{k} y_k \cdot ec{w} \cdot \mathbf{ ilde{x}}_k \geq 1$

$$rac{1}{\|ec{oldsymbol{w}}\|}$$
 over $\displaystyle \min_{k} y_k \cdot ec{oldsymbol{w}} \cdot \mathbf{ ilde{x}}_k \geq 1$

Can be turned into

$$||\underline{\vec{w}}||$$
 over $\min_{k} y_k \cdot \vec{w} \cdot \tilde{\mathbf{x}}_k \ge 1$

$$\dfrac{1}{\| ec{ ec{w}} \|}$$
 over $\displaystyle \min_k y_k \cdot ec{w} \cdot \mathbf{ ilde{x}}_k \geq 1$

Can be turned into

To maximize the margin, find \vec{w} minimizing

$$||\underline{\vec{w}}||$$
 over $\min_{k} y_k \cdot \vec{w} \cdot \tilde{\mathbf{x}}_k \ge 1$

And, finally,

To maximize the margin, find \vec{w} minimizing

$$\underline{\vec{w}} \cdot \underline{\vec{w}}$$
 over $y_k \cdot \vec{w} \cdot \mathbf{\tilde{x}}_k \ge 1$ for all k

Indeed, just note that $||\underline{\vec{w}}|| = \sqrt{\underline{\vec{w}} \cdot \underline{\vec{w}}}$.

SVM - Optimization

Assume a given training set

$$D = \{ (\vec{x}_1, y_1), (\vec{x}_2, y_2), \dots, (\vec{x}_p, y_p) \}$$

Here
$$\vec{x}_k = (x_{k1} \dots, x_{kn}) \in X \subseteq \mathbb{R}^n$$
 and $y_k \in \{-1, 1\}$. (recall $\tilde{x}_k = (x_{k0}, x_{k1}, \dots, x_{kn})$ where $x_{k0} = 1$)

Margin maximization as a quadratic optimization problem:

Find \vec{w} minimizing

$$\vec{w} \cdot \vec{w}$$

under the constraints

$$y_k \cdot \vec{w} \cdot \tilde{\mathbf{x}}_k \geq 1$$
 for all k

Support vectors are vectors \vec{x}_k closest to the optimal separating hyperplane, i.e., those satisfying $y_k \cdot \vec{w} \cdot \vec{x}_k = 1$ for a minimizing \vec{w} .

Example

Training set:

$$D = \{((0,0),-1),((1,1),1),((0,3),1)\}$$

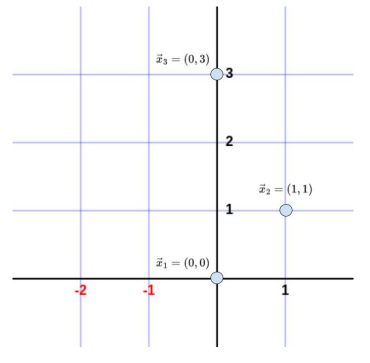
That is

$$\vec{x}_1 = (0,0)$$
 $\vec{x}_1 = (1,0,0)$ $\vec{x}_2 = (1,1)$ $\vec{x}_3 = (0,3)$ $\vec{x}_3 = (1,0,3)$

$$y_1 = -1$$

$$y_2 = 1$$

$$y_3 = 1$$



Find \vec{w} minimizing $w_1^2+w_2^2$ under the constraints $(-1)\cdot (1w_0+0w_1+0w_2)=-w_0\geq 1$ $1\cdot (1w_0+1w_1+1w_2)=w_0+w_1+w_2\geq 1$ $1\cdot (1w_0+0w_1+3w_2)=w_0+3w_2\geq 1$

It can be solved using a quadratic programming solver.

Find \vec{w} minimizing $w_1^2 + w_2^2$ under the constraints

$$(-1) \cdot (1w_0 + 0w_1 + 0w_2) = -w_0 \ge 1$$
$$1 \cdot (1w_0 + 1w_1 + 1w_2) = w_0 + w_1 + w_2 \ge 1$$
$$1 \cdot (1w_0 + 0w_1 + 3w_2) = w_0 + 3w_2 \ge 1$$

It can be solved using a quadratic programming solver.

To solve by hand, assume that we know that $\vec{x_1}$ and $\vec{x_2}$ are support vectors.

Find
$$\vec{w}$$
 minimizing $w_1^2+w_2^2$ under the constraints
$$-w_0=1$$

$$w_0+w_1+w_2=1$$

$$w_0+3w_2\geq 1$$

Note that the equality constraints correspond to our assumption that \vec{x}_1 and \vec{x}_2 are support vectors.

Find
$$\vec{w}$$
 minimizing $w_1^2+w_2^2$ under the constraints

$$-w_0 = 1$$

$$w_0 + w_1 + w_2 = 1$$

$$w_0 + 3w_2 \ge 1$$

Find
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$$w_0 + 3w_2 \ge 1$$

Can be transformed to

Find
$$\vec{w}$$
 minimizing $w_1^2 + w_2^2$ under the constraints

$$w_1 + w_2 = 2$$
$$3w_2 \ge 2$$

Find \vec{w} minimizing $w_1^2 + w_2^2$ under the constraints

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Find \vec{w} minimizing $w_1^2 + w_2^2$ under the constraints

$$w_1 + w_2 = 2$$
$$3w_2 \ge 2$$

Substituting $w_2 = 2 - w_1$ into the quadratic function we obtain

$$w_1^2 + (2 - w_1)^2 = w_1^2 + w_1^2 - 4w_1 + 4 = 2w_1^2 - 4w_1 + 4$$

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substituting $w_2=2-w_1$ into the inequality $3w_2\geq 2$ we obtain

$$6 - 3w_1 \ge 2$$

Find \vec{w} minimizing $w_1^2 + w_2^2$ under the constraints

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substituting $w_2 = 2 - w_1$ into the inequality $3w_2 \ge 2$ we obtain

$$6 - 3w_1 \ge 2$$

This reduces our problem to

Find \vec{w} minimizing $2w_1^2 - 4w_1 + 4$ under the constraint $w_1 \leq \frac{4}{3}$

Is solved by

$$w_1 = 1$$

Is solved by

$$w_1 = 1$$

From $w_2 = 2 - w_1$ we obtain

$$w_2=2-1=1$$

Is solved by

$$w_1 = 1$$

From $w_2 = 2 - w_1$ we obtain

$$w_2 = 2 - 1 = 1$$

From $-w_0 = 1$ we obtain

$$w_0 = -1$$

Is solved by

$$w_1 = 1$$

From $w_2 = 2 - w_1$ we obtain

$$w_2 = 2 - 1 = 1$$

From $-w_0 = 1$ we obtain

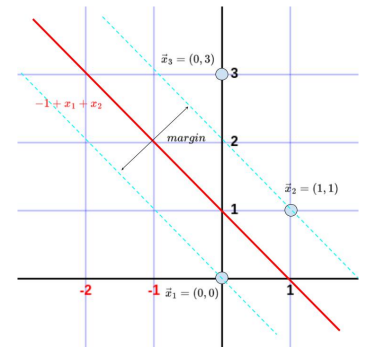
$$w_0 = -1$$

The final model is

$$h[\vec{w}](\vec{x}) = -1 + x_1 + x_2$$

The separating hyperplane is determined by

$$-1 + x_1 + x_2 = 0$$



SVM – Optimization

Need to optimize a quadratic function subject to linear constraints.

SVM – Optimization

- Need to optimize a quadratic function subject to linear constraints.
- Quadratic optimization problems are a well-known class of mathematical programming problems for which efficient methods (and tools) exist.

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SVM – Optimization

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The answer lies in their ability to deal with non-linearly separable sets efficiently using the *kernel trick* (see a later lecture).

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 - Afterwards, only support vectors matter in the solution! Leave only them in the training set, and add new training examples.
 - ► This iterative procedure decreases the (general) cost function.

Soft-margin SVM

Trade-off few misclassifications with a wide margin for the rest.

Find \vec{w} minimizing

$$\underline{\vec{w}} \cdot \underline{\vec{w}} + C \sum_{k} \zeta_{k}$$

C is a hyperparameter

under the constraints

$$y_k \cdot \vec{w} \cdot \tilde{\mathbf{x}}_k \ge 1 - \zeta_k$$
 for all k

$$\zeta_k \ge 0$$
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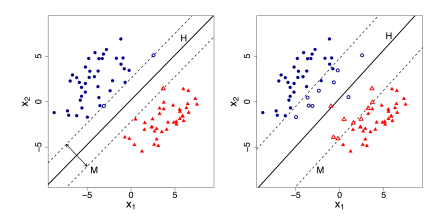
$$\zeta_k > 0$$
 for all k

Which is the same as the following unconstrained optimization:

Find \vec{w} minimizing the hinge loss

$$\underline{\vec{w}} \cdot \underline{\vec{w}} + C \sum_{k} \max(0, 1 - y_k \cdot \vec{w} \cdot \tilde{\mathbf{x}}_k)$$

Hard vs Soft Margin SVM



Source: Dishaa Agarwal

https://www.analyticsvidhya.com/blog/2021/04/insight-into-svm-support-vector-machine-along-with-code/

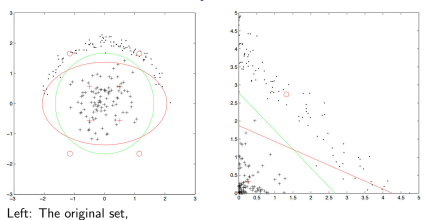
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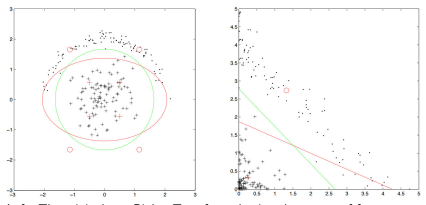
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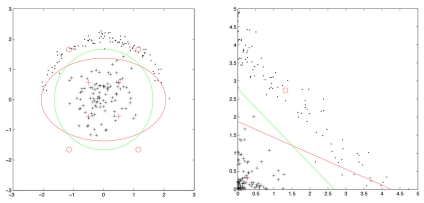
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- ➤ SVMs can be applied to complex data types beyond feature vectors (e.g., graphs, sequences, relational data) by designing kernel functions for such data.
- SVM techniques have been extended to several tasks, such as regression [Vapnik et al. '97], principal component analysis [Schölkopf et al. '99], etc.

Kernel Methods



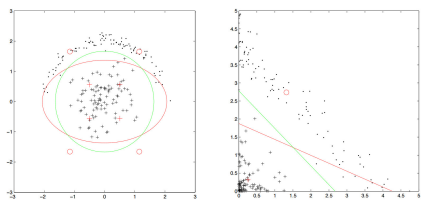


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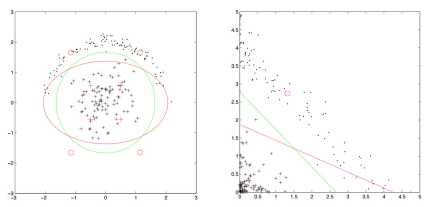
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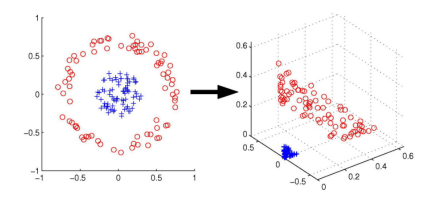
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How to classify (in the original space): First, transform a given feature vector by squaring the features, then use a linear classifier.

Another Solution



Mapping from \mathbb{R}^2 to \mathbb{R}^3 so that there is "more space" for linear separation.

Do We Need to Map Explicitly?

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But first, we need to *dualize* our learning algorithm.

Linear Regression

Given a set D of training examples:

$$D = \{ (\vec{x}_1, f_1), (\vec{x}_2, f_2), \dots, (\vec{x}_p, f_p) \}$$

Here $\vec{x}_k = (x_{k1} \dots, x_{kn}) \in \mathbb{R}^n$ and $f_k \in \mathbb{R}$.

- ▶ Our goal: Find \vec{w} so that $h[\vec{w}](\vec{x_k}) = \vec{w} \cdot \tilde{\mathbf{x}}_k$ is close to f_k for every $k = 1, \ldots, p$. Recall that $\tilde{\mathbf{x}}_k = (x_{k0}, x_{k1}, \ldots, x_{kn})$ where $x_{k0} = 1$.
- Squared Error Function:

$$E(\vec{w}) = \frac{1}{2} \sum_{k=1}^{p} (\vec{w} \cdot \tilde{\mathbf{x}}_{k} - f_{k})^{2} = \frac{1}{2} \sum_{k=1}^{p} \left(\sum_{i=0}^{n} w_{i} x_{ki} - f_{k} \right)^{2}$$

Regularized Linear Regression

Regularized Squared Error Function:

$$E(\vec{w}) = \frac{1}{2} \sum_{k=1}^{p} (\vec{w} \cdot \tilde{\mathbf{x}}_{k} - f_{k})^{2} + \vec{w} \cdot \vec{w}$$

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The Representer Theorem: The weight vector \vec{w}^* minimizing the regularized squared error function can be written as

$$\vec{w}^* = \sum_{i=1}^p \alpha_i f_i \, \tilde{\mathbf{x}}_i$$
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Substituting this expression for weights in *E* gives

$$E'(\vec{w}^*) = \frac{1}{2} \sum_{k=1}^{p} \left(\sum_{i=1}^{p} \alpha_i f_i \, \tilde{\mathbf{x}}_i \cdot \tilde{\mathbf{x}}_k - f_k \right)^2 + \sum_{i=1}^{p} \sum_{j=1}^{p} \alpha_i \alpha_j f_i f_j \, \tilde{\mathbf{x}}_i \cdot \tilde{\mathbf{x}}_j$$

and we minimize E' w.r.t. $\alpha_1, \ldots, \alpha_p$. What is this good for??

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The resulting coefficients $\alpha_1, \ldots, \alpha_p$ give a weight vector

$$\vec{w}^* = \sum_{i=1}^p \alpha_i f_i \, \tilde{\mathbf{x}}_i$$

which in turn gives a linear model

$$h[\vec{w}^*](\vec{x}) = \vec{w}^* \tilde{\mathbf{x}} = \sum_{i=1}^{p} \alpha_i f_i \, \tilde{\mathbf{x}}_i \cdot \tilde{\mathbf{x}}$$

Note that all $\tilde{\mathbf{x}}, \tilde{\mathbf{x}}_i, \tilde{\mathbf{x}}_i, \tilde{\mathbf{x}}_k$ occur in dot products with themselves!

Find $\vec{\alpha} = (\alpha_1, \dots, \alpha_p)$ minimizing dual regularized squared error

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Non-linear model: $h[\vec{\alpha}](\vec{x}) = \sum_{i=1}^{p} \alpha_i f_i \kappa(\tilde{\mathbf{x}}_i, \tilde{\mathbf{x}})$

Here κ is a **kernel function**. But now what is the trick?

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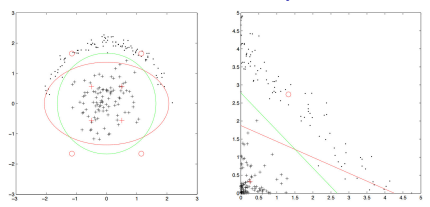
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The trick is that suitable kernel functions κ correspond to dot products in transformed spaces!

Recall the Quadratic Decision Boundary



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Right: the green line is a separating hyperplane in the transformed space.

Left: the green ellipse maps exactly to the green line.

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But now consider a mapping ϕ to \mathbb{R}^6 defined by

$$\phi(\tilde{\mathbf{x}}_k) = (1, x_{k1}^2, x_{k2}^2, \sqrt{2}x_{k1}x_{k2}, \sqrt{2}x_{k1}, \sqrt{2}x_{k2})$$

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THE Idea: Using the kernel $\kappa(\tilde{\mathbf{x}}_k, \tilde{\mathbf{x}}_\ell) = (\tilde{\mathbf{x}}_k \cdot \tilde{\mathbf{x}}_\ell)^2$ in the kernel, dual regularized squared error corresponds to using the regularized squared error after the transformation ϕ .

Quadratic Decision Boundary

Given a set D of training examples:

$$D = \{ (\vec{x}_1, f_1), (\vec{x}_2, f_2), \dots, (\vec{x}_p, f_p) \}$$

Assume that $f_i \in \{1, -1\}$ indicates the class of $\vec{x_i}$.

Yes, I know that squared error regression should not be used for classification!

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Considering $\kappa(\tilde{\mathbf{x}}_k, \tilde{\mathbf{x}}_\ell) = (\tilde{\mathbf{x}}_k \cdot \tilde{\mathbf{x}}_\ell)^2$ in our kernel dual regularized squared error we obtain

Find
$$\vec{\alpha} = \alpha_1, \dots, \alpha_p$$
 minimizing

$$E'(\vec{w}) = \frac{1}{2} \sum_{k=1}^{p} \left(\sum_{i=1}^{p} \alpha_{i} f_{i} \left(\mathbf{\tilde{x}}_{i} \cdot \mathbf{\tilde{x}}_{k} \right)^{2} - f_{k} \right)^{2} + \sum_{i=1}^{p} \sum_{j=1}^{p} \alpha_{i} \alpha_{j} f_{i} f_{j} \left(\mathbf{\tilde{x}}_{i} \cdot \mathbf{\tilde{x}}_{j} \right)^{2}$$

Non-linear classifier: $h[\vec{\alpha}](\vec{x}) = \sum_{i=1}^{p} \alpha_i f_i (\tilde{\mathbf{x}}_i \cdot \tilde{\mathbf{x}})^2$

Intuitively, minimizing E' in \mathbb{R}^2 gives a separating hyperplane for the input vectors transformed into \mathbb{R}^5 . This means, that in \mathbb{R}^2 it searches for a quadratic (i.e., non-linear) boundary.

▶ Linear: $\kappa(\mathbf{\tilde{x}}_{\ell}, \mathbf{\tilde{x}}_{\mathbf{k}}) = \mathbf{\tilde{x}}_{\ell} \cdot \mathbf{\tilde{x}}_{\mathbf{k}}$

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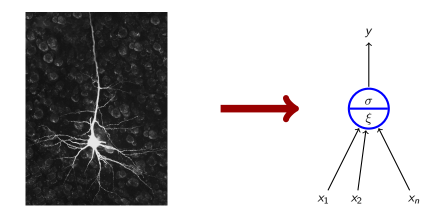
. . . .

Choosing kernels remains to be the black magic of kernel methods. They are usually chosen based on trial and error (of course, experience and additional insight into data help).

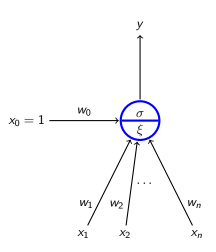
A similar trick can be done with (soft-margin) support vector machines.

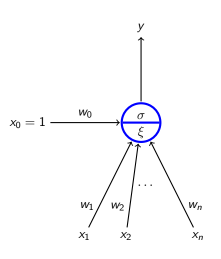
Neural Networks

(Primitive) Mathematical Model of Neuron

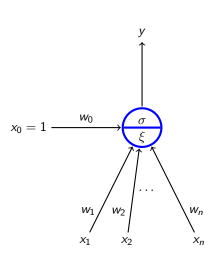


 $ightharpoonup x_1, \dots, x_n$ real *inputs*

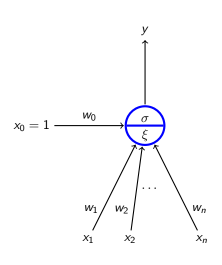




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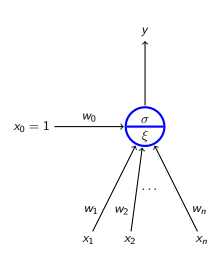


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In general, other potentials are considered (e.g. Gaussian), more on this in PV021.



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- y output defined by $y = \sigma(\xi)$ where σ is an activation function. We consider several activation functions.
 - e.g., linear threshold function

$$\sigma(\xi) = sgn(\xi) = egin{cases} 1 & \xi \geq 0 \ 0 & \xi < 0. \end{cases}$$

Formal Neuron vs Linear Models

 \triangleright If σ is a linear threshold function

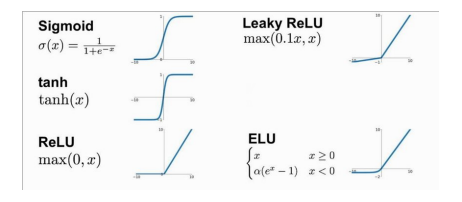
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We obtained a linear classifier.

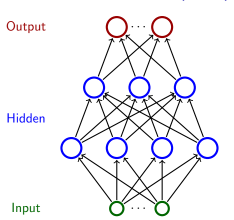
- If σ is identity, i.e., $\sigma(\xi) = \xi$, we obtain a linear (affine) function.
- ▶ If $\sigma(\xi) = 1/(1 + e^{-\xi})$ we obtain the logistic regression.

Also, other activation functions are used in neural networks!

Activation Functions

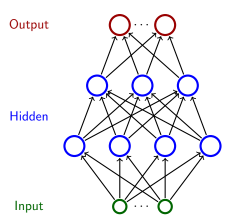


Multilayer Perceptron (MLP)



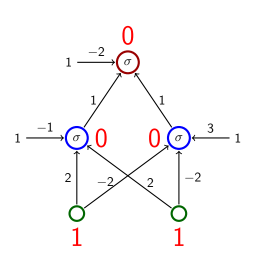
- Neurons are organized in *layers* (input layer, output layer, possibly several hidden layers)
- ► Layers are numbered from 0; The input is 0-th
- Neurons in the ℓ -th layer are connected with all neurons in the $\ell+1$ -th layer

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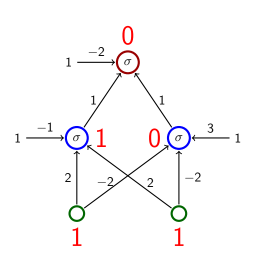


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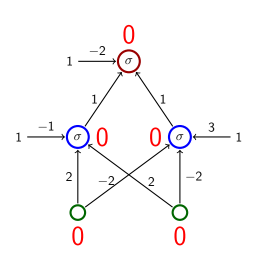
Intuition: The network computes a function: Assign input values to the input neurons and 0 to the rest. Proceed upwards through the layers, one layer per step. In the $\ell\text{-th}$ step consider output values of neurons in $\ell-1\text{-th}$ layer as inputs to neurons of the $\ell\text{-th}$ layer. Compute output values of neurons in the $\ell\text{-th}$ layer.



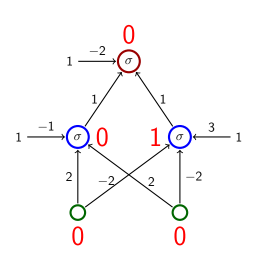
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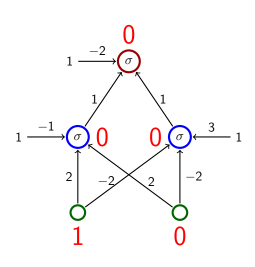
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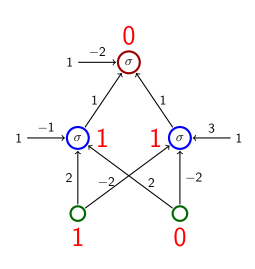
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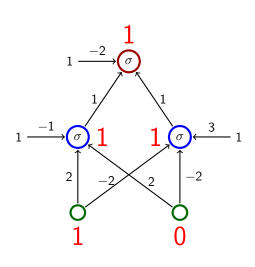
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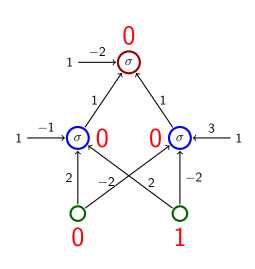
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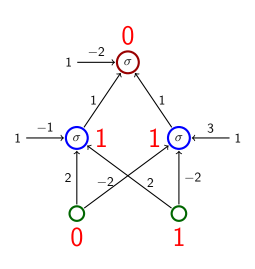
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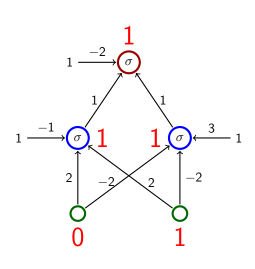
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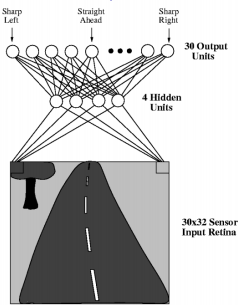


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Classical Example – ALVINN



- One of the first autonomous car. driving systems (in the 90s) ALVINN drives a car
- ▶ The net has $30 \times 32 = 960$ input neurons (the input space is \mathbb{R}^{960}).
- ► The value of each input captures the shade of gray of the corresponding pixel.
- Output neurons indicate where to turn (to the center of gravity).

Source: http://jmvidal.cse.sc.edu/talks/ann/alvin.html

A Bit of History

▶ Perceptron (Rosenblatt et al., 1957)



- Single layer (i.e., no hidden layers), the activation function linear threshold
 (i.e., a bit more general linear classifier)
- Perceptron learning algorithm
- Used to recognize digits

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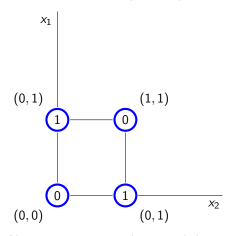
 Single layer, the activation function identity (i.e., a bit more linear function)

- Online version of the gradient descent
- ▶ Used a new circuitry element called *memristor* which was able to "remember" history of current in form of resistance

In both cases, the expressive power is somewhat limited- it can only express linear decision boundaries and linear (affine) functions.

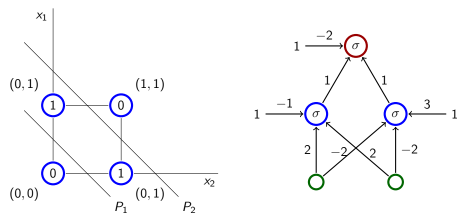
A Bit of History

One of the famous (counter)-examples: XOR



No perceptron can distinguish between ones and zeros.

XOR vs Multilayer Perceptron



(Here, σ is a linear threshold function.)

$$P_1: -1 + 2x_1 + 2x_2 = 0$$
 $P_2: 3 - 2x_1 - 2x_2 = 0$

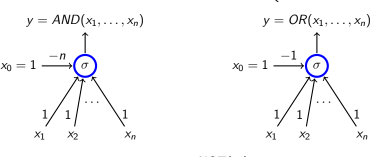
The output neuron performs an intersection of half-spaces.

Boolean functions

Activation function: unit step function
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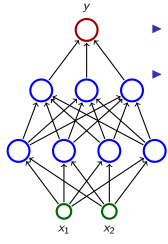


$$y = NOT(x_1)$$

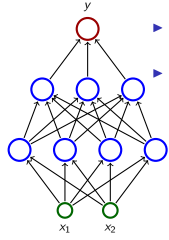
$$x_0 = 1 \xrightarrow{0} \xrightarrow{\sigma}$$

$$-1 \uparrow$$

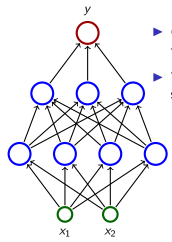
$$x_1$$



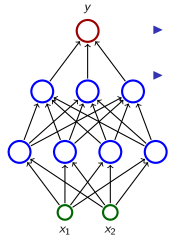
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 - The third layer may, e.g., make unions of some convex sets.

Consider a triangle T in \mathbb{R}^2 determined by three vertices (-1,-1),(1,-1),(-1,2).

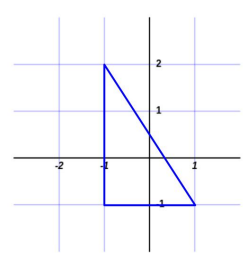
Give an example of a multilayer perceptron (MLP) with two input neurons and a single output neuron computing the function $F: \mathbb{R}^2 \to \{0,1\}$ defined as follows:

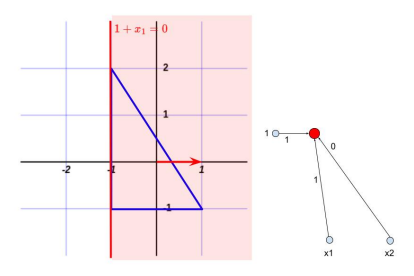
 $F(x_1,x_2)=1$ iff (x_1,x_2) lies either inside, or on the border of T

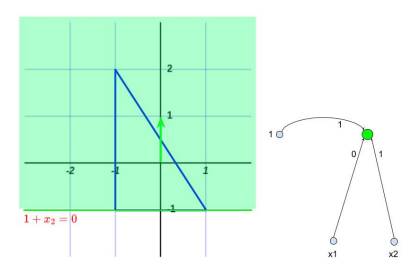
All activation functions in the network should be

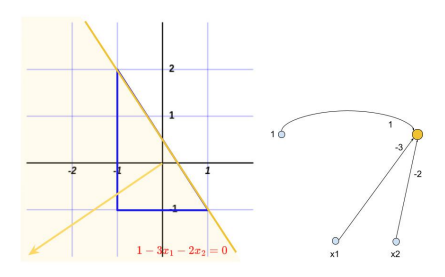
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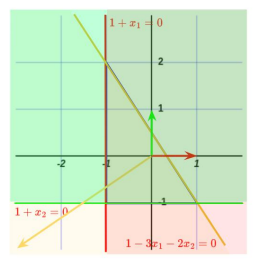
Homework: Consider $F(x_1, x_2) = 1$ iff (x_1, x_2) lies inside of T (but *not* on the border)

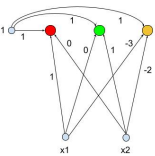


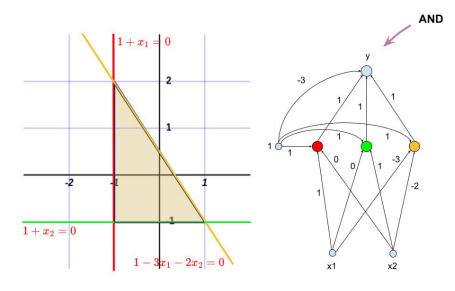












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An efficient way of using the gradient descent was published in 1986!

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▶ Fixing weights of all neurons, the network computes a function $F[\vec{w}]: \mathbb{R}^{|X|} \to \mathbb{R}^{|Y|}$ as follows: Assign values of a given vector $\vec{x} \in \mathbb{R}^{|X|}$ to the input neurons, evaluate the network, then $F[\vec{w}](\vec{x})$ is the vector of values of the output neurons.

Here, we implicitly assume a fixed ordering on input and output vectors.

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▶ Given a set *D* of training examples:

$$D = \left\{ \left(\vec{x}_k, \vec{d}_k \right) \mid k = 1, \dots, p \right\}$$

Here $\vec{x}_k \in \mathbb{R}^{|X|}$ and $\vec{d}_k \in \mathbb{R}^{|Y|}$. We write d_{kj} to denote the value in \vec{d}_k corresponding to the output neuron j.

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Here $\vec{x}_k \in \mathbb{R}^{|X|}$ and $\vec{d}_k \in \mathbb{R}^{|Y|}$. We write d_{kj} to denote the value in \vec{d}_k corresponding to the output neuron j.

Error Function: $E(\vec{w})$ where \vec{w} is a vector of all weights in the network. The choice of E depends on the solved task (classification vs regression etc.).

Example (Squared error):

$$E(\vec{w}) = \sum_{k=1}^{p} E_k(\vec{w})$$

where

$$E_k(\vec{w}) = \frac{1}{2} \sum_{j \in Y} (y_j [\vec{w}] (\vec{x}_k) - d_{kj})^2$$

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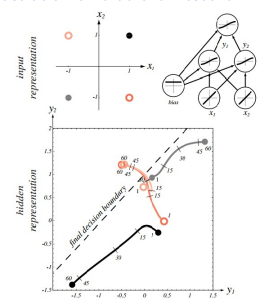
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Note that $\frac{\partial E}{\partial w_{ji}}(\vec{w}^{(t)})$ is a component of ∇E , i.e., the weight change in the step t+1 can be written as follows: $\vec{w}^{(t+1)} = \vec{w}^{(t)} - \varepsilon(t) \cdot \nabla E(\vec{w}^{(t)})$.

Illustration of Gradient Descent - XOR



Source: Pattern Classification (2nd Edition); Richard O. Duda, Peter E. Hart, David G. Stork

Stochastic Gradient Descent (SGD)

Assume that $E(\vec{w}) = \sum_{k=1}^{p} E_k(\vec{w})$ where $E_k(\vec{w})$ is an error w.r.t. the single training example (\vec{x}_k, \vec{d}_k) .

- weights in $\vec{w}^{(0)}$ are randomly initialized to values close to 0
- ▶ in the step t+1 (here t=0,1,2...), weights $\vec{w}^{(t+1)}$ are computed as follows:
 - ▶ Choose (randomly) a set of training examples $T \subseteq \{1, ..., p\}$
 - Compute

$$\vec{w}^{(t+1)} = \vec{w}^{(t)} + \Delta \vec{w}^{(t)}$$

where

$$\Delta \vec{w}^{(t)} = -\varepsilon(t) \cdot \sum_{k \in T} \nabla E_k(\vec{w}^{(t)})$$

- ▶ $0 < \varepsilon(t) \le 1$ is a *learning rate* in step t + 1
- $\triangleright \nabla E_k(\vec{w}^{(t)})$ is the gradient of the error of the example k

Note that the random choice of the minibatch is typically implemented by randomly shuffling all data and then choosing minibatches sequentially.

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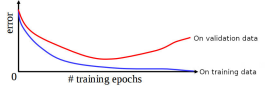
Comments on Training Algorithm

- Not guaranteed to converge to zero training error, may converge to a local minimum or oscillate indefinitely.
- In practice, does converge to low error for many large networks on (big) real data.
- Many epochs (thousands) may be required, hours or days of training for large networks.

There are many issues concerning learning efficiency (data normalization, selection of activation functions, weight initialization, learning rate, efficiency of the gradient descent itself, etc.) – see PV021.

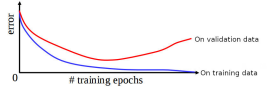
Overfitting

▶ Due to their expressive power, neural networks are quite sensitive to overfitting.



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► Keep a hold-out validation set and test the error of the network on this set after every epoch. Stop training when additional epochs increase the validation error.

The validation error can be measured by completely different means than the training error E.

Hidden Neurons Representations

Trained hidden neurons can be seen as newly constructed features.

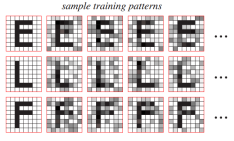
E.g., in a two-layer network used for classification, the hidden layer transforms the input so that important features become explicit (and hence the result may become linearly separable).

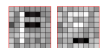
Hidden Neurons Representations

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E.g., in a two-layer network used for classification, the hidden layer transforms the input so that important features become explicit (and hence the result may become linearly separable).

Consider a two-layer MLP, 64-2-3, for classifying letters (three output neurons, each corresponding to one of the letters).





learned input-to-hidden weights

Optimal Architecture?

For MLP: Too few hidden neurons prevent the network from adequately fitting the data. Too many hidden units can result in overfitting.

(There are advanced methods that prevent overfitting even for rich models, such as regularization, where the error function penalizes overfitting – see PV021.)

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- For MLP: Too few hidden neurons prevent the network from adequately fitting the data. Too many hidden units can result in overfitting.
 - (There are advanced methods that prevent overfitting even for rich models, such as regularization, where the error function penalizes overfitting see PV021.)
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Optimal Architecture?

- For MLP: Too few hidden neurons prevent the network from adequately fitting the data. Too many hidden units can result in overfitting.
 - (There are advanced methods that prevent overfitting even for rich models, such as regularization, where the error function penalizes overfitting see PV021.)
- There are (almost) infinitely many types of architectures of neural networks (convolutional, recurrent, transformers, adversarial, etc.) suitable for various tasks.
- ► **Transfer learning:** Start with a known solution to a related problem.

Simplified view: Preserve lower parts of the network trained to solve the related problem (feature extractors). Add your top part and then train only the new top part (or train the whole network but carefully).

How to Choose Activation Functions & Error

▶ **Hidden neurons:** "Almost" linear activations such as (leaky) ReLU $(y = max(0, \xi))$

Better than sigmoidal that saturate more often.

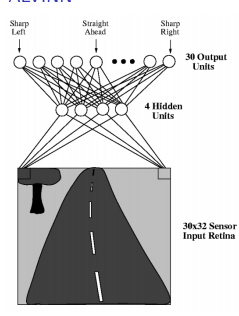
- **Output neurons:** Single output:
 - Regression: Typically "linear" output, i.e., no activation on the output neuron.
 - ▶ Binary classification: Logistic sigmoid $y = 1/(1 + e^{-\xi})$
- **Error:** Single output:
 - Regression: (Mean) squared error
 - Binary classification: Binary cross-entropy

For multiple outputs and classification, use softmax output and cross-entropy.

Applications

- Image recognition, segmentation, etc.
- Machine translation and other text processing
- ► Text generation, image generation, movie generation, theatre plays generation
- Text to Speech and vice versa
- ► Finance, business predictions, fraud detection
- Game playing (backgammon is a classic example, AlphaGo is the famous one, computer games are the big ones, **bridge** is the hard one)
- (artificial brain and intelligence)
- ..

Text and image processing are possibly the most advanced deep learning applications.





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- Direction corresponds to the center of gravity.

, I.e., output neurons are considered as points of mass evenly distributed along a line. The weight of each neuron corresponds to its value.

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 - $ightharpoonup ec{x}_k = {\sf image} \ {\sf of} \ {\sf the} \ {\sf road}$
 - $\vec{d}_k \approx$ corresponding direction of the steering wheel set by the driver
- ▶ the values $\vec{d_k}$ computed using Gaussian distribution:

$$d_{ki} = e^{-D_i^2/10}$$

Where D_i is the distance between the *i*-th output from the one corresponding to the steering wheel's real direction.

(The authors claimed this approach is better than the binary output because similar road directions induce similar driver reactions.)

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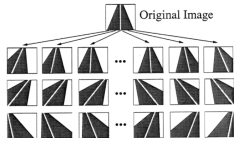
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 - turn the learning off momentarily, deviate from the right track, then turn on the learning and let the network learn how to solve the situation.
 - let the driver go crazy! (a bit dangerous, expensive, unreliable)
- Images are very similar (the network sees the road from the right lane), overfitting.

Problems with too good driver were solved as follows:

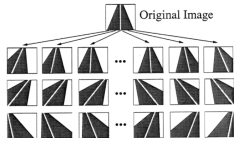
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every image of the road has been has been transformed to 15 slightly different copies



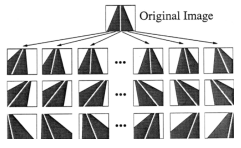
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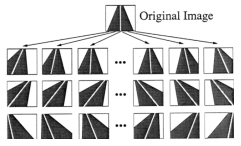
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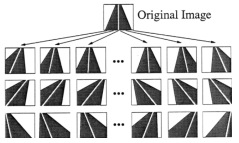


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▶ the system has a buffer of 200 images (including the 15 copies of the current one), in every round trains on these images

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The repetitiveness of images was solved as follows:

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- ▶ afterward, a new image is captured, 15 copies made, and these new 15 substitute 15 selected from the buffer (10 with the smallest training error, five randomly)

ALVINN - Training

▶ gradient descent

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The result:

➤ Training took 5 minutes, and the speed was 4 miles per hour (The speed was limited by the hydraulic controller of the steering wheel, not the learning algorithm.)

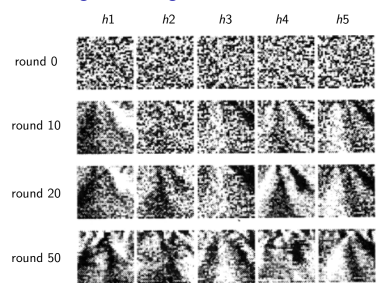
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The result:

- Training took 5 minutes, and the speed was 4 miles per hour (The speed was limited by the hydraulic controller of the steering wheel, not the learning algorithm.)
- ► ALVINN was able to go through roads it had never "seen" and in different weather

ALVINN - Weight Learning



Here $h1, \ldots, h5$ are values of hidden neurons.

Backpropagation

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To implement a single step of the gradient descent, we need to compute the partial derivatives $\frac{\partial E}{\partial w_{ii}}$ of E w.r.t. all weights w_{ij} .

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► Single element training set

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- ▶ The error function is the squared error.
 - Assume that 1 is the output neuron,
 - which means that $y_1 = y_1[\vec{w}](x)$ is the output of the network with weights \vec{w} and the input x,
 - and the error on D is then

$$E(\vec{w}) = \frac{1}{2}(y_1 - d)^2$$

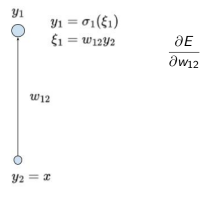
$$D=\{(x,d)\}$$

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$$y_1 = \sigma_1(\xi_1)$$
 $\xi_1 = w_{12}y_2$
 w_{12}
 $y_2 = x$

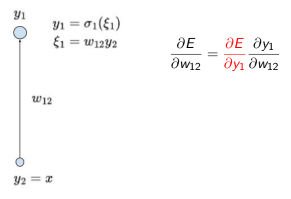
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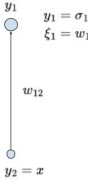
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$$= \frac{\partial E}{\partial y_1} \sigma'_1(\xi_1)y_2$$
Here σ'_1 is just the plain derivative of σ' as a function of a single variable
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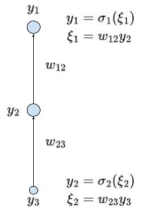
 y_1 $y_1=\sigma_1(\xi_1)$ Here σ_1' is just the plain derivative of σ' as a $\xi_1=w_{12}y_2$ function of a single variable

$$\frac{\partial E}{\partial y_1} = y_1 - d$$

Note that if σ_1 is identity, we obtain exactly the gradient from the linear regression method. Considering σ_1 equal to the logistic sigmoid and E the cross-entropy, we get the logistic regression gradient.

$$D=\{(x,d)\}$$

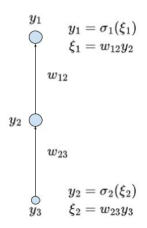
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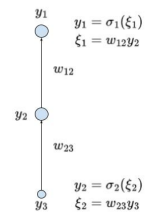
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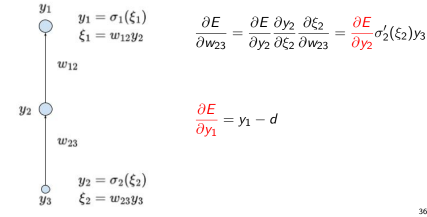


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 w_{14}
 w_{15}
 w_{15}

$$\frac{\partial E}{\partial w_{12}} = \frac{\partial E}{\partial y_1} \frac{\partial y_1}{\partial \xi_1} \frac{\partial \xi_1}{\partial w_{12}} = \frac{\partial E}{\partial y_1} \sigma_1'(\xi_1) y_2$$

$$\frac{\partial E}{\partial w_{13}} = \frac{\partial E}{\partial y_1} \frac{\partial y_1}{\partial \xi_1} \frac{\partial \xi_1}{\partial w_{13}} = \frac{\partial E}{\partial y_1} \sigma_2'(\xi_2) y_3$$

$$D = \{(x,d)\}$$
 $E = \frac{1}{2}(y_1 - d)^2$
 $y_1 = \sigma_1(\xi_1)$
 $y_1 \quad \xi_1 = w_{12}y_2 + w_{13}y_3$
 w_{12}
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 w_{15}
 w_{15}

$$\frac{\partial E}{\partial w_{12}} = \frac{\partial E}{\partial y_1} \frac{\partial y_1}{\partial \xi_1} \frac{\partial \xi_1}{\partial w_{12}} = \frac{\partial E}{\partial y_1} \sigma'_1(\xi_1) y_2$$

$$\frac{\partial E}{\partial w_{13}} = \frac{\partial E}{\partial y_1} \frac{\partial y_1}{\partial \xi_1} \frac{\partial \xi_1}{\partial w_{13}} = \frac{\partial E}{\partial y_1} \sigma_2'(\xi_2) y_3$$

$$\frac{\partial E}{\partial w_{24}} = \frac{\partial E}{\partial y_2} \frac{\partial y_2}{\partial \xi_2} \frac{\partial \xi_2}{\partial w_{24}} = \frac{\partial E}{\partial y_2} \sigma_2'(\xi_2) y_4$$

$$D = \{(x,d)\}$$
 $E = \frac{1}{2}(y_1 - d)^2$
 $y_1 = \sigma_1(\xi_1)$
 $y_1 \quad \xi_1 = w_{12}y_2 + w_{13}y_3$
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$$\frac{\partial E}{\partial w_{12}} = \frac{\partial E}{\partial y_1} \frac{\partial y_1}{\partial \xi_1} \frac{\partial \xi_1}{\partial w_{12}} = \frac{\partial E}{\partial y_1} \sigma_1'(\xi_1) y_2$$

$$\frac{\partial E}{\partial w_{13}} = \frac{\partial E}{\partial y_1} \frac{\partial y_1}{\partial \xi_1} \frac{\partial \xi_1}{\partial w_{13}} = \frac{\partial E}{\partial y_1} \sigma_2'(\xi_2) y_3$$

$$\frac{\partial E}{\partial w_{24}} = \frac{\partial E}{\partial y_2} \frac{\partial y_2}{\partial \xi_2} \frac{\partial \xi_2}{\partial w_{24}} = \frac{\partial E}{\partial y_2} \sigma_2'(\xi_2) y_4$$

$$\frac{\partial E}{\partial w_{34}} = \frac{\partial E}{\partial y_3} \frac{\partial y_3}{\partial \xi_3} \frac{\partial \xi_3}{\partial w_{34}} = \frac{\partial E}{\partial y_3} \sigma_3'(\xi_3) y_4$$

$$D = \{(x,d)\}$$
 $E = \frac{1}{2}(y_1 - d)^2$
 $y_1 = \sigma_1(\xi_1)$
 $y_1 \quad \xi_1 = w_{12}y_2 + w_{13}y_3$
 w_{12}
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$$\frac{\partial E}{\partial w_{12}} = \frac{\partial E}{\partial y_1} \frac{\partial y_1}{\partial \xi_1} \frac{\partial \xi_1}{\partial w_{12}} = \frac{\partial E}{\partial y_1} \sigma_1'(\xi_1) y_2$$

$$\frac{\partial E}{\partial w_{13}} = \frac{\partial E}{\partial y_1} \frac{\partial y_1}{\partial \xi_1} \frac{\partial \xi_1}{\partial w_{13}} = \frac{\partial E}{\partial y_1} \sigma_2'(\xi_2) y_3$$

$$\frac{\partial E}{\partial w_{24}} = \frac{\partial E}{\partial y_2} \frac{\partial y_2}{\partial \xi_2} \frac{\partial \xi_2}{\partial w_{24}} = \frac{\partial E}{\partial y_2} \sigma_2'(\xi_2) y_4$$

$$\frac{\partial E}{\partial w_{34}} = \frac{\partial E}{\partial y_3} \frac{\partial y_3}{\partial \xi_3} \frac{\partial \xi_3}{\partial w_{34}} = \frac{\partial E}{\partial y_3} \sigma_3'(\xi_3) y_4$$

$$\frac{\partial E}{\partial w_{45}} = \frac{\partial E}{\partial y_4} \frac{\partial y_4}{\partial \xi_4} \frac{\partial \xi_3}{\partial w_{45}} = \frac{\partial E}{\partial y_4} \sigma_4'(\xi_4) y_5$$

$$D = \{(x,d)\}$$

$$E = \frac{1}{2}(y_1 - d)^2$$

$$y_1 = \sigma_1(\xi_1)$$

$$y_1 \quad \xi_1 = w_{12}y_2 + \frac{\partial E}{\partial y_1} = y_1 - d$$

$$w_{13}y_3$$

$$w_{12}$$

$$w_{13}$$

$$w_{13}$$

$$y_2$$

$$w_{13}$$

$$y_3$$

$$w_{24}$$

$$y_4 = \sigma_4(\xi_4)$$

$$\xi_4 = w_{45}y_5$$

$$w_{45}$$

$$y_5 = x$$

$$y_2 = \sigma_2(\xi_2)$$

$$y_3 = \sigma_3(\xi_3)$$

$$\xi_2 = w_{24}y_4$$

$$\xi_3 = w_{34}y_4$$

$$D = \{(x,d)\}$$
 $E = \frac{1}{2}(y_1 - d)^2$
 $y_1 = \sigma_1(\xi_1)$
 $y_1 \quad \xi_1 = w_{12}y_2 + w_{13}y_3$
 w_{12}
 w_{13}
 w_{13}
 w_{14}
 w_{15}
 w_{15}

$$\frac{\partial E}{\partial y_1} = y_1 - d$$

$$\frac{\partial E}{\partial y_2} = \frac{\partial E}{\partial y_1} \frac{\partial y_1}{\partial \xi_1} \frac{\partial \xi_1}{\partial y_2} = \frac{\partial E}{\partial y_1} \sigma_1'(\xi_1) w_{12}$$

$$D = \{(x,d)\}$$
 $E = \frac{1}{2}(y_1 - d)^2$
 $y_1 = \sigma_1(\xi_1)$
 $y_1 \quad \xi_1 = w_{12}y_2 + w_{13}y_3$
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 w_{13}
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$$\frac{\partial E}{\partial y_2} = \frac{\partial E}{\partial y_1} \frac{\partial y_1}{\partial \xi_1} \frac{\partial \xi_1}{\partial y_2} = \frac{\partial E}{\partial y_1} \sigma_1'(\xi_1) w_{12}$$

$$\frac{\partial E}{\partial y_3} = \frac{\partial E}{\partial y_1} \frac{\partial y_1}{\partial \xi_1} \frac{\partial \xi_1}{\partial y_3} = \frac{\partial E}{\partial y_1} \sigma_1'(\xi_1) w_{13}$$

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$$\frac{\partial E}{\partial y_3} = \frac{\partial E}{\partial y_1} \frac{\partial y_1}{\partial \xi_1} \frac{\partial \xi_1}{\partial y_3} = \frac{\partial E}{\partial y_1} \sigma_1'(\xi_1) w_{13}$$

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$$\frac{\partial E}{\partial y_4}$$

$$D = \{(x,d)\}$$
 $E = \frac{1}{2}(y_1 - d)^2$
 $y_1 = \sigma_1(\xi_1)$
 $y_1 \quad \xi_1 = w_{12}y_2 + w_{13}y_3$
 $w_{12} \quad w_{13}$
 w_{13}
 $w_{12} \quad w_{13}$
 w_{13}
 $w_{14} \quad w_{15}$
 $w_{15} \quad$

$$\frac{\partial E}{\partial y_1} = y_1 - d$$

$$\frac{\partial E}{\partial y_2} = \frac{\partial E}{\partial y_1} \frac{\partial y_1}{\partial \xi_1} \frac{\partial \xi_1}{\partial y_2} = \frac{\partial E}{\partial y_1} \sigma_1'(\xi_1) w_{12}$$

$$\frac{\partial E}{\partial y_3} = \frac{\partial E}{\partial y_1} \frac{\partial y_1}{\partial \xi_1} \frac{\partial \xi_1}{\partial y_3} = \frac{\partial E}{\partial y_1} \sigma_1'(\xi_1) w_{13}$$

$$\frac{\partial E}{\partial y_4} = \frac{\partial E}{\partial y_2} \frac{\partial y_2}{\partial y_4} + \frac{\partial E}{\partial y_3} \frac{\partial y_3}{\partial y_4}$$

$$D = \{(x,d)\}$$
 $E = \frac{1}{2}(y_1 - d)^2$
 $y_1 = \sigma_1(\xi_1)$
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 w_{12}
 w_{13}
 w_{13}
 w_{14}
 w_{15}
 w_{15}

$$\frac{\partial E}{\partial y_1} = y_1 - d$$

$$\frac{\partial E}{\partial y_2} = \frac{\partial E}{\partial y_1} \frac{\partial y_1}{\partial \xi_1} \frac{\partial \xi_1}{\partial y_2} = \frac{\partial E}{\partial y_1} \sigma_1'(\xi_1) w_{12}$$

$$\frac{\partial E}{\partial y_3} = \frac{\partial E}{\partial y_1} \frac{\partial y_1}{\partial \xi_1} \frac{\partial \xi_1}{\partial y_3} = \frac{\partial E}{\partial y_1} \sigma_1'(\xi_1) w_{13}$$

$$\frac{\partial E}{\partial y_4} = \frac{\partial E}{\partial y_2} \frac{\partial y_2}{\partial y_4} + \frac{\partial E}{\partial y_3} \frac{\partial y_3}{\partial y_4}
= \frac{\partial E}{\partial y_2} \frac{\partial y_2}{\partial \xi_2} \frac{\partial \xi_2}{\partial y_4} + \frac{\partial E}{\partial y_3} \frac{\partial y_3}{\partial \xi_3} \frac{\partial \xi_3}{\partial y_4}$$

$$D = \{(x,d)\}$$
 $E = \frac{1}{2}(y_1 - d)^2$
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$$\frac{\partial E}{\partial y_1} = y_1 - d$$

$$\frac{\partial E}{\partial y_2} = \frac{\partial E}{\partial y_1} \frac{\partial y_1}{\partial \xi_1} \frac{\partial \xi_1}{\partial y_2} = \frac{\partial E}{\partial y_1} \sigma_1'(\xi_1) w_{12}$$

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$$\frac{\partial E}{\partial y_4} = \frac{\partial E}{\partial y_2} \frac{\partial y_2}{\partial y_4} + \frac{\partial E}{\partial y_3} \frac{\partial y_3}{\partial y_4}
= \frac{\partial E}{\partial y_2} \frac{\partial y_2}{\partial \xi_2} \frac{\partial \xi_2}{\partial y_4} + \frac{\partial E}{\partial y_3} \frac{\partial y_3}{\partial \xi_3} \frac{\partial \xi_3}{\partial y_4}
= \frac{\partial E}{\partial y_2} \sigma_2'(\xi_2) w_{24} + \frac{\partial E}{\partial y_3} \sigma_3'(\xi_3) w_{34}$$

MLP - Gradient Computation

Under our simplifying assumptions $D = \{(x, d)\}$ and $E = (y_1 - d)^2$ the gradient computation proceeds as follows:

Applying the chain rule, we obtain

$$\frac{\partial E_k}{\partial w_{ji}} = \frac{\partial E_k}{\partial y_j} \cdot \sigma'_j(\xi_j) \cdot y_i$$

where (after more applications of the chain rule)

$$\frac{\partial E_k}{\partial y_1} = y_1 - d$$

Keep in mind that 1 is the only output neuron which means that y_1 is the value of the network.

$$\frac{\partial E_k}{\partial y_j} = \sum_{r \in j^{\to}} \frac{\partial E_k}{\partial y_r} \cdot \sigma'_r(\xi_r) \cdot w_{rj} \qquad \text{for } j \in Z \setminus (Y \cup X)$$

Here $y_r = y[\vec{w}](x)$ where \vec{w} are the current weights and x is the example input.

MLP - Gradient Computation - General!

Let us drop our simplifying assumptions!

▶ Given a set *D* of training examples:

$$D = \left\{ \begin{pmatrix} \vec{x}_k, \vec{d}_k \end{pmatrix} \mid k = 1, \dots, p \right\}$$

Here $\vec{x}_k \in \mathbb{R}^{|X|}$ and $\vec{d}_k \in \mathbb{R}^{|Y|}$. We write d_{kj} to denote the value in \vec{d}_k corresponding to the output neuron j.

MLP - Gradient Computation - General!

Let us drop our simplifying assumptions!

▶ Given a set *D* of training examples:

$$D = \left\{ \left(\vec{x}_k, \vec{d}_k \right) \mid k = 1, \dots, p \right\}$$

Here $\vec{x}_k \in \mathbb{R}^{|X|}$ and $\vec{d}_k \in \mathbb{R}^{|Y|}$. We write d_{kj} to denote the value in \vec{d}_k corresponding to the output neuron j.

Error Function: $E(\vec{w})$ where \vec{w} is a vector of all weights in the network. The choice of E depends on the solved task (classification vs regression, etc.).

Example (Squared error):

$$E(\vec{w}) = \sum_{k=1}^{p} E_k(\vec{w})$$

where

$$E_k(\vec{w}) = \frac{1}{2} \sum_{i \in Y} (y_j[\vec{w}](\vec{x}_k) - d_{kj})^2$$

MLP - Gradient Computation

For every weight w_{ii} we have (obviously)

$$\frac{\partial E}{\partial w_{ji}} = \sum_{k=1}^{p} \frac{\partial E_k}{\partial w_{ji}}$$

So now it suffices to compute $\frac{\partial E_k}{\partial w_{ji}}$, that is the error for a fixed training example (\vec{x}_k, \vec{d}_k) .

MLP - Gradient Computation

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Applying the chain rule, we obtain

$$\frac{\partial E_k}{\partial w_{ji}} = \frac{\partial E_k}{\partial y_j} \cdot \sigma'_j(\xi_j) \cdot y_i$$

where (more applications of the chain rule)

 $\frac{\partial E_k}{\partial y_j}$ is computed directly for the output neurons $j \in Y$

$$\frac{\partial \mathsf{E}_k}{\partial y_j} = \sum_{r \in i \to j} \frac{\partial \mathsf{E}_k}{\partial y_r} \cdot \sigma_r'(\xi_r) \cdot w_{rj} \qquad \text{for } j \in Z \setminus (Y \cup X)$$

(Here $y_r = y[\vec{w}](\vec{x}_k)$ where \vec{w} are the current weights and \vec{x}_k is the input of the k-th training example.)

MLP - Backpropagation

Input: A training set $D = \left\{ \left(\vec{x}_k, \vec{d}_k \right) \mid k = 1, \dots, p \right\}$ and the current vector of weights \vec{w} .

Note that the backprop. is repeated in every iteration of the gradient descent!

Evaluate all values y_i of neurons using the standard bottom-up procedure with the input \vec{x}_k .

MLP - Backpropagation

Input: A training set $D = \left\{ \left(\vec{x}_k, \vec{d}_k \right) \mid k = 1, \dots, p \right\}$ and the current vector of weights \vec{w} .

Note that the backprop. is repeated in every iteration of the gradient descent!

- Evaluate all values y_i of neurons using the standard bottom-up procedure with the input \vec{x}_k .
- For every training example $(\vec{x_k}, \vec{d_k})$ compute $\frac{\partial E_k}{\partial y_j}$ using backpropagation through layers top-down:
 - For all $j \in Y$ compute $\frac{\partial E_k}{\partial y_j}$ by taking the derivative of the error. e.g., in the case of the squared error, we have $\frac{\partial E_k}{\partial y_i} = y_j d_{kj}$.

MLP - Backpropagation

Input: A training set $D = \{ (\vec{x_k}, \vec{d_k}) \mid k = 1, ..., p \}$ and the current vector of weights \vec{w} .

Note that the backprop. is repeated in every iteration of the gradient descent!

- Evaluate all values y_i of neurons using the standard bottom-up procedure with the input \vec{x}_k .
- For every training example $(\vec{x_k}, \vec{d_k})$ compute $\frac{\partial E_k}{\partial y_j}$ using backpropagation through layers top-down:
 - For all $j \in Y$ compute $\frac{\partial E_k}{\partial y_j}$ by taking the derivative of the error. e.g., in the case of the squared error, we have $\frac{\partial E_k}{\partial y_i} = y_j d_{kj}$.
 - ▶ In the layer ℓ , assuming that $\frac{\partial E_k}{\partial y_r}$ has been computed for all neurons r in the layer $\ell + 1$, compute

$$\frac{\partial E_k}{\partial y_j} = \sum_{r \in i \to j} \frac{\partial E_k}{\partial y_r} \cdot \sigma'_r(\xi_r) \cdot w_{rj}$$

for all j from the ℓ -th layer. Here σ'_r is the derivative of σ_r .

Output:
$$\frac{\partial E}{\partial w_{ji}} = \sum_{k=1}^{p} \frac{\partial E_k}{\partial w_{ji}}$$
.

MLP Learning Example

Training set:

$$y_1 = \sigma_1(\xi_1)$$
 $D = \{(x,d)\} = \{(1,1)\}$
 $O = 1/(1+e^{-\xi_1})$ That is
 $y_3 = x = 1$
 $O = \{(x,d)\} = \{(1,1)\}$
 $O = 1/(1+e^{-\xi_1})$ That is
 $O = 1/(1+e^{-\xi_1})$ Th

MLP Learning Example

Training set:
$$D = \{(x,d)\} = \{(1,1)\}$$

$$= x$$

 $=-\log(v_1)$

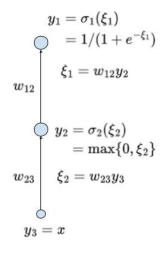
Assume the initial weight vector $\vec{w}^{(0)} = (w_{12}^{(0)}, w_{23}^{(0)}) = (\frac{1}{4}, 2).$

Consider the learning rate $\varepsilon = 0.5$.

Error cross-entropy:

 $E(\vec{w}) = -(d \log(y_1) + (1 - d) \log(1 - y_1))$

MLP Learning Example - Gradient Descent



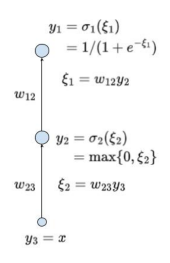
To make the gradient descent step:

$$w_{12}^{(1)} = w_{12}^{(0)} - \varepsilon \frac{\partial E}{\partial w_{12}} (\vec{w}^{(0)})$$

$$w_{23}^{(1)} = w_{23}^{(0)} - \varepsilon \frac{\partial E}{\partial w_{23}} (\vec{w}^{(0)})$$

we need to compute the partial derivatives $\frac{\partial E}{\partial w_{12}}$ and $\frac{\partial E}{\partial w_{23}}$.

MLP Learning Example - Forward Pass



We have
$$x = 1, w_{12}^{(0)} = 1/4, w_{23}^{(0)} = 2$$

MLP Learning Example – Forward Pass

$$y_1 = \sigma_1(\xi_1)$$
 $0 = 1/(1 + e^{-\xi_1})$
 $\xi_1 = w_{12}y_2$
 w_{12}
 $y_2 = \sigma_2(\xi_2)$
 $= \max\{0, \xi_2\}$
 w_{23}
 $\xi_2 = w_{23}y_3$
 $y_3 = x$

We have
$$x = 1$$
, $w_{12}^{(0)} = 1/4$, $w_{23}^{(0)} = 2$

First, compute the forward pass

$$y_3 = x = 1$$

$$\xi_2 = w_{23}^{(0)} y_3 = 2y_3 = 2$$

$$y_2 = \max\{0, \xi_2\} = 2$$

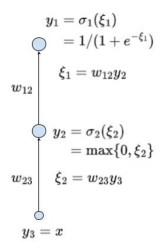
$$\xi_1 = w_{12}^{(0)} y_2 = \frac{1}{4} 2 = \frac{1}{2}$$

$$y_1 = 1/(1 + e^{-\xi_1}) = 1/(1 + e^{-(1/2)})$$

$$= 0.6225$$

MLP Learning Example - Backward Pass

We have
$$w_{12}^{(0)} = 1/4$$
, $w_{23}^{(0)} = 2$, $y_1 = 0.6225$, $y_2 = 2$, $y_3 = 1$.



MLP Learning Example - Backward Pass

$$y_1 = \sigma_1(\xi_1)$$
 $\bigcirc = 1/(1+e^{-\xi_1})$
 $\xi_1 = w_{12}y_2$
 w_{12}
 $y_2 = \sigma_2(\xi_2)$
 $= \max\{0, \xi_2\}$
 w_{23}
 $\xi_2 = w_{23}y_3$
 $y_3 = x$

We have
$$w_{12}^{(0)} = 1/4$$
, $w_{23}^{(0)} = 2$, $y_1 = 0.6225$, $y_2 = 2$, $y_3 = 1$.

Proceed with the backward pass:

$$\frac{\partial E}{\partial y_1} = \frac{\partial (-\log(y_1))}{\partial y_1} = -\frac{1}{y_1} = -1.6065$$

MLP Learning Example - Backward Pass

We have $w_{12}^{(0)} = 1/4$, $w_{23}^{(0)} = 2$, $y_1 = 0.6225$, $y_2 = 2$, $y_3 = 1$.

Proceed with the backward pass:

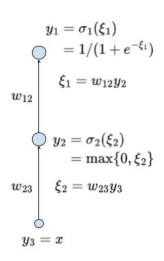
$$\frac{\partial E}{\partial y_1} = \frac{\partial (-\log(y_1))}{\partial y_1} = -\frac{1}{y_1} = -1.6065$$

Since
$$\sigma_1' = \sigma_1(1-\sigma_1)$$

$$\frac{\partial E}{\partial y_2} = \frac{\partial E}{\partial y_1} \sigma_1'(\xi_1) w_{12}^{(0)}
= \frac{\partial E}{\partial y_1} \sigma_1(\xi_1) (1 - \sigma_1(\xi_1)) w_{12}^{(0)}
= \frac{\partial E}{\partial y_1} y_1 (1 - y_1) w_{12}^{(0)}
= -1.6065 \cdot 0.6225 \cdot 0.3775 \cdot (1/4)
= -0.09438$$

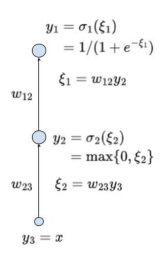
$$y_1 = \sigma_1(\xi_1)$$
 $= 1/(1 + e^{-\xi_1})$
 $\xi_1 = w_{12}y_2$
 w_{12}
 $y_2 = \sigma_2(\xi_2)$
 $= \max\{0, \xi_2\}$
 w_{23}
 $\xi_2 = w_{23}y_3$
 $y_3 = x$

MLP Learning Example - The Gradient



We have
$$\begin{aligned} &w_{12}^{(0)} = 1/4, w_{23}^{(0)} = 2, y_1 = 0.6225, y_2 = \\ &2, y_3 = 1, \frac{\partial E}{\partial y_1} = -1.6065, \frac{\partial E}{\partial y_2} = -0.09438. \end{aligned}$$

MLP Learning Example - The Gradient



We have
$$w_{12}^{(0)}=1/4, w_{23}^{(0)}=2, y_1=0.6225, y_2=2, y_3=1, \frac{\partial E}{\partial y_1}=-1.6065, \frac{\partial E}{\partial y_2}=-0.09438.$$

Compute derivatives of E w.r.t. weights:

$$\frac{\partial E}{\partial w_{12}} = \frac{\partial E}{\partial y_1} \sigma'_1(\xi_1) y_2$$
$$= \frac{\partial E}{\partial y_1} y_1 (1 - y_1) y_2$$
$$= -0.755$$

MLP Learning Example - The Gradient

$$y_1 = \sigma_1(\xi_1) \ = 1/(1+e^{-\xi_1}) \ \xi_1 = w_{12}y_2 \ y_2 = \sigma_2(\xi_2) \ = \max\{0,\xi_2\} \ \xi_2 = w_{23}y_3 \ y_3 = x$$

We have $w_{12}^{(0)} = 1/4$, $w_{23}^{(0)} = 2$, $y_1 = 0.6225$, $y_2 = 2$, $y_3 = 1$, $\frac{\partial E}{\partial y_1} = -1.6065$, $\frac{\partial E}{\partial y_2} = -0.09438$.

Compute derivatives of E w.r.t. weights:

$$\frac{\partial E}{\partial w_{12}} = \frac{\partial E}{\partial y_1} \sigma_1'(\xi_1) y_2$$

$$= \frac{\partial E}{\partial y_1} y_1 (1 - y_1) y_2$$

$$= -0.755$$

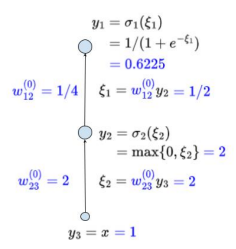
$$\frac{\partial E}{\partial w_{23}} = \frac{\partial E}{\partial y_2} \sigma_2'(\xi_2) y_3$$

$$= \frac{\partial E}{\partial y_2} y_3$$

$$= -0.09438$$

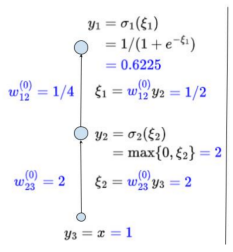
Backpropagation – Example – Summary

Forward pass (bottom up)



Backpropagation – Example – Summary

Forward pass (bottom up)



Backward pass (top down)

$$y_{1} = \sigma_{1}(\xi_{1})$$

$$= 1/(1 + e^{-\xi_{1}})$$

$$= 0.6225$$

$$\xi_{1} = w_{12}^{(0)} y_{2} = 1/2$$

$$y_{2} = \sigma_{2}(\xi_{2})$$

$$= \max\{0, \xi_{2}\} = 2$$

$$\xi_{2} = w_{23}^{(0)} y_{3} = 2$$

$$\frac{\partial E}{\partial w_{12}} = \frac{\partial E}{\partial y_{1}} y_{1} (1 - y_{1}) y_{2}$$

$$= -0.755$$

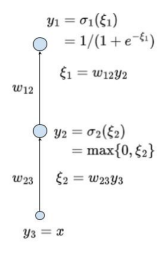
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Note that WE HAVE NOT YET CHANGED ANY WEIGHTS!

MLP Learning Example - Gradient Descent Step



So **ONLY NOW** we can make the **learning step** and change the weights:

$$w_{12}^{(1)} = w_{12}^{(0)} - \varepsilon \frac{\partial E}{\partial w_{12}} (\vec{w}^{(0)})$$

$$= \frac{1}{4} - 0.5 \cdot (-0.755)$$

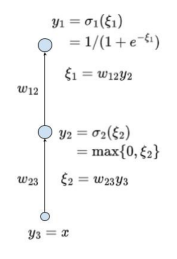
$$= 0.627$$

$$w_{23}^{(1)} = w_{23}^{(0)} - \varepsilon \frac{\partial E}{\partial w_{23}} (\vec{w}^{(0)})$$

$$= 2 - 0.5 \cdot (-0.09438)$$

$$= 2.047$$

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We have made just a **single** gradient descent step!

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- ► Training of neural networks in practice is tricky due to many reasons comprising in particular
 - ▶ highly complex non-linear shape of the error function,
 - tendency to overfit very quickly,
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An advice: Always concentrate the main effort on the solved problem formulation and, afterward, on the data you have at your disposal (and honestly separate the Test set right at the beginning).

Cybenko's theorem shows that two-layer networks are omnipotent – such results nearly killed NN when support vector machines were found to be easier to train in 00's.

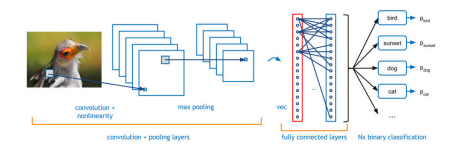
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- ... but how to train them? The gradient descent suffers from a so-called vanishing gradient; intuitively, updates of weights in lower layers are very slow.
- ▶ In 2006, a solution was found by Hinton et al.:
 - ▶ Use *unsupervised* methods to initialize the weights layer by layer to capture important data features.
 - More precisely: The lowest hidden layer learns patterns in data, the second lowest learns patterns in data transformed through the first layer, and so on.

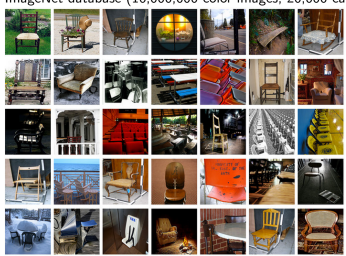
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 More precisely: The lowest hidden layer learns patterns in data, the second lowest learns patterns in data transformed through the first layer, and so on.
 - Then use a supervised learning algorithm to only *fine tune* the weights to the desired input-output behavior.
- ... but the actual revolution started with convolutional networks trained on several GPUs.

Convolutional network



ImageNet Large-Scale Visual Recognition Challenge (ILSVRC)

ImageNet database (16,000,000 color images, 20,000 categories)



ImageNet Large-Scale Visual Recognition Challenge (ILSVRC)

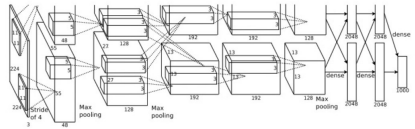
Competition in classification over a subset of images from ImageNet.

In 2012, training saw 1,200,000 images and 1000 categories. Validation set 50,000, Test set 150,000.

Many images contain several objects \rightarrow typical rule is top-5 highest probability assigned by the net.

KSH net

ImageNet classification with deep convolutional neural networks, by Alex Krizhevsky, Ilya Sutskever, and Geoffrey E. Hinton (2012).



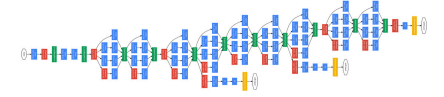
Trained on two GPUs (NVIDIA GeForce GTX 580)

Results:

- Accuracy 84.7% in top-5 (second best alg. at the time: 73.8%)
- ▶ 63.3% in "perfect" classification (top-1)

The same set of images as in 2012, top-5 criterium.

GoogLeNet: deep convolutional net, 22 layers



Results:

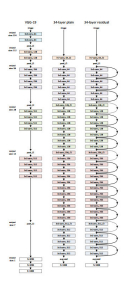
▶ 93.33% in top-5

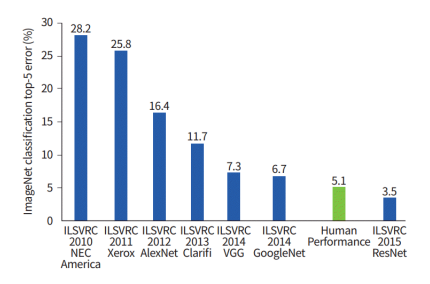
Superhuman power?

Superhuman GoogLeNet?!

Andrei Karpathy: ...the task of labeling images with 5 out of 1000 categories quickly turned out to be highly challenging, even for some friends in the lab who have been working on ILSVRC and its classes for a while. First, we thought we would put it up on [Amazon Mechanical Turk]. Then, we thought we could recruit paid undergrads. Then, I organized a labeling party of intense labeling effort only among the (expert labelers) in our lab. Then, I developed a modified interface that used GoogLeNet predictions to prune the number of categories from 1000 to only about 100. It was still too hard - people kept missing categories and getting up to ranges of 13-15% error rates. In the end, I realized that to get anywhere competitively close to GoogLeNet, it would be most efficient if I sat down and went through the painfully long training process and the subsequent careful annotation process myself. The labeling happened at a rate of about 1 per minute, but this decreased over time... Some images are easily recognized, while some pictures (such as those of fine-grained breeds of dogs, birds, or monkeys) can require multiple minutes of concentrated effort. I became very good at identifying breeds of dogs... Based on the sample of images I worked on, the GoogLeNet classification error turned out to be 6.8%... In the end, my error turned out to be 5.1%

- ► Microsoft network ResNet: 152 layers, complex architecture
- ► Trained on 8 GPUs
- ▶ 96.43% accuracy in top-5





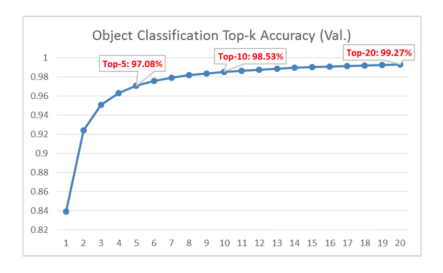
Trumps-Soushen (The Third Research Institute of Ministry of Public Security)

There is no new innovative technology or novelty by Trimps-Soushen.

Ensemble of the pre-trained models from previous years.

Each model is strong at classifying some categories but weak at categorizing others.

Test error: 2.99%



https://towards datascience.com/review-trimps-soushen-winner-in-ilsvrc-2016-image-classification-dfbc 423111 dd and the state of the

Top-20 typical errors

Out of 1458 misclassified images in Top-20:

Error Categories	Numbers	Percentages(%)
Label May Wrong	221	15.16
Multiple Objects (>5)	118	8.09
Non-Obvious Main Object	355	24.35
Confusing Label	206	14.13
Fine-grained Label	258	17.70
Obvious Wrong	234	16.05
Partial Object	66	4.53

https://towards datascience.com/review-trimps-soushen-winner-in-ilsvrc-2016-image-classification-dfbc 423111dd. A state of the control of t

Predict:

1 pencil box

2 diaper

3 bib

4 purse

5 running shoe

Ground Truth: sleeping bag



https://towardsdatascience.com/review-trimps-soushen-winner-in-ilsvrc-2016-image-classification-dfbc423111dd

Predict:

- 1 dock
- 2 submarine
- 3 boathouse
- 4 breakwater
- 5 lifeboat

Ground Truth: paper towel



Predict:

1 bolete

2 earthstar

3 gyromitra

4 hen of the woods

5 mushroom

Ground Truth: stinkhorn



Predict:

1 apron

2 plastic bag

3 sleeping bag

4 umbrella

5 bulletproof vest

Ground Truth: **poncho**



Anomaly Detection

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This lecture presents algorithms detecting specific data instances that may be anomalous w.r.t. the used algorithm.

It is up to the context-knowing user to decide whether such data are anomalous.

What is the Anomaly Detection Good For?

There are two main sources of motivation for anomaly detection in machine learning:

► *Modeling perspective*: Many models are sensitive to anomalous data.

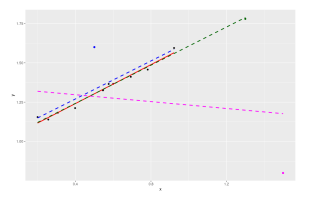
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- ► Application perspective: Many tasks are based on searching for anomalies in data.

Modeling Perspective

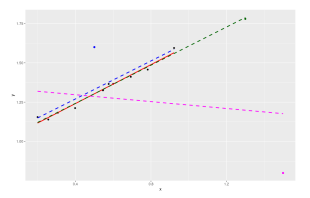
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Modeling Perspective

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On the other hand, we will see that the sensitivity of some models can be used to *detect* the anomalies.

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- Detect atypical natural world events.
- Prediction and understanding of hurricanes, floods, droughts, heat waves, and fires.

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Ecosystem Disturbances

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Medicine

- Unusual patient symptoms or test results may indicate health problems.
- Balancing the need for further tests with the potential costs and risks.

...

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The anomalies mentioned above should be inspected by domain experts and possibly corrected/removed.

Approaches to Anomaly Detection

- ► Model-based techniques:
 - Build a model of data.
 - Anomalies should not fit the model very well.
 For example, using a clustering model, an anomaly may lie far away from larger clusters.
 - ► Alternatively, removing anomalies should have the strongest impact on the model parameters (more on this later).

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- Proximity-based techniques: Given distance between objects, anomalies are objects distant from others.
- Density-based techniques: Estimate the density distribution of the objects. Anomalies would probably be outside of dense regions.

I would count the most recent deep learning-based anomaly detection among the model-based techniques even though a combination of the above approaches is usually used.

We will make the above ideas more precise in the rest of this lecture.

Anomaly Detection Learning

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 - A tumor detection system trained on healthy people detects tumors as anomalies.
 - In this case, we detect anomalies that are dissimilar to normal instances. We may train a model representing the normal instances and detect instances that do not fit the model.
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 - We may have information about (some) normal and some anomalous instances. Here, we can use methods for semi-supervised classification.
- Unsupervised: Create a model of all instances and hope that anomalous instances will still be dissimilar to instances typical for the model.

Task-specific issues comprise:

➤ Single vs. multi-attribute anomaly: The question is whether an object is anomalous due to a single attribute (200 kg person) or a combination of attributes (100 kg person of 1 meter height).

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Another issue is evaluation: How can we determine how good an anomaly detector is?

In the supervised case, we have a ground truth, but usually, we just observe detected anomalies.

Various Approaches to Anomaly Detection

We shall have a look at five approaches to the unsupervised anomaly detection:

- Statistical
- Proximity-based
- Density-based
- Cluster-based
- Autoencoders

The above approaches are also used in the semi-supervised setting where we have a dataset of normal instances. However, some issues discussed further will not appear in this case.

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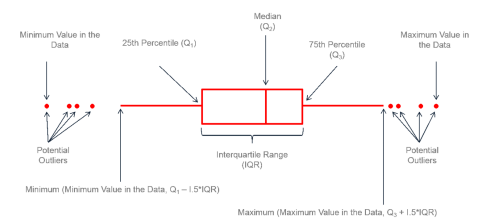
However, in most cases, we do not know P and have to resort to a model of P created using a dataset of feature vectors.

There are many more or less sophisticated tests based on models of P in the literature.

Some use rather advanced statistical methods.

Let us have a look at a few simple examples.

Box Plot



A simple method for outlier detection in the *univariate* case, that is, for values of a single attribute.

Consider a numeric attribute whose values x are normally distributed with mean μ and standard deviation σ . We write $N(\mu, \sigma^2)$.

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$$z = (x - \mu)/\sigma$$

Then z has the distribution N(0,1).

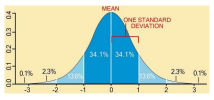
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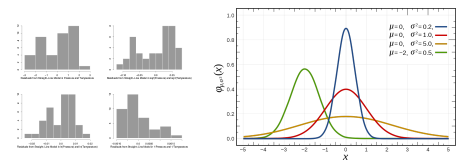
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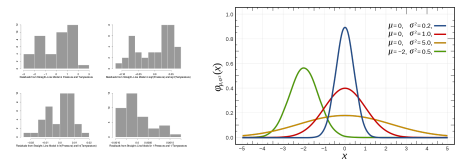
Now, choose c so the probability $P(|z| \ge c)$ is small enough for z to be an outlier w.r.t. N(0,1).



Given an attribute value x, we may decide whether x is an outlier by deciding whether $|z| = |(x - \mu)/\sigma| \ge c$.



Problem 1: The attribute does not have a normal distribution.



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Before starting, use a normality test (observe the histogram, use a specialized test such as Shapiro-Wilk).

Some transformations may sometimes succeed in normalizing an attribute (Box-Cox transformation, etc.)

Different distributions can be used to model the attribute (Log Normal, Weibull, etc.), but be aware of the assumptions of anomaly detection tests!

Problem 2: We typically do not know the population mean μ and the population variance σ^2 .

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They can be estimated from a dataset by the sample mean and the sample variance:

$$\bar{\mu} = \frac{1}{p} \sum_{i=1}^{p} x_i$$
 $s^2 = \frac{1}{p-1} \sum_{i=1}^{p} (x_i - \bar{\mu})^2$

However, then $z=(x-\bar{\mu})/s$ does no longer have the distribution N(0,1).

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Note that $\bar{\mu}$ and s^2 are unbiased estimates of μ and σ^2 . Also, $\bar{\mu}$ converges to μ and s^2 converges to σ^2 with growing sample size.

Problem 3: The outliers distort the estimates $\bar{\mu}$ and s^2 of the mean and the standard deviation.

For example, a millionaire would not look like an outlier in a group containing a billionaire.

There is a circularity here: To get outliers using the normal distribution model, we need to remove the outliers to have a good model.

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There is a circularity here: To get outliers using the normal distribution model, we need to remove the outliers to have a good model.

One possible heuristic is to remove the outliers one by one, starting from the most extreme, hoping the most extreme outlier will be detected in every step.

One such heuristic is the Grubb's test.

Consider a dataset $D = \{x_1, \dots, x_p\}$ of values of a normally distributed attribute and choose $\alpha > 0$.

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Now
$$P(G \ge c_{\alpha,p}) = \alpha$$
 if

$$c_{\alpha,p} = \frac{p-1}{\sqrt{p}} \sqrt{\frac{t^2}{p-2+t^2}}$$

Here t is such a value that makes $P(T \ge t) = \alpha/(2p)$ for T with the t-distribution with p-2 degrees of freedom.

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Grubb's test simply iteratively tests whether $G \ge c_{\alpha,p}$ and, if yes, removes an x_i maximizing $|x_i - \bar{x}|$ from D.

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Mahalanobis distance:

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$$(\vec{x}, \vec{z}) = (\vec{x} - \vec{z}) \Sigma^{-1} (\vec{x} - \vec{z})^{\top}$$

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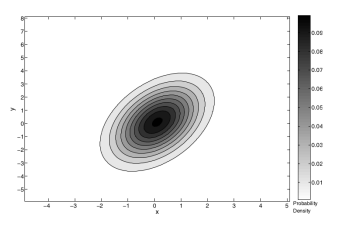
Here Σ is the covariance matrix of the data.

Intuitively, the Mahalanobis distance generalizes the squared Euclidean distance:

$$(\vec{x}-\vec{z})(\vec{x}-\vec{z})^{\top}$$

By taking into account the "spread" of the data.

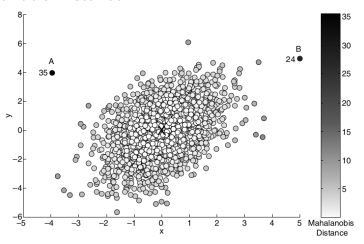
Mahalanobis Distance



Here, we assume multivariate normal data distribution. The covariance matrix is

$$\Sigma = \left(\begin{array}{cc} 1.00 & 0.75 \\ 0.75 & 3.00 \end{array}\right)$$

Mahalanobis Distance



Notice that A has a larger Mahalanobis distance from the cluster's center than B even though it is closer in the Euclidean distance.

The reason is that the density function falls more rapidly in the direction of A than in the direction of B.

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Now, we compute the total likelihood of the dataset before and after removing each element. If the likelihood changes significantly, we have probably eliminated an anomaly.

$$L(U, V) = \left((1 - \alpha)^{|U|} \prod_{x_i \in U} P_M(x_i) \right) \left(\alpha^{|V|} \prod_{x_i \in V} P_A(x_i) \right)$$

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- Randomly choose between normal and anomaly, where the probability of choosing anomaly is α .
- ▶ If normal has been chosen, choose x_i from the distribution M.
- \triangleright If anomaly has been chosen, choose x_i from the distribution A.

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- ▶ If anomaly has been chosen, choose *x_i* from the distribution *A*.

To eliminate the "large" product, we consider the log-likelihood:

$$LL(U, V) = \log(L(U, V))$$

$$= |U| \log(1 - \alpha) + \sum_{x_i \in U} \log P_M(x_i)$$

$$+ |V| \log \alpha + \sum_{x_i \in V} \log P_A(x_i)$$

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$$\Delta_i = |LL(M_k, A_k) - LL(M_k \setminus \{x_i\}, A_k \cup \{x_i\})|$$

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▶ Consider *i* maximizing Δ_i . If $\Delta_i \geq c$, then

$$M_{k+1} = M_k \setminus \{x_i\}$$
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Else, stop.

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Ultimately, the A_k will contain the anomalies the algorithm detects.

Note that we may also move all anomalies detected in a single iteration from M_k to A_k . This would result in a different method (also valid).

Summary of Statistical Anomaly Detection

- ▶ Build on strong statistical foundations.
- Tests are very effective if the dataset is sufficiently large (informative).
- Lots of tests for univariate data, the area has developed for a long time.
- ► Fewer options for multivariate data and problematic for highly dimensional data (curse of dimensionality).

The likelihood-based approach does not assume a particular distribution shape: It can be used with arbitrary models of P_M and P_A , including deep learning ones.

Proximity-Based Methods

Proximity-Based Outlier Detection

Assume a distance measure d. That is, given two feature vectors \vec{x} , \vec{z} their distance is $d(\vec{x}, \vec{z})$.

We consider the Euclidean distance for simplicity.

Definition 2

The outlier score of an object is given by the distance to its k-nearest neighbor.

Proximity-Based Outlier Detection

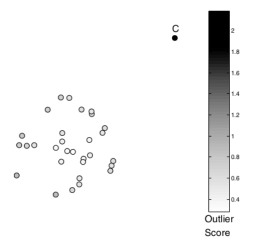
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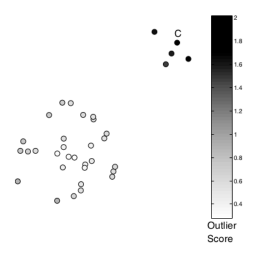
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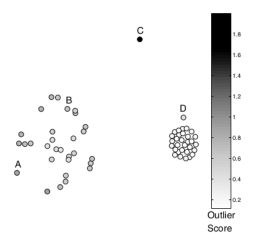
A threshold on the minimum distance of an outlier can be set on a training set.



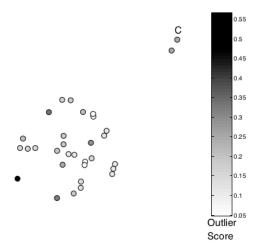
The outlier score is based on the distance to the fifth nearest neighbor.



The outlier score is based on the distance to the fifth nearest neighbor.



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The outlier score is based on the distance to the first nearest neighbor.

Proximity-Based Approaches

- Conceptually simple and easy to implement.
- ▶ Time complexity typically $\mathcal{O}(p^2)$ where p is the number of feature vectors in the dataset.
- Sensitivity to the choice of parameters (the number of neighbors).
- Density/compactness not taken explicitly into account.

Density-Based Methods

Density-Based Approaches

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The outlier score of an object is the inverse of the density around the object.

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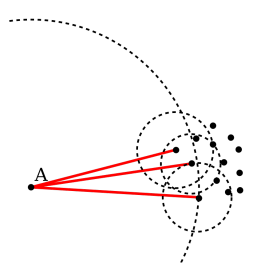
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- computing the average D of the local densities of k-nearest neighbors of A,
- ▶ dividing the average local density with the local density of A, that is, compute \bar{D}/D_A .

The question is, how exactly can the local density be defined?

LOF - Illustration



Here, the density around A is smaller than the densities around the other points.

LOF - Formally

Let us fix $k \in \mathbb{N}$.

Define k-distance(A) as the distance of the k-th nearest neighbor.

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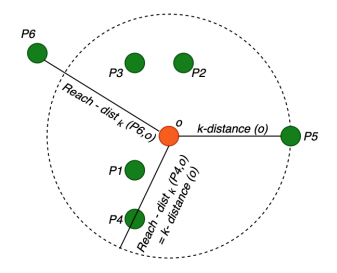
Note that if there are objects in the same distance from A, we may have more than k elements in $N_k(A)$.

Define

reachability-distance_k
$$(A, B) = \max\{k - \text{distance}(B), d(A, B)\}$$

The reachability distance of A from B is the distance between A and B but at least the distance from B to the k-nearest neighbor.

Reachability Distance



LOF

Define local reachability density

$$\operatorname{Ird}_k(A) = \left(\frac{\sum_{B \in N_k(A)} \operatorname{reachability-distance}_k(A, B)}{|N_k(A)|}\right)^{-1}$$

That is the reciprocal of the average reachability distance of A from its k-nearest neighbors.

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We define *local outlier factor* of an object A by

$$LOF_k(A) = \left(\frac{\sum_{B \in N_k(A)} Ird_k(B)}{|N_k(A)|}\right) / Ird_k(A)$$

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$$\mathsf{LOF}_k(A) = \left(\frac{\sum_{B \in N_k(A)} \mathsf{Ird}_k(B)}{|N_k(A)|}\right) / \mathsf{Ird}_k(A)$$

Now

- ▶ LOF_k(A) ≈ 1 similar density as the neighbors have
- ▶ LOF_k(A) < 1 higher density of neighbors than the neighbors have (inlier?)
- ► LOF_k(A) > 1 smaller density of neighbors than the neighbors have (outlier?)

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$$D = \{A, B, C, D\} = \{5, 2, 1, 0\}$$

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	k-distance(.)	d(A,.)	$reach-dist_k(A,.)$
В	2	3	3
C	1	4	4

$$\operatorname{Ird}_k(A) = \left(\frac{1}{2}\left(\operatorname{reach-dist}_k(A,B) + \operatorname{reach-dist}_k(A,C)\right)\right)^{-1} = 2/7$$

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$$N_k(B) = \{C, D\}$$

	k-distance(.)	d(B,.)	$reach-dist_k(B,.)$
С	1	1	1
D	2	2	2

$$\operatorname{Ird}_k(B) = \left(\frac{1}{2}\left(\operatorname{reach-dist}_k(B,C) + \operatorname{reach-dist}_k(B,D)\right)\right)^{-1} = 2/3$$

Consider a dataset

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$$N_k(C) = \{B, D\}$$

	k-distance(.)	d(C,.)	reach-dist $_k(C,.)$
В	2	1	2
D	2	1	2

$$\operatorname{Ird}_k(C) = \left(\frac{1}{2}(\operatorname{\mathsf{reach-dist}}_k(C,B) + \operatorname{\mathsf{reach-dist}}_k(C,D))\right)^{-1} = 1/2$$

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$$N_k(D) = \{B, C\}$$

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В	2	2	2
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$$\operatorname{Ird}_k(A) = 2/7$$

 $\operatorname{Ird}_k(B) = 2/3$
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$$\begin{split} & \operatorname{Ird}_k(A) = 2/7 \\ & \operatorname{Ird}_k(B) = 2/3 \\ & \operatorname{Ird}_k(C) = 1/2 \\ & \operatorname{Ird}_k(D) = 2/3 \end{split}$$

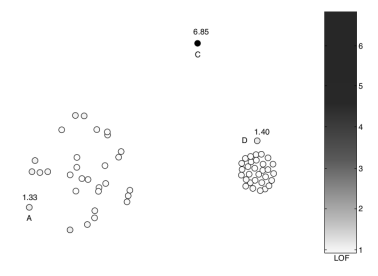
$$\begin{aligned} & \operatorname{LOF}_k(A) = \frac{1}{2} (\operatorname{Ird}_k(B) + \operatorname{Ird}_k(C)) / \operatorname{Ird}_k(A) \\ & = (1/2)(2/3 + 1/2) / (2/7) = 2.041 \end{split}$$

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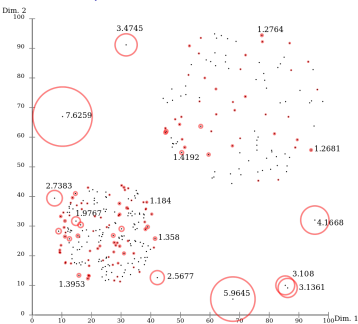
$$\begin{aligned} & \operatorname{LOF}_k(C) = \frac{1}{2} (\operatorname{Ird}_k(B) + \operatorname{Ird}_k(D)) / \operatorname{Ird}_k(B) \\ & = (1/2)(2/3 + 2/3) / (2/3) = 1 \end{aligned}$$

$$\begin{aligned} & \operatorname{LOF}_k(D) = 0.875 \end{aligned}$$

Another Example



Yet Another Example



Comments on Density Based Methods

- As opposed to the distance-based methods, quantifies the distance of the neighborhood of the (potential) outliers.
- ▶ Time complexity still $\mathcal{O}(p^2)$ but may be reduced for low dimensional data (to $\mathcal{O}(p \log p)$) using special data structures.
- ▶ Still need to determine the parameter *k*.

Clustering-Based Methods

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A better approach would be to asses how strongly objects belong to clusters.

Definition 4

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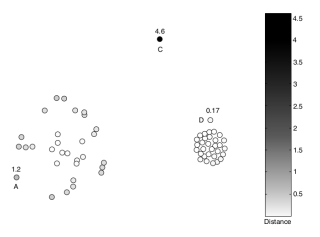
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Depending on the particular clustering algorithm, we have (at least some) information about the cohesion of objects.

Anomaly Detection Using k-Means Clustering

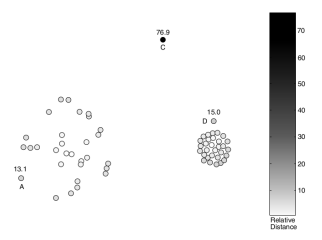


In k-means, the clusters are represented by centroids.

We may measure the anomaly of a given object using the distance to its cluster centroid.

This approach does not take into account the density of clusters.

Anomaly Detection Using k-Means Clustering



Here, we compute the relative distance to the cluster center, that is, the ratio of the object's distance to the centroid and the median distance of all objects of the same cluster to the centroid.

Clustering Based Anomaly Detection

Other algorithms may provide different information (probability density of the clusters, etc.).

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The usual problem: Outliers have an impact on the clustering itself.

Clustering Based Anomaly Detection

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The usual problem: Outliers have an impact on the clustering itself.

Some algorithms can be modified to treat potential outliers specially.

For example, during the k-means clustering, put objects very far away from the current cluster centroids into a special category not used to move the centroids. After every step, test whether some of these objects became close enough to at least one centroid (in which case these objects will be removed from the special category).

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Setting the proper number of clusters is an issue as well.

For anomaly detection, a larger number of smaller clusters may be beneficial as they may be more cohesive. If an object seems to be an outlier with small clusters, it is probably an outlier.

Comments on Clustering Based Methods

Some clustering methods have a sub-quadratic complexity.

Conceptually, clustering complements anomaly detection, so it is natural to compute both together.

On the other hand, the results strongly depend on the number of clusters, and anomalies may distort the clustering.

Autoencoders as Anomaly Detectors

... just a short comment

Neural Networks in Anomaly Detection

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However, what should we do with data with high or even variable dimensions?

- Images
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- Video
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Train a neural network that takes the large input and returns a smaller representation, which (hopefully) preserves crucial features of the input.

An autoencoder consists of two parts:

- $\phi: \mathbb{R}^n \to \mathbb{R}^m$ the encoder
- $\psi: \mathbb{R}^m \to \mathbb{R}^n$ the decoder

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$$\mathcal{T} = \{\vec{x}_1, \dots, \vec{x}_p\}$$

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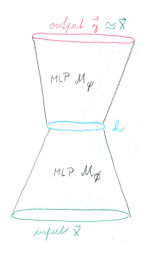
Minimize the reconstruction error

$$E = \sum_{i=1}^{p} (\vec{x}_i - \psi(\phi(\vec{x}_i)))^2$$

Autoencoders - neural networks

Both ϕ and ψ can be represented using MLP \mathcal{M}_{ϕ} and \mathcal{M}_{ψ} , respectively.

 \mathcal{M}_{ϕ} and \mathcal{M}_{ψ} can be connected into a single network.



► Compression/feature extraction – from \vec{x} to \vec{h} .

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- ► Anomaly detection (see the next slide)

Anomaly Detection with Autoencoders

Straightforward approach:

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More general approach:

- Train the autoencoder and transform the normal data to their low dimensional latent representations.
- Use one of the previous approaches to anomaly detection to detect anomalies in the latent representations.
 - The assumption is that the latent representation of an anomaly will substantially differ from the latent representations of normal instances.

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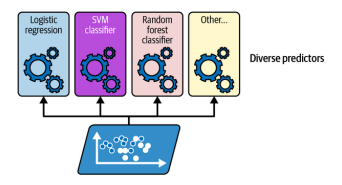
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- ► There are many more methods: One class SVM, isolation forests, etc.

Ensemble Methods

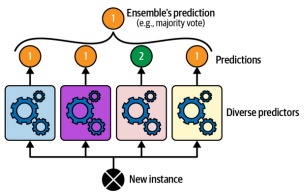
Voting Classifiers



Train several models. They may differ in

- Structure (completely different models)
- ► Training data (subsample the training set)

Voting Classifiers



During the inference, ensemble the predictions. For binary classifiers, you may do the following:

- Take the majority vote.
- Summarize the output probabilities (e.g., by averaging)
- ▶ If logistic regression or neural networks with logistic output activation are used, summarize the outputs before the application of the last logistic sigmoid.

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The ensemble methods work best when the models are as independent as possible.

Use different learning methods or independently chosen training sets (see later slides).

Let us try to formalize the above intuition.

Consider a random function $g(x) + \varepsilon$ where $g : \mathbb{R} \to \mathbb{R}$ and ε is random noise with mean 0 and variance σ^2 .

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Consider a dataset for regression:

$$D = \{(x_1, d_1), \dots, (x_p, d_p)\}\$$

Here x_1, \ldots, x_p are fixed inputs, and each $d_k = g(x_k) + \varepsilon_k$ where $\varepsilon_1, \ldots, \varepsilon_p$ are independent samples from the noise ε .

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Let us denote by $h_D: \mathbb{R} \to \mathbb{R}$ a model trained on the dataset D by minimizing the squared error $\sum_{k=1}^{p} (h_D(x_k) - d_k)^2$.

Let us consider the expected value of the squared error:

$$\mathbb{E}\left(\sum_{k=1}^{p} (h_D(x_k) - d_k)^2\right) = \sum_{k=1}^{p} \mathbb{E}(h_D(x_k) - d_k)^2$$

The expectation is computed over the distribution of the datasets D.

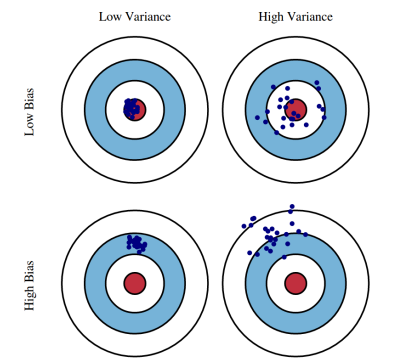
One can prove that

$$\sum_{k=1}^{p} \mathbb{E} (h_D(x_k) - d_k)^2 =$$

$$\sum_{k=1}^{p} \mathbb{E} (h_D(x_k) - \mathbb{E}[h_D(x_k)])^2 + (\mathbb{E}[h_D(x_k)] - g(x_k))^2 + \sigma^2$$

Here

- ▶ $\mathbb{E}(h_D(x_k) \mathbb{E}[h_D(x_k)])^2$ is the *Variance* of the model on x_k how much the model's value jumps around its average on x_k .
- ▶ $\mathbb{E}[h_D(x_k)] g(x_k)$ is the *Bias* of the model. how much the average model's value differs from the true values.
- $ightharpoonup \sigma^2$ is the noise variance.



B/V Decomposition Proof (Optional)

Let us study $\mathbb{E}(h_D(x_k) - d_k)^2$. To simplify notation we drop the index k and write $\mathbb{E}_D(h_D(x) - d)^2$.

$$(h_D(x) - d)^2 = (h_D(x) - \mathbb{E}[h_D(x)] + \mathbb{E}[h_D(x)] - d)^2$$

$$= (h_D(x) - \mathbb{E}[h_D(x)])^2$$

$$+ (\mathbb{E}[h_D(x)] - d)^2$$

$$+ 2(h_D(x) - \mathbb{E}[h_D(x)]) (\mathbb{E}[h_D(x)] - d)$$

Now, just apply ${\mathbb E}$ to both sides. As

$$\mathbb{E}(d) = \mathbb{E}(g(x) + \varepsilon) = g(x)$$

$$\mathbb{E}(d^2) = \mathbb{E}(g(x) + \varepsilon)^2 = \mathbb{E}(g(x)^2 + 2\varepsilon g(x) + \varepsilon^2) = g(x)^2 + \sigma^2$$

We obtain ...

B/V Decomposition Proof (Optional)

... this

$$\mathbb{E} (\mathbb{E}[h_D(x)] - d)^2$$

$$= \mathbb{E} (\mathbb{E}[h_D(x)]^2 - 2d\mathbb{E}[h_D(x)] + d^2)$$

$$= \mathbb{E}[h_D(x)]^2 - 2\mathbb{E}(d)\mathbb{E}[h_D(x)] + \mathbb{E}d^2$$

$$= \mathbb{E}[h_D(x)]^2 - 2g(x)\mathbb{E}[h_D(x)] + g(x)^2 + \sigma^2$$

$$= \mathbb{E} (\mathbb{E}[h_D(x)] - g(x))^2 + \sigma^2$$

and this

$$\mathbb{E} [2 (h_{D}(x) - \mathbb{E}[h_{D}(x)]) (\mathbb{E}[h_{D}(x)] - d)]$$

$$= \mathbb{E} [2 (h_{D}(x) - \mathbb{E}[h_{D}(x)]) (\mathbb{E}[h_{D}(x)] - g(x) - \varepsilon)]$$

$$= \mathbb{E} [2 (h_{D}(x) - \mathbb{E}[h_{D}(x)]) (\mathbb{E}[h_{D}(x)] - g(x)) - 2 (h_{D}(x) - \mathbb{E}[h_{D}(x)]) \varepsilon]$$

$$= \mathbb{E} [2 (h_{D}(x) - \mathbb{E}[h_{D}(x)]) (\mathbb{E}[h_{D}(x)] - g(x))]$$

$$= (\mathbb{E}[h_{D}(x)] - g(x)) \mathbb{E} [2 (h_{D}(x) - \mathbb{E}[h_{D}(x)])]$$

$$= 0$$

Here, the third equality follows from the zero mean of ε .

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- ▶ The Bias does not change $(\mathbb{E}[h_{D_{\ell}}(x_k)])$ is independent of ℓ):

$$\mathbb{E}\left[\frac{1}{m}\sum_{\ell=1}^{m}h_{D_{\ell}}(x_k)\right]-g(x_k)=\mathbb{E}[h_{D_{\ell}}(x_k)]-g(x_k)$$

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ight] - \mathsf{g}(\mathsf{x}_k) = \mathbb{E}[h_{D_\ell}(\mathsf{x}_k)] - \mathsf{g}(\mathsf{x}_k)$$

▶ Only the Variance changes, it is *reduced*:

$$\mathbb{E}\left(h_{ens}(x_k) - \mathbb{E}[h_{ens}(x_k)]\right)^2 = \frac{1}{m}\mathbb{E}(h_{D_\ell}(x_k) - \mathbb{E}[h_{D_\ell}(x_k)])^2$$

The equality follows from basic facts about the variance of sums of independent variables.

Now, having m independently sampled datasets is a real luxury!

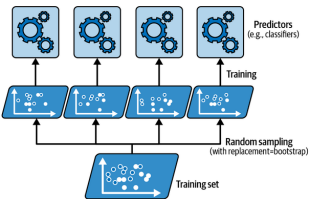
Bagging

In practice, we use sampled subsets of a given dataset.

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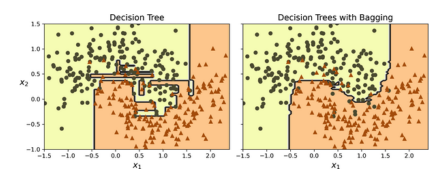
In practice, we use sampled subsets of a given dataset.

One example of using different subsets of the training set is bagging (Bootstrap Aggregating).



- ▶ Bootstrap sampling = sampling data subsets with replacement
- Aggregating = majority voting (classification) or averaging (regression)

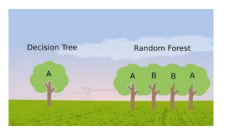
Bagging Decision Trees



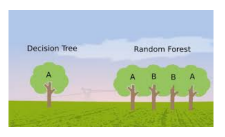
Left: A single decision tree (overfit) Right: A bagging ensemble of 500 trees

Summary of Bagging

- Reduces overfitting (variance)
- Can work with any type of classifier
- Easy to parallelize
 Just train each model independently, inference can also be parallelized.
- ► Loses (some) interpretability even for interpretable models (how could you read 500 decision trees?)

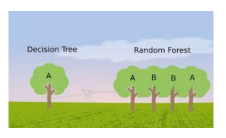


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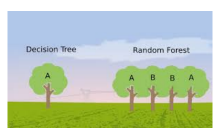


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When searching for a split attribute, look only into a randomly sampled subset of attributes (by default \sqrt{n} of n attributes).



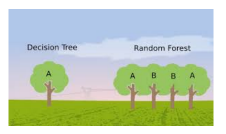
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The notion of the feature importance can be easily generalized to random forests (computed over all trees).

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To implement this kind of learning algorithm, we need to be able to train models on weighted datasets.

We are going to use weights on samples.

Note that these are different from the weights used in neural-like models!

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Linear regression: Assume a given dataset $D = \{(\vec{x}_1, f_1), \dots, (\vec{x}_p, f_p)\}$ and weights a_1, \dots, a_p associated to examples from D.

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Let \vec{w} be a vector of model weights. We minimize the weighted mean squared error

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$$p_c = \sum \{a_i \mid i\text{-th sample belongs to } c\} / \sum_{i=1}^p a_i$$

Everything else (Gini impurity, etc.) is the same.

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$$h_{ens}^k(x) = \operatorname{sign}\left(\sum_{\ell=1}^k \alpha_\ell h_\ell(x)\right)$$

Shift the weights so that the "most wrong" training instances for the current ensemble have the largest weights.

Consider a training example (x,c) with $c\in\{-1,1\}$. Then h_{ens}^k misclassifies x iff $c\cdot\sum_{\ell=1}^k\alpha_\ell h_\ell(x)<0$. The more negative this number is, the more wrong the ensemble classifier is.

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Consider a training example (x,c) with $c\in\{-1,1\}$. Then h_{ens}^k misclassifies x iff $c\cdot\sum_{\ell=1}^k\alpha_\ell h_\ell(x)<0$. The more negative this number is, the more wrong the ensemble classifier is.

After K iterations, the ensemble classifier h_{ens}^{K} is the output.

- Train classifiers using weighted samples in the training dataset.
- Start with uniform weights, that is, each sample in the dataset has the same weight.
- ► In *k*-th iteration:
 - ightharpoonup train a new classifier h_k on weighted samples,
 - obtain a coefficient $\alpha_k > 0$ of the classifier h_k ,
 - Consider the current ensemble classifier

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For details, see more advanced courses in Machine Learning.

Summary of Ensemble Methods

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- ► Technically, the ensembling decreases the model variance.
- ► There are several approaches to ensembling:
 - ► Bagging training of several models on bootstrap subsamples of the training dataset (random forest is an example)
 - ▶ Boosting ensemble is built sequentially, the newly added model possibly solves the hardest instances
 - There are many more algorithms (gradient boosting, etc.)

THE END