Machine learning for graphs and with graphs Course 1: introduction

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CR09: Machine learning for graphs and with graphs

From theory ...

- 1. Basics of machine learning
- 2. The graph framework
- 3. Community detection/ graph clustering
- 4. Graph signal processing
- 5. Kernels for graphs
- 6. Graph neural networks
- 7. Optimal transport for graphs
- 8. Learning graphs from (unstructured) data

Full description:

https://tvayer.github.io/courses/ coursegraph.html

... to practice We will use Python and various librairies



Some references for machine learning Shai Shalev-Shwartz and Shai Ben-David (2014). Understanding Machine Learning - From Theory to Algorithms.

Francis Bach (2022). Learning Theory from First Principles.

 Trevor Hastie, Robert Tibshirani, and Jerome Friedman

 (2001). The Elements of Statistical Learning.

- ▶ 50 % oral presentation on a selected research article.
- ▶ 50 % code associated to the article applied on real data.

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Bonus.

Python installations

- The practical sessions of the course will require to run jupyter notebooks.
- We recommend that you install python through the Anaconda distribution (python 3.7, 3.8 or 3.9 is preferrable) available at https://www.anaconda.com/products/distribution

You should check that you are able to create and open a jupyter notebook, and inside, run the following imports:

```
1 import matplotlib
```

- 2 import numpy
- 3 import sklearn
- 4 import pytorch
- 5 import pandas
- 6 import scipy

If any of these packages is missing, it can be installed with 'conda install numpy', the command being run in a terminal or in Anaconda prompt for Windows user.

Basics of machine learning

What is machine learning ?

Data in machine learning

From training data to prediction Loss functions Empirical risk minimization

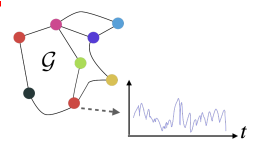
Model selection and validation Split your dataset !

The problems with structured data

Motivating examples A primer on graph theory Why "classical ML" struggles with stuctured data

Some applications

1. Energy networks, disease propagation

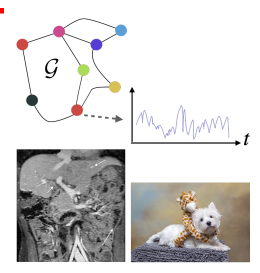


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- 1. Energy networks, disease propagation
- 2. Image analysis (medical application, web)

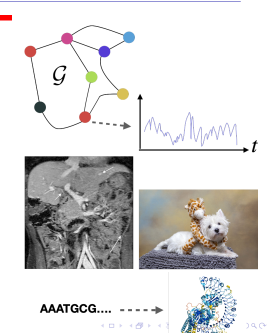


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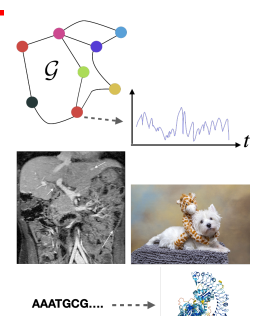
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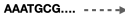
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- 4. Generative models https: //stablediffusionweb.com/



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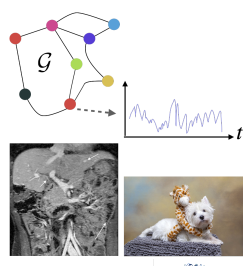
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- For art https://www.youtube. com/watch?v=MwtVkPKx3RA



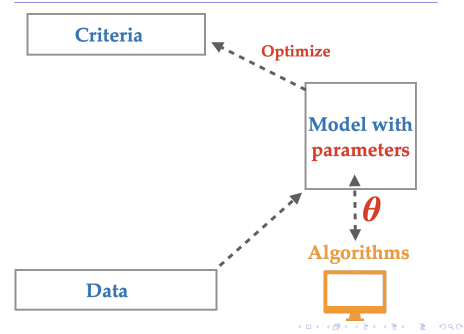




Criteria

Data

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Teach a machine to process automatically a some data in order to solve a given problem.

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Teach a machine to process automatically a some data in order to solve a given problem.

Unsupervised learning: understanding the data

- Clustering & probability density estimation
- Dimensionality reduction



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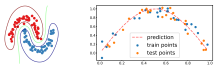
Unsupervised learning: understanding the data

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Supervised learning: learning to predict

- Classification: classify points according to some labels
- Regression: predict real (vector) values

Some images and slides have been obtained by the courtesy of Rémi Flamary



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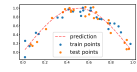
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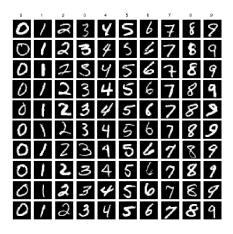
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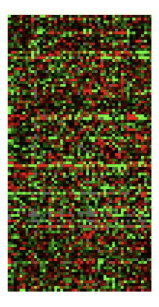






Supervised classification examples

- e.g. to identify the numbers on images from a 16 × 16 gray level image (image classification)
- ► SPAM, fraud detection, disease classification ...



Clustering example

- Analyse n sequences (individuals) of d genetical responses
- Groups of similar samples ? Gene with similar expressions ?

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Plan

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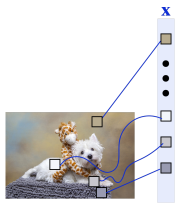
Store a data point

Vectorial representation

One "sample", "data point", "individual":

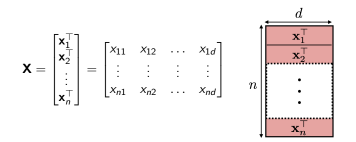
$$\mathbf{x} = (x_1, \cdots, x_d)^{ op} \in \mathbb{R}^d$$

- d is the dimension, x_i is the *i*th information *i* of
 x
- Can describe information about an individual
- For an image x: each pixel of an image
- Descriptors of a cell, word embedding ...



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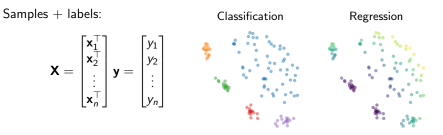
Unsupervised dataset



Unsupervised learning

- The dataset contains the samples (x_i)ⁿ_{i=1} where n is the number of samples of size d.
- d and n define the dimensionality of the learning problem.
- Data stored as a matrix X ∈ ℝ^{n×d} that contains the training samples as rows.

Supervised dataset

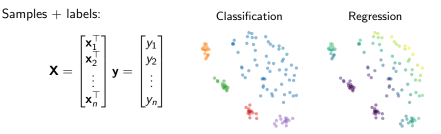


Supervised learning

- ► The dataset contains the samples (x_i, y_i)ⁿ_{i=1} where x_i is the feature sample and y_i ∈ 𝔅 its label.
- ▶ The values to predict (label) can be concatenated in a vector $\mathbf{y} \in \mathcal{Y}^n$

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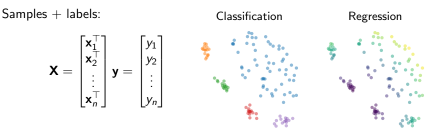
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- Prediction space *Y* can be:
 - $\mathcal{Y} = \{-1, 1\}$ or $\mathcal{Y} = \{1, \dots, K\}$ for classification problems.
 - $\mathcal{Y} = \mathbb{R}$ for regression problems (\mathbb{R}^{p} for multi-output regression).

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 - $\mathcal{Y} = \mathbb{R}$ for regression problems (\mathbb{R}^{p} for multi-output regression).
- Semi-supervised learning: few labeled points are available, but a large number of unlabeled points are given.

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Training data

• We have access to *n* samples $(\mathbf{x}_1, y_1), \cdots, (\mathbf{x}_n, y_n) \sim p$

- $p \in \mathcal{P}(\mathcal{X} \times \mathcal{Y})$ is the data distribution
- p is unknown ! We only have access to samples.

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- ▶ For unsupervised problem we only have $\mathbf{x}_1, \cdots, \mathbf{x}_n \sim p$ and $p \in \mathcal{P}(\mathcal{X})$

Objective

- ▶ We have a task to solve: classification, regression, clustering ...
- Most ML problems formulate as finding some function f that "best" solves our task
- f is called an hypothesis and is implemented by a computer
- Most of the time f depends on some parameter $\theta \in \Theta$

Regression



Objective

$$(\mathbf{x}_i, y_i)_{i=1}^n \quad \Rightarrow \quad f: \mathbb{R}^d \to \mathbb{R}$$

▶ Train a function $f(\mathbf{x}) = y \in \mathcal{Y}$ predicting a continuous value $(\mathcal{Y} = \mathbb{R})$.

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• Can be extended to multi-value prediction $(\mathcal{Y} = \mathbb{R}^p)$.

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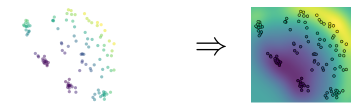
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Hyperparameters

- Type of function (linear, kernel, neural network).
- Performance measure.
- Regularization.

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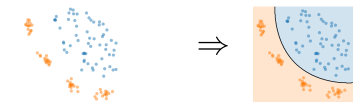
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Methods

- ► Least Square (LS).
- ► Ridge regression, Lasso.
- Kernel regression.
- ► Deep learning.

Binary classification



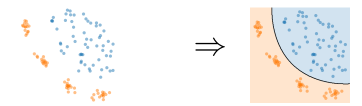
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$$(\mathbf{x}_i, y_i)_{i=1}^n \quad \Rightarrow \quad f: \mathbb{R}^d \to \{-1, 1\}$$

Train a function f(x) = y ∈ 𝔅 predicting a binary value (𝔅 = {−1,1}).
f(x) = 0 defines the boundary on the partition of the feature space.

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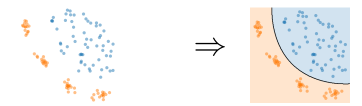
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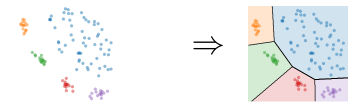
Hyperparameters

- Type of function (linear, kernel, neural network).
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Methods

- Bayesian classifier (LDA, QDA)
- Linear and kernel discrimination
- Decision trees, random forests.
- ► Deep learning.

Multiclass classification



Objective

$$(\mathbf{x}_i, y_i)_{i=1}^n \quad \Rightarrow \quad f: \mathbb{R}^d \to \{1, \dots, K\}$$

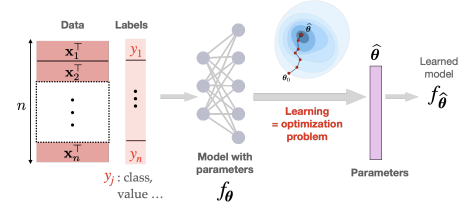
- ► Train a function f(x) = y ∈ 𝔅 predicting an integer value (𝔅 = {1,..., 𝔅}).
- In practice K continuous score functions f_k are estimated and the prediction is

$$f(\mathbf{x}) = \arg \max_{k} f_k(\mathbf{x})$$

Softmax can be used instead of argmax to get probability estimates.

The big picture of (parametrized) ML

But how to find this function ?



The goal in the learning step will be to find the parameters $\hat{\theta}$ (hence the function) that minimizes a measure of error on the data

A loss function is $\ell:\mathcal{Y}\times\mathcal{Y}\rightarrow\mathbb{R}$ so that:

 ℓ (true value , predicted value) = how good is my prediction



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$$y_i \in \mathbb{R} \ \ell(y_i, f(\mathbf{x}_i)) = (y_i - f(\mathbf{x}_i))^2$$
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Classification problems

► E.g.
$$y_i \in \{-1, 1\} \ \ell(y_i, f(\mathbf{x}_i)) = \mathbf{1}_{y_i \neq f(\mathbf{x}_i)} \ (0/1 \text{ loss})$$

Loss functions

A focus on classification problems $\mathcal{Y} = \{-1,1\}$

 $\ell(y_i, f(\mathbf{x}_i)) = \Phi(y_i f(\mathbf{x}_i))$ with Φ non-increasing.

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• $y_i f(\mathbf{x}_i)$ is **the margin** (on the board).

$$\blacktriangleright \ \ell(y_i, f(\mathbf{x}_i)) = \mathbf{1}_{y_i f(\mathbf{x}_i) \leq 0} \ (0/1 \text{ loss})$$

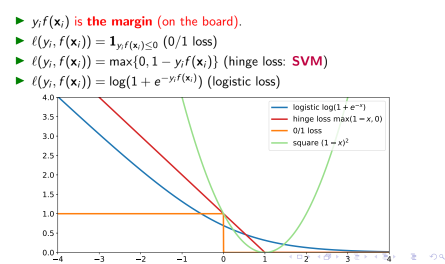
• $\ell(y_i, f(\mathbf{x}_i)) = \max\{0, 1 - y_i f(\mathbf{x}_i)\}$ (hinge loss: **SVM**)

•
$$\ell(y_i, f(\mathbf{x}_i)) = \log(1 + e^{-y_i f(\mathbf{x}_i)})$$
 (logistic loss)

Loss functions

A focus on classification problems $\mathcal{Y} = \{-1, 1\}$

 $\ell(y_i, f(\mathbf{x}_i)) = \Phi(y_i f(\mathbf{x}_i))$ with Φ non-increasing.



To find f the idea is to **minimize the averaged error** on the training samples:

$$\min_{f} \frac{1}{n} \sum_{i=1}^{n} \ell(y_i, f(\mathbf{x}_i))$$
(ERM)

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- It is called empirical risk minimization (ERM)
- Given the loss, finds the "best" f on the training data
- Same idea applies for unsupervised problem (minimizing reconstruction error)

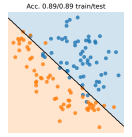
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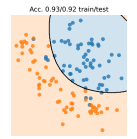
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Once solved how do I know if my model is good ?

Underfitting and overfitting

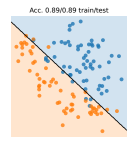


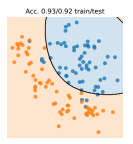


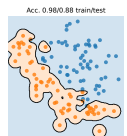
Acc. 0.98/0.88 train/test

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Underfitting and overfitting

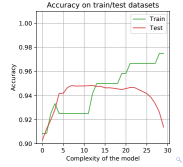






Complexity of a model

- Under-fitting when the model is too simple.
- Over-fitting occurs when the model is too complex
- Training data performance is not a good proxy for testing performance.
- We want to predict well on new data!
- Parameter and model validation.



To find f the idea is to **minimize the averaged error** on the training samples:

$$\min_{f} \frac{1}{n} \sum_{i=1}^{n} \ell(y_i, f(\mathbf{x}_i)) + \lambda \operatorname{Reg}(f)$$
(ERM)

- It is called empirical risk minimization (ERM)
- Given the loss, finds the "best" f on the training data
- Teacher/student analogy
- Same idea applies for unsupervised problem

... but we want generalization !

- We want f to be good outside the training samples
- Add regularization ! (limit the complexity of f)

Plan

What is machine learning ?

Data in machine learning

From training data to prediction Loss functions Empirical risk minimization

Model selection and validation Split your dataset !

The problems with structured data

Motivating examples A primer on graph theory Why "classical ML" struggles with stuctured data

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Bias-complexity tradeoff

generalization error = estimation error + approximation error

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Select a model that is not too complex but not too simple !

Bias-complexity tradeoff

generalization error = estimation error + approximation error

Select a model that is not too complex but not too simple !

General principle

Estimate the generalization error on data not seen during training

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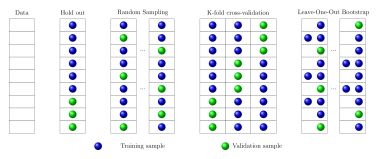
- Is a rough estimate of the test error
- Select the model with the lowest "approximate" test error



Principle of Hold-Out cross-validation

- Split the training data in a training and validation sets (non overlapping).
- Train different models (different methods and/or hyperparameters) on the train data.
- Evaluate performance on the validation data and select the method/parameters with best performance.
- Validation set acts as a proxy of test data
- But only one split is a poor proxy !

Different ways to split the data



Data splitting for cross-validation Arlot and Celisse 2010

- The training data is split in non-overlapping training/validation sets.
- Hold-Out uses a unique split.
- More robust cross-validation approaches actually investigate several splits of the data and compute the average performance:
 - K-fold (split in K sets and use one split as test for all k)
 - Random sampling (aka Shuffle split) draws several random splittings.
- Scikit-learn implementation : sklearn.model_selection.cross_validate

Plan

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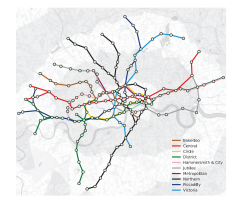
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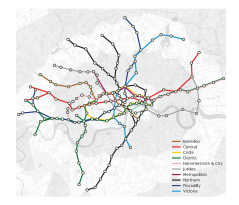
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 Traffic forecasting (e.g. ETA estimation): GNN for Google Maps Derrow-Pinion et al. 2021.



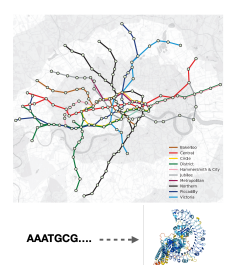
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- Traffic forecasting (e.g. ETA estimation): GNN for Google Maps Derrow-Pinion et al. 2021.
- Chemistry and Drug Design: space of chemically synthesisable molecules is very large (estimated around 10⁶⁰).
- ► Drug Repositioning: action of drugs and their interactions → graph Barabási, Gulbahce, and Loscalzo 2011.



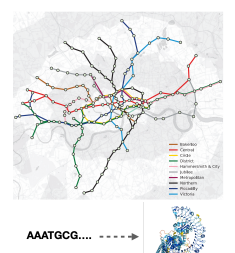
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- Healthcare.



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- and more...



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Definition

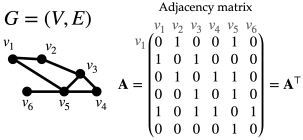
A graph G = (V, E) is defined as a set of **vertices** V, which are connected by a set of **edges** $E \subset V \times V$.

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Definition

A graph G = (V, E) is defined as a set of **vertices** V, which are connected by a set of **edges** $E \subset V \times V$.

Example of undirected graph



Adjacency matrix

The adjacency $\boldsymbol{\mathsf{A}} \in \mathbb{R}^{|\mathcal{V}| \times |\mathcal{V}|}$ is defined as

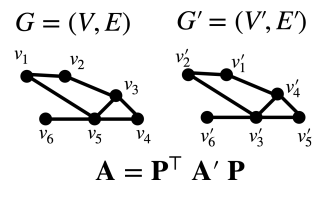
$$[\mathbf{A}]_{ij} = \begin{cases} 1 \text{ if } (v_i, v_j) \in E(\text{often noted as } v_i \sim v_j) \\ 0 \text{ otherwise} \end{cases}$$

Definition

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Isomorphic graphs

The definition depends on the ordering of the nodes.



Definition

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Some special structures





Complete graph



Bipartite graph



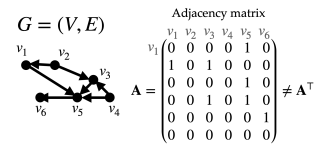
Star graph



Circular graph

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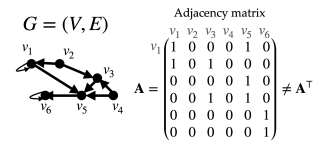
Example of **directed** graph



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Example of directed graph with self-loops.



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Weighted graph

A weighted graph G = (V, E) associates <u>non-negative</u> weights to each edge.

Example of weighted graph

$$G = (V, E)$$
Weight matrix
$$\begin{array}{c} & & & & \\ V_1 & 0.5 & V_2 \\ \bullet & & \\ & & & & \\ & & & & \\ & & & \\ & & & & \\ & & & & \\ & & & \\ & & & & \\ & & & & \\ & &$$

Degree of a node

The degree of a node v_i is

$$d_i = |\{v \in V : v \sim v_i\}| = \sum_{j=1}^{|V|} A_{ij}$$

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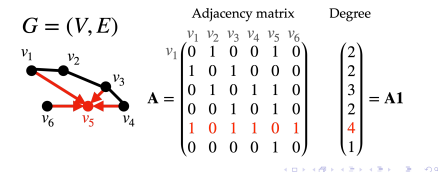
The degree matrix is $\mathbf{D} = \text{diag}(d_1, \cdots, d_{|V|})$.

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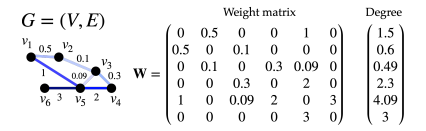
The degree matrix is $\mathbf{D} = \text{diag}(d_1, \cdots, d_{|V|})$.



Degree of a node

The degree of a node v_i in a weighted graph is

$$d_i = \sum_{j=1}^{|V|} W_{ij}$$



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Laplacian matrix

The Laplacian matrix of a undirected graph is defined as

 $\mathbf{L}=\mathbf{D}-\mathbf{W}$ where \mathbf{D} is the degree matrix

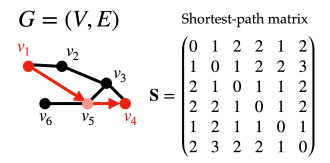
Properties

On the board

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Shortest-path matrix

The shortest-path between $v, v' \in V$ is the path that connects v, v' such that the sum of the weights of its constituent edges is minimized.

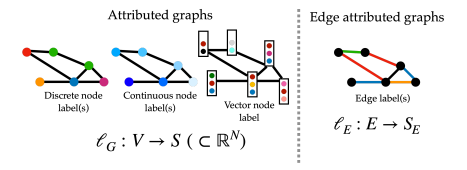


Dijkstra's algorithm computes all the shortest paths from a single node in O(|E| + |V| log(|V|)).

▶ All-pairs shortest paths with Floyd–Warshall algorithm in $O(|V|^3)$.

Attributed graphs

Most graphs encountered in ML also have attributes.



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Problems

• Can we encode a graph G as a vector $\in \mathbb{R}^d$ to use standard ML methods ?



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How can we handle the combinatoric nature of graphs ?

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- ML outputs should be permutation invariant ? equivariant ?

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- ML outputs should be permutation invariant ? equivariant ?
- When data = vectors in one graph how can we take into account the structure of the graph ?
- When data = vectors can we find an (interesting) graph that represent these data ?

References I

Arlot, Sylvain and Alain Celisse (2010). "A survey of cross-validation procedures for model selection". In: Statistics Surveys 4. ā. Bach, Francis (2022). Learning Theory from First Principles. Barabási, Albert-László, Natali Gulbahce, and Joseph Loscalzo (2011). "Network medicine: a network-based approach to human disease". In: Nature reviews genetics 12.1, pp. 56-68. Derrow-Pinion, Austin et al. (2021). "Eta prediction with graph neural networks in google maps". In: Proceedings of the 30th ACM International Conference on Information & Knowledge Management, pp. 3767–3776. Hastie, Trevor, Robert Tibshirani, and Jerome Friedman (2001). The Elements of Statistical Learning. Jumper, John et al. (2021). "Highly accurate protein structure prediction with AlphaFold". In: *Nature* 596.7873, pp. 583–589. Shalev-Shwartz, Shai and Shai Ben-David (2014). Understanding Machine Learning - From Theory to Algorithms.