

# Homework 2: Similarity measures on time series and graphs

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## Objectives

The goals of this homework are:

- to learn how to model data as time series and graphs.
- to implement similarity measures on time series and graphs.
- to apply similarity measures on a heartbeat time series dataset and a molecular graphs dataset.
- to explore theoretical aspects of these dissimilarity or similarity measures.

## Problem overview

In this homework you will implement the Dynamic Time Warping (DTW) distance function [5] on *time series* in **Part I** and the Shortest Path Kernel (SPKernel) similarity measure [1] on *graphs* in **Part II**. Additionally, you will be asked to address some theoretical questions about DTW and SPKernel.

## Homework Part 1: time series

### Introduction

A time series is a sequence of measurements over a time interval. There are many methods to compute distances between time series. Dynamic Time Warping (DTW) is one of the most famous and widely-used methods. In this homework, you will implement the DTW distance and use it on a heartbeat time series dataset.

Suppose we have two time series  $t_1$  and  $t_2$  of lengths  $m$  and  $n$ , respectively. We define an  $(m, n)$ -warping path to be a sequence  $(w_l)_{l=1}^L$  of index pairs such that each term  $w_l = (w_l^{(1)}, w_l^{(2)}) \in \llbracket 1, m \rrbracket \times \llbracket 1, n \rrbracket$  indexes an element  $t_1[w_l^{(1)}]$  of time series  $t_1$  and an element  $t_2[w_l^{(2)}]$  of time series  $t_2$  for all  $l = 1, \dots, L$ . The length  $L$  of the warping path satisfies  $\max(m, n) \leq L < m + n - 1$ . An  $(m, n)$ -warping path can be understood as a sequence of correspondences between elements of  $t_1$  and  $t_2$ , thus defining an alignment between the two time series. In DTW, the set of allowable  $(m, n)$ -warping paths is restricted to those which satisfy the following three properties:

**Boundary conditions:** The first and last terms of the warping path must be  $w_1 = (1, 1)$  and  $w_L = (m, n)$ , respectively. That is, the warping path must start comparing the first element of  $t_1$  with the first element of  $t_2$  and end comparing the last element of  $t_1$  with the last element of  $t_2$ .

**Monotonicity:** Let  $w_l = (i, j)$  and  $w_{l+1} = (i', j')$  be two consecutive elements in the warping path. Then,  $i \leq i'$  and  $j \leq j'$  must hold. That is, the warping path cannot go “backwards”.

**Unit-size steps:** Let  $w_l = (i, j)$  and  $w_{l+1} = (i', j')$  be two consecutive elements in the warping path. Then,  $i' - i \leq 1$  and  $j' - j \leq 1$  must hold, with at least one of them being a strict inequality. That is, if  $w_l = (i, j)$ , the next element in the warping path  $w_{l+1}$  must be one of  $(i + 1, j)$ ,  $(i, j + 1)$  or  $(i + 1, j + 1)$ .

If we denote by  $\mathcal{W}(m, n)$  the set of all  $(m, n)$ -warping paths satisfying all three conditions above, the DTW distance between two time series  $t_1$  and  $t_2$  of lengths  $m$  and  $n$  is defined by the following optimization problem:

$$\text{DTW}_d(t_1, t_2) = \min_{w \in \mathcal{W}(m, n)} \sum_{l=1}^{L(w)} d\left(t_1[w_l^{(1)}], t_2[w_l^{(2)}]\right), \quad (1)$$

where  $d : \mathbb{R} \times \mathbb{R} \rightarrow \mathbb{R}_+$  is an element-wise, user-defined distance function.

$\text{DTW}_d(t_1, t_2)$  can be efficiently calculated using dynamic programming. Define  $C$  to be a  $(m + 1) \times (n + 1)$  matrix such that  $C_{i+1, j+1}$  equals the DTW distance between the time series  $t_1[1 : i]$  containing the first  $i$  elements of time series  $t_1$  and the time series  $t_2[1 : j]$  containing the first  $j$  elements of time series  $t_2$ . Each entry  $C_{i, j}$  can be obtained in  $O(1)$  time provided the values of  $C_{i-1, j-1}$ ,  $C_{i, j-1}$  and  $C_{i-1, j}$  are known:

$$C_{i, j} = d(t_1[i], t_2[j]) + \min(C_{i-1, j-1}, C_{i, j-1}, C_{i-1, j}). \quad (2)$$

With  $C$  defined in such a way, it is evident that  $\text{DTW}_d(t_1, t_2) = C_{m+1, n+1}$ . Note that if  $i = 1$ , then  $C_{i-1, j-1}$  and  $C_{i-1, j}$  are undefined and  $C_{i, j} = d(t_1[i], t_2[j]) + C_{i, j-1}$ . Analogously, if  $j = 1$ , then  $C_{i, j} = d(t_1[i], t_2[j]) + C_{i-1, j}$ . In order to avoid explicitly dealing with these particular cases, we define the sentinel values

$$\begin{aligned} C_{0,0} &= 0, \\ C_{i,0} &= \infty \quad \forall i = 1, \dots, m, \\ C_{0,j} &= \infty \quad \forall j = 1, \dots, n, \end{aligned} \quad (3)$$

which render Equation (2) valid for all  $i = 1, \dots, m$  and  $j = 1, \dots, n$ .

## Dataset

You will work on the “ECG” heartbeat time series data from the UCR time series archive [2], that has been widely used as one of the benchmark datasets for evaluating time series classification. Each record of “ECG” is a time series of measurements recorded by one electrode during one heartbeat. The data has been annotated by a cardiologist and a label of normal or abnormal is assigned to each data record.

In Exercise 1, you will work on the dataset stored in `ECG200_TRAIN.txt`. There are 100 time series samples and the length of each sample is 96 (same length for all samples). The first column indicates the class of samples, where  $-1$  means the abnormal heartbeat and  $1$  the normal heartbeat. The other 96 columns are records of time series samples.

## Exercise 1

**Exercise 1.a** A common variant of DTW adds an additional requirement in the definition of valid  $(m, n)$ -warping paths, namely:

**$w$ -constrained warping:** Any element  $w_l = (i, j)$  in the warping path must satisfy  $|i - j| \leq w$ , where  $w \geq |m - n|$  is a user-defined hyperparameter.

Create a Python module named `dynamic_time_warping.py` in which you will implement the DTW distance function with  $w$ -constrained warping:

```
constrained_dtw(t1, t2, w)
```

**Please note:**

- The first two arguments of the `constrained_dtw` function, namely `t1` and `t2`, are both lists of type `float`, while the third argument `w` is a non-negative integer not smaller than the difference in length between the two time series `t1` and `t2` to be compared.
- The name of the function and its arguments should match the above description.
- The return value is a `float`.
- Use the Manhattan distance  $d(x, y) = |x - y|$  as the base element-wise distance for DTW, that is,  $d(t_1[i], t_2[j]) = |t_1[i] - t_2[j]|$ .
- Do not use libraries or external source code that implement the DTW distance function, with or without  $w$ -constrained warping.

**Exercise 1.b** Since the lengths of different time series samples in the “ECG” dataset are the same, they can also be treated as vectors. In such a case, distance functions on vectors can also be used to compare time series. In this exercise, you will exploit this to compare DTW with the **Manhattan** distance on vectors.

Create a Python script named `compute_dtw.py` to read the `ECG200_TRAIN.txt` dataset and compute the average distances between groups using:

- i) The **Manhattan** distance, treating the time series as vectors.

- ii) The DTW distance with  $w$ -constrained warping and the Manhattan distance as element-wise metric, for values of  $w$  in  $\{0, 10, 25, \infty\}$ .

**Please note:**

- Your program `compute_dtw.py` will receive 2 command line arguments:
  - `--datadir path`: the path to the directory where the input file is stored.
  - `--outdir path`: the path to the directory where the output file will be stored.
- The program will compute the average DTW and `Manhattan` distances between groups and output them to the file `timeseries_output.txt`. Its format is shown in Figure 1. Your output file should include the header as well and output rows and columns in the specified order.

Pair of classes	Manhattan	DTW, $w = 0$	DTW, $w = 10$	DTW, $w = 25$	DTW, $w = \infty$
abnormal:abnormal	99.9	99.9	99.9	99.9	99.9
abnormal:normal	99.9	99.9	99.9	99.9	99.9
normal:normal	99.9	99.9	99.9	99.9	99.9

Figure 1: Format of output file `timeseries_output.txt`.

**Exercise 1.c** Compare and discuss the results of the DTW and `Manhattan` distances on separating abnormal and normal heartbeats.

**Exercise 1.d** Discuss the effect that hyperparameter  $w$  has on the DTW distance and its ability to separate abnormal and normal heartbeats.

**Exercise 1.e** Is the DTW distance a metric? If not, give examples showing which conditions are not satisfied.

**Exercise 1.f** What is the runtime complexity of computing the DTW distance with  $w$ -constrained warping? You might consider that  $m = n$ .

## Homework Part 2: Graphs

### Introduction

Graphs provide a flexible way to represent complex data, such as molecules. As a result, they play a central role in data mining. Firstly, we consider the similarity between nodes in the graph. The most popular distance function is the length of the shortest path between nodes. Secondly, we compute the similarity between graphs. Graph kernels have been proposed in order to do this job. The Shortest Path kernel (`SPKernel`) is one that uses shortest paths to compare graphs.

Suppose we have two graphs `G1` with `m` vertices and `G2` with `n` vertices (for simplicity, in this homework, we only consider unweighted and undirected graphs). The corresponding adjacency matrices are `A1` and `A2`. Then the shortest path kernel (`SPKernel`) between `G1` and `G2` can be calculated by the following steps:

- Transform the adjacency matrix **A1** and **A2** into shortest path matrix **S1** and **S2**, where  $S(i, j)$  is the length of shortest path between node  $i$  and  $j$ .
- The **SPKernel** is calculated with **S1** and **S2**:

$$K_{sp}(S1, S2) = \sum_{e_1 \in S1} \sum_{e_2 \in S2} K_{walk}^1(e_1, e_2)$$

$$K_{walk}^1(e_1, e_2) = \begin{cases} 1, & \text{if } weight(e_1) == weight(e_2) \\ 0, & \text{otherwise} \end{cases}$$

- $e_1$  is an edge walk of length 1 in **S1**, which is a non-zero entry of **S1**. Since **S1** is symmetric, only the upper or lower triangular matrix of **S1** should be considered.

## Dataset

You will work on the “MUTAG” dataset of 188 mutagenic aromatic and heteroaromatic nitro compounds [3]. As stated in the dataset description page:

*“...Sample graphs are labeled according to whether or not they have a mutagenic effect on the Gram-negative bacterium Salmonella typhimurium.”*<sup>1</sup>

Graphs with a mutagenic effect are labeled 1, otherwise -1. Each sample graph is represented by its adjacency matrix. The adjacency matrices and labels are stored in **MUTAG.mat**. This dataset will be used in Exercise 2. Instructions for how to load this file in Python will be introduced later in Exercise 2.b.

## Exercise 2

**Exercise 2.a** Create a Python script named `shortest_path_kernel.py` in which you will implement (i) Floyd-Warshall’s algorithm [4] to compute shortest path lengths and (ii) **SPKernel**. The pseudo-code of Floyd-Warshall’s algorithm can be found in the lecture slides *Part1: Introduction and Distance Functions*.

```
floyd_warshall(A)
spkernel(S1, S2)
```

Please note:

- The argument of the function `floyd_warshall` is the adjacency matrix **A** of a graph. If there is no link between two nodes  $i$  and  $j$ , then  $A(i, j)$  will be 0. For such pairs of nodes  $i$  and  $j$ , the shortest distance should be initialized as infinite. Therefore, in practice, you will need to convert all the zero entries except the diagonal of **A** to a very large value before calculating the shortest path lengths.
- The return value of `floyd_warshall` is the shortest path matrix **S** of a graph.
- The parameters of the function `spkernel` are two shortest path matrices **S1** and **S2**.
- The return value of `spkernel` is a float.
- Do not use libraries or external source code that implement `spkernel` and `floyd_warshall`.

<sup>1</sup><https://www.bsse.ethz.ch/mlcb/research/machine-learning/graph-kernels.html>

**Exercise 2.b** Create a script named `compute_spkernel.py` to read the `MUTAG.mat` dataset and compute the average `SPKernel` similarities between the abnormal and normal class.

With `scipy.io.loadmat` the adjacency matrices and label information can be loaded using the following script:

```
import scipy.io
mat = scipy.io.loadmat('PATH to MUTAG.mat')
label = np.reshape(mat['lmutag'], (len(mat['lmutag']), ))
data = np.reshape(mat['MUTAG']['am'], (len(label), ))
```

Please make sure that the `scipy` module is installed before using it.

**Please note:**

- Your program `compute_spkernel.py` will receive 2 command line arguments:
  - `--datadir path`: the path to the directory where the input file is stored.
  - `--outdir path`: the path to the directory where the output file will be stored.
- The program will compute the average similarities and output them to a file named `graphs_output.txt`. Its format is shown in Figure 2.

Pair of classes	SPKernel
mutagenic:mutagenic	99.9
mutagenic:non-mutagenic	99.9
non-mutagenic:non-mutagenic	99.9

Figure 2: Format of output file `graphs_output.txt` file.

**Exercise 2.c** What is the runtime complexity of Floyd-Warshall's algorithm? What is the runtime complexity of `SPKernel`?

## Grading and submission guidelines

This homework is worth a total of 100 points. Table 1 shows the points assigned to each exercise.

**WARNING:** Submitted scripts that do not run will result in zero points for that exercise and will not be corrected.

Table 1: Grading key for Homework 2

<b>60 pts.</b>	<b>Exercise 1</b>
35 pts.	Exercise 1.a
5 pts.	Exercise 1.b
5 pts.	Exercise 1.c
5 pts.	Exercise 1.d
5 pts.	Exercise 1.e
5 pts.	Exercise 1.f
<b>40 pts.</b>	<b>Exercise 2</b>
25 pts.	Exercise 2.a
5 pts.	Exercise 2.b
10 pts.	Exercise 2.c

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## References

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