

# EE 046202 - Technion - Unsupervised Learning & Data Analysis

Tal Daniel (https://taldatech.github.io)

# Tutorial 06 - Dimensionality Reduction - t-SNE



#### Agenda

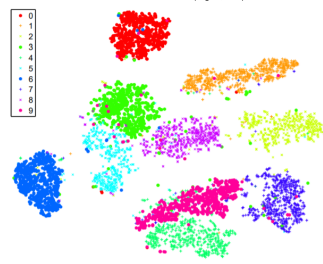
- Motivation and Introduction
- Stochastic Neighbor Embedding (SNE))
- t-Distributed Stochastic Neighbor Embedding (t-SNE))
  - Symetric SNE
  - The Crowding Problem
  - Student t-distribution
  - Algorithm and Optimization
  - Weaknesses
- Examples
- Recommended Videos
- Credits

```
In [1]:  # imports for the tutorial
import numpy as np
import pandas as pd
import matplotlib.pyplot as plt
from mpl_toolkits.mplot3d import Axes3D
from sklearn.manifold import TSNE
from sklearn.preprocessing import StandardScaler
from scipy.stats import t
%matplotlib notebook
```



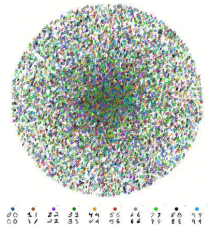
- We want to have a dimensionality reduction method that may help us understand the data, and there is no better way to "explain" data than seeing it visually!
- t-SNE is a method to visualize high-dimensional data, which is easier to optimize than many other methods.
- One of its most important traits is that it mitigates the crowding problem which we will discuss, but many other dimensionality reduction algorithms suffer from.
  - In short, it is the tendency of dimensionality reduction algorithms to crowd points together in the center of the map.

• It looks really (!) good. Here, have a look at the result on the MNIST dataset (digits, 0-9):



- The goal of dimensionality reduction is to preserve as much of the significant structure of the high-dimensional data as possible in the low-dimensional map.
  - For example, recall that PCA is a linear method that focuses on keeping the low-dimensional representations of dissimilar points far apart. It is very common to chain together PCA and t-SNE to achieve good results (first perform PCA and then apply t-SNE).
  - On the other hand, high-dimensional data that lies on or near a low-dimensional, non-linear manifold, it is usually more important to keep the low-dimensional representations of very similar datapoints close together, which is usually not possible with linear mapping.
- So what is the problem with other techniques like Sammon, SNE, Isomap, LLE, Laplacian Eigenmaps and etc...?
  - Despite performing really well on artificial datasets, they are often not very successful at visualizing real, high-dimensional data.
  - Most of them are not capable of retaining both the local and the global structure of the data in a single map (while t-SNE does well!).
- · t-SNE is based (how surprising...) on SNE which we introduce shortly.
- Formulation:
  - $\begin{array}{l} \blacksquare \mbox{ High-dimensional dataset: } \mathcal{X} = \{x_1, x_2, \ldots, x_n\} \mbox{ (total of } n \mbox{ points)} \\ \blacksquare \mbox{ Low-dimensional dataset: } \mathcal{Y} = \{y_1, y_2, \ldots, y_n\} \\ \blacksquare \mbox{ The goal is to find mapping } \mathcal{X} \to \mathcal{Y} \end{array}$

· Not convinced yet? look at this:



Source: Google Al Blog (https://ai.googleblog.com/2018/06/realtime-tsne-visualizations-with.html)



## Stochastic Neighbor Embedding (SNE)

We will now introduce the base algorithm of t-SNE, called Stochastic Neighbor Embedding (SNE), and present how it works.

- · SNE starts by converting Euclidean distances between datapoints into conditional probabilities that represent similarities.
  - ullet The **similarity** of datapoint  $x_j$  to datapoint  $x_i$  is denoted  $p_{j|i}$ , that is, the conditional probability that  $x_i$  would pick  $x_j$  as its neighbor, if neighbors were picked in proportion to their probability density under a **Gaussian** centered (its mean) at  $x_i$ .
  - As expected, for nearby datapoints this probability would be high whereas for widely separated datapoints it would be very small.
- · Mathematically:

$$p_{j|i} = rac{e^{-||x_i - x_j||^2/2\sigma_i^2}}{\sum_{k 
eq i} e^{-||x_i - x_k||^2/2\sigma_i^2}}$$

- ullet  $\sigma_i$  is the variance of the Gaussian that is centererd on datapoint  $x_i$
- ullet  $p_{i|i}=0$ , as we are interested only in the pairwise relation.
- Similarly, for the **low-dimensional** counterparts,  $y_i$  and  $y_j$  we denote the conditional probability:

$$q_{j|i} = rac{e^{-||y_i - y_j||^2}}{\sum_{k 
eq i} e^{-||y_i - y_k||^2}}$$

- $q_{j|i} = \frac{e^{-||y_i y_j||^2}}{\sum_{k \neq i} e^{-||y_i y_k||^2}}$   $\blacksquare \text{ We set the } \textit{variance} \text{ of the Gaussian to } \frac{1}{\sqrt{2}} \text{ which results in } 2 \cdot (\frac{1}{\sqrt{2}})^2 = 1 \text{ (in case you were wondering where did the } \sigma \text{ go)}.$   $\bullet \text{ Why } \frac{1}{\sqrt{2}} \text{ ? setting to another value only changes the scale in the final map. Using the } \mathbf{same} \text{ variance for each datapoint we lose}$ some property of the data (as it has  $\sigma_i$  in the high-dimensional space).
- Intuition if the mapping  $\mathcal{X} o \mathcal{Y}$  correctly models the similarity between datapoints, then  $p_{j|i}$  and  $q_{j|i}$  will be equal.

- Choosing σ<sub>i</sub>:
  - In **dense** regions a smaller value of  $\sigma_i$  is usually more appropriate than in **sparser** regions.
  - SNE performs binary search for the value of  $\sigma_i$  that produces a  $P_i$  with a fixed perplexity that is specified by the user. The perplexity is

$$Perp(P_i) = 2^{H(P_i)}$$

$$Perp(P_i)=2^{H(P_i)}$$
  $\circ~H(P_i)$  is the **Shannon entropy** of  $P_i$  measured in bits 
$$H(P_i)=-\sum_j p_{j|i}\log_2 p_{j|i}$$

- Intuition for perplexity a smooth measure of the effective number of neighbors.
- Typical values are between 5 and 50 (SNE is fairly robust for this hyper-parameter).
- The **goal** of SNE is to find a mapping that minimizes the mismatches between  $p_{j|i}$  and  $q_{j|i}$ .
  - How do we measure "distance between probabilities"? As you probably have guessed, KL-divergence comes to the rescue!
- SNE minimizes the sum of Kullback-Leibler divergences over all datapoints using a gradient descent method. We denote the cost function C:

$$C = \sum_i KL(P_i||Q_i) = \sum_i \sum_{j 
eq i} p_{j|i} \log rac{p_{j|i}}{q_{j|i}}$$

- ullet  $P_i$  the conditional probability distribution over all other *datapoints* given datapoint  $x_i$
- ullet  $Q_i$  the conditional probability distribution over all other *map points* given point  $y_i$ .
- · KL-divergence is not a metric as it is not symmetric, and many errors are due to the fact the distances are not weighted equally.
  - There is a **large cost** for using **widely separated map points** to represent nearby datapoints (using a small  $q_{i|i}$  to model a large  $p_{i|i}$ ), but only a **small cost** for using **nearby map points** to represent widely separated datapoints (using a large  $q_{i|i}$  to model a small  $p_{i|i}$ ).
    - In other words, SNE's lost function focuses on retaining the local structure of the data in the map (which is made possible thanks to this asymmetry).



#### Exercise - Gradient of SNE's Cost Function

The minimization of the cost function C is performed using gradient descent. Calculate the gradient of C w.r.t.  $y_i$ .



The derivative calculation is not so trivial, so we will break it up to parts:

· We denote the following:

$$q_{j|i} = rac{e^{-||y_i - y_j||^2}}{\sum_{k 
eq i} e^{-||y_i - y_k||^2}} = rac{E_{ij}}{\sum_{k 
eq i} E_{ik}} = rac{E_{ij}}{Z_i}$$

- Notice that  $E_{ij} = E_{ji}$  (because of the  $||\cdot||^2$ )
- ullet We express C as follows:

$$C = \sum_{i,j \neq i} p_{j|i} \log \frac{p_{j|i}}{q_{j|i}} = \sum_{i,j \neq i} p_{j|i} \log p_{j|i} - p_{j|i} \log q_{j|i} = \sum_{i,j \neq i} p_{j|i} \log p_{j|i} - p_{j|i} \log E_{ij} + p_{j|i} \log Z_i$$

• Notice that we got 3 expressions, and notice that the first one  $(p_{i|i} \log p_{i|i})$  does not depend on  $y_i$  and thus we can work only on the

We will now derive the derivative for the 2 parts separately and we change the indices from i, j to k, l to avoid confusion.

The second term:

$$rac{\partial}{\partial y_i} \sum_{k,l 
eq k} -p_{l|k} \log E_{kl}$$

- Now we ask: when is this derivative non-zero? Only when k=i or l=i

$$\begin{split} &= \sum_k -p_{k|i} \frac{\partial}{\partial y_i} \log E_{ik} - p_{i|k} \frac{\partial}{\partial y_i} \log E_{ki} \\ \bullet & \frac{\partial}{\partial y_i} E_{ik} = E_{ik} \cdot (-2(y_i - y_k)) \rightarrow \frac{\partial}{\partial y_i} \log E_{ik} = \frac{E_{ik} \cdot (-2(y_i - y_k))}{E_{ik}} = -2(y_i - y_k) \\ & \qquad \rightarrow \sum_k -p_{k|i} \frac{\partial}{\partial y_i} \log E_{ik} - p_{i|k} \frac{\partial}{\partial y_i} \log E_{ki} = \\ & \qquad \sum_k -p_{k|i} (-2(y_i - y_k)) - p_{i|k} \cdot 2(y_k - y_i) = 2 \sum_k (p_{k|i} + p_{i|k})(y_i - y_k) \end{split}$$

#### The Third Term

The third term:

$$rac{\partial}{\partial y_i} \sum_{k,l 
eq k} p_{l|k} \log Z_k$$

- · Notice that:
  - ullet  $\sum_{l 
    eq k} p_{l|k} = 1$  (as we modeled it as probability, it is not suprising)

$$\begin{array}{l} \blacksquare \ \, \overline{Z_k} \ \text{does not depend} \ \text{on} \ l \ \text{(in the} \ \underline{\sum} \ \text{operation), thus we can write the following:} \\ \frac{\partial}{\partial y_i} \sum_{k,l \neq k} p_{l|k} \log Z_k = \sum_{k,l \neq k} p_{l|k} \frac{\partial}{\partial y_i} \log Z_k = \sum_k \frac{\partial}{\partial y_i} \log Z_k \sum_{l \neq k} p_{l|k} = \sum_k \frac{\partial}{\partial y_i} \log Z_k \\ \end{array}$$

• Now we ask: when is this derivative non-zero? **Only** when k=i or m=i. ( $Z_k=\sum_{m 
eq k} E_{km}$ )

$$egin{aligned} & o \sum_k rac{\partial}{\partial y_i} \mathrm{log}\, Z_k = \sum_k rac{1}{Z_k} \sum_{m 
eq k} rac{\partial}{\partial y_i} E_{km} \ & = \sum_k rac{E_{ki} \cdot (2(y_k - y_i))}{Z_k} + \sum_k rac{E_{ik} (-2(y_i - y_k))}{Z_i} \ & = 2 \sum_k (-q_{k|i} - q_{i|k}) (y_i - y_k) \end{aligned}$$

Changing the index k o j, we get in total:

$$rac{\partial C}{\partial y_i} = 2\sum_j (p_{j|i} - q_{j|i} + p_{i|j} - q_{i|j})(y_i - y_j)$$

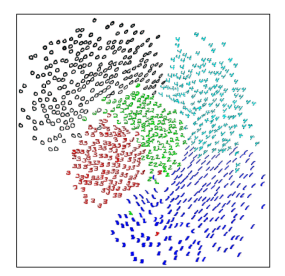
• Physics intuition: the expression of the gradient may be interpreted as the resultant force created by a set of springs between the map point  $y_i$ and all other map points  $y_j$ . The direction of the force is  $(y_i-y_j)$ . The spring between  $y_i$  and  $y_j$  repels or attracts the map points depending on whether the distance between the in the map is too small or too large to represent the similarities between the two high-dimensional points.



- The gradient descent is initialized by sampling map points randomly from an isotropic Gaussian with small variance that is centered around the origin.
- · In order speed up optimization and to avoid poor local minima, a relatively large momentum term is added to the gradient.
  - Or, the *current* gradient is added to an exponentially decaying sum of *previous gradient* in order to determine the changes in the coordinates of map points at each iteration of the gradient search.

$$\mathcal{Y}^{(t)} = \mathcal{Y}^{(t-1)} + \eta rac{\partial C}{\partial \mathcal{Y}} + lpha(t) ig( \mathcal{Y}^{(t-1)} - \mathcal{Y}^{(t-2)} ig)$$

- $\eta$  is the learning rate
- $\alpha(t)$  is the momentum at iteration t
- The weakness of SNE is that in order to avoid poor local minima, Gaussian noise is added to the map points after each iteration, which
  requires some sensible tuning.
  - In case of high-dimensionality, far values are accumulated in a "block around". This slippage is due to our willingness to adequately represent the neighbour's structure.
- The main problem with SNE: although it constructs reasonably good visualizations, its cost function is rather difficult to optimize by a problem called "the crowding problem". t-SNE aims to alleviate this problem.
- · Results on MNIST:

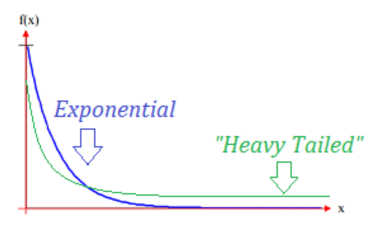




## t-Distributed Stochastic Neighbor Embedding (t-SNE)

As mentioned, SNE's cost function is problematic and hard to optimize. t-SNE's cost function differs from the one used by SNE in 2 ways:

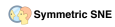
- 1. Symmetry it uses a symmetrized version of the SNE cost function with simpler gradients.
- Student-t Distribution it uses a Student-t distribution instead of a Gaussian to compute the similarity between two points in the low-dimensional space. t-SNE employs a heavy-tailed distribution in the low-dimensional space to alleviate both the crowding problem and the optimization problems of SNE.
- What is a heavy-tailed distribution?
  - A heavy tailed distribution has a tail that's heavier than an *exponential distribution*. In other words, a distribution that is heavy tailed goes to zero slower than one with heavy tails.
  - Heavy tailed distributions tend to have many outliers with very high values. The heavier the tail, the larger the probability that you'll get one or more disproportionate values in a sample.



Source: Data Science Central (https://www.statisticshowto.datasciencecentral.com/heavy-tailed-distribution/)

- Two important properties of heavy-tailed distributions:
  - The Central Limit (CLT) Theorem doesn't work.
  - Some moments don't exist, so order statistics are used instead.

We will now perform in-depth analysis of t-SNE's characteristics and improvements over SNE.



• Recall that in SNE we minimize the **sum of KL divergences** between the conditional probabilities  $p_{j|i}$  and  $q_{j|i}$ . It is also possible to minimize a **single KL divergence** between a *joint probability distribution* P in the high-dimensional space and a joint probability distribution Q in the low-dimensional space:

$$C = KL(P||Q) = \sum_i \sum_{j 
eq i} p_{ij} \log rac{p_{ij}}{q_{ij}}$$

- lacksquare Again  $p_{ii}=0$  and  $q_{ii}=0$
- ullet Symmetry notice the property that  $p_{ij}=p_{ji}, q_{ij}=q_{ji}$
- In symmetric SNE, the **pairwise similarities** in the *low-dimensional* map  $q_{ij}$  are given by

$$q_{ij} = rac{e^{-||y_i - y_j||^2}}{\sum_k \sum_{l 
eq k} e^{-||y_k - y_l||^2}}$$

- lacktriangle WAIT! How is this different from the expression for  $q_{j|i}$ ????
  - · Look again, whereas the nominator is the same, the denominator is different, and includes a sum of all possible distances!
- Similarly, the pairwise similarities in the high-dimensional space:

$$p_{ij} = rac{e^{-||x_i - x_j||^2/2\sigma^2}}{\sum_k \sum_{l 
eq k} e^{-||x_k - x_l||^2/2\sigma^2}}$$

#### **Exercise - Symmetric SNE Weakness**

Look again at the mapping in the low-dimensional and high-dimensional spaces. What will happen if the point in the high dimensional-space,  $x_i$ , is an **outlier** (i.e., all pairwise distances  $||x_i - x_j||^2$  are large)?



In that case, the values of  $p_{ij}$  are extremely small for all j, so the location of its low-dimensional map point  $y_i$  has very little effect on the cost function (remember the KL divegence form). As a result, the position of the map point is not well-determined by the positions of other map points.

ullet So how do we solve this **outliers** problem? We redefine the joint probabilities  $p_{ij}$  in the high-dimensional space to be the **symmetrized** conditional probabilities:

$$p_{ij} = rac{p_{j|i} + p_{i|j}}{2n}$$

- $p_{ij} = \frac{p_{j|i} + p_{i|j}}{2n}$  This ensures  $\sum_j p_{ij} > \frac{1}{2n}$  for all datapoints  $x_i$ , and thus each datapoint  $x_i$  makes a significant contribution to the cost function. The definition for  $q_{ij}$  is **the same as before**.
- The main advantage of the symmetric version of SNE is the simpler form of its gradient, which is faster (!) to compute:

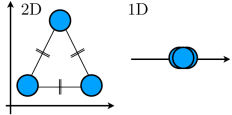
$$rac{\partial C}{\partial y_i} = 4 \sum_j (p_{ij} - q_{ij}) (y_i - y_j)$$



#### 🌋 The Crowding Problem

- The Crowding Problem the area of low-dimensional (2D, 3D) map that is available to accommodate moderately (medium) distant datapoints will not be nearly large enough compared with the area available to accommodate nearby datapoints.
  - ullet Thus, if we wish to model the small distances accurately in the map, most of the points that are at a moderate distance from datapoint i will have to be placed much too far away in the low-dimensional map.
  - In SNE, the "force" that connects datapoint i to each of these too-distant map points will be fairly small.
    - The very large number of such forces collapses together the points in the center of the map and prevents gaps from forming between the natural clusters.
  - The problem is not specific to SNE and also occurs in other local methods for dimensionality reduction such as Sammon mapping.
- Imaginative explanation: consider a set of datapoints that lie on a 2D curved manifold which is approximately linear on a small scale, and which is embedded within a higher-dimensional space. It is possible to model the small pairwise distances between datapoints fairly well in a 2D map (look at the "Swiss Roll" example). Now let's assume that the manifold has 10 intrinsic dimensions (that is, it can be represented with 10 variables, like the MNIST dataset which is comprised of images of 10 digits, so each digit can be represented as a variable, e.g. a one-hot vector of size 10). There are several reasons why the pairwise distances in a 2D map cannot faithfuly model distances between poins on the ten-dimensional manifold.
- For instance, in 10 dimensions, it is possible to have 11 datapoints that are evenly spaced and there is no way to model this faithfully in a 2 dimensions.



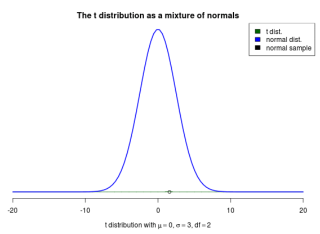


· There were attempts to solve this problem for SNE. The method is named UNI-SNE and it has a very complicated process of optimization. t-SNE also solves this problem as we will soon see.

- I
- Due to matching the joint probabilities of pairs of datapoints in the high-dimensional and low-dimensional spaces instead of their distances, there is a natural way to alleviate the **crowding problem** which is described in this section.
- In the high-dimensional space we convert distances to probabilities using a Gaussian distribution.
- In the low-dimensional map we can use a probability distribution that has much heavier tails than a Gaussian to convert distances into
  probabilities.
  - Why is this helpful? This allows a moderate (medium) distance in the high-dimensional space to be faithfully modeled by a much larger distance in the map. As a result, it eliminates unwanted attractive forces between map points that represent moderately dissimilar datapoints.
- In t-SNE a **Student-t** distribution with *one* degree of freedom (which turns it to a **Cauchy** distribution) is used as the heavy-tailed distribution in the low-dimensional map.
- The joint probabilities  $q_{ij}$  are defined as

$$q_{ij} = rac{(1 + ||y_i - y_j||^2)^{-1}}{\sum_k \sum_{l 
eq k} (1 + ||y_k - y_l||^2)^{-1}}$$

- Why **one** degree of freedom? It has a nice property that  $(1+||y_i-y_j||^2)^{-1}$  approaches an **inverse square law** for large pairwise distances  $||y_i-y_j||$  in the low-dimensional map. This makes the map's representation of joint probabilities (almost) invariant to changes in the scale of the map for map points that are far apart.
  - Also, it means that large clusters of points that are far apart interact the same way as individual points, so the optimization operates in the same way at all but the finest scales.
- Theoretically, the selection of Student-t is justified due to the fact that it is closely related to the Gaussian distribution, as it is an **infinite** mixture of Gaussians.
- Illustration:



Source: Rasmus Bååth's Research Blog (http://www.sumsar.net/blog/2013/12/t-as-a-mixture-of-normals/)

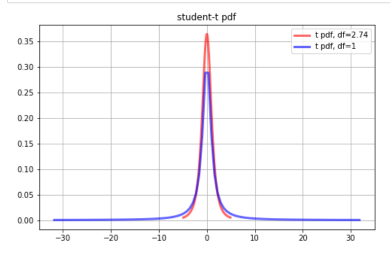
- Generally, Student-t proability density function (PDF) with u degrees of freedom is defined as:

$$f(x) = rac{\Gamma(rac{
u+1}{2})}{\sqrt{
u}\pi\Gamma(rac{
u}{2})}(1+rac{x^2}{
u})^{-rac{
u+1}{2}}$$

 $lacksquare \Gamma(z)$  is the Gamma function:  $\Gamma(z)=\int_0^\infty x^{z-1}e^{-x}dx$ 

```
In [2]: # student-t example, look how it looks like a normal distribution
        # doc @ https://docs.scipy.org/doc/scipy/reference/generated/scipy.stats.t.html
        def plot_student_t_example():
            # calculate the moments
            df = 2.74 # degrees of freedom, does not have to be discrete
            df 1 = 1
            mean, var, skew, kurt = t.stats(df, moments='mvsk')
            # press Shift+TAB inside the parenthesis to see argumnets description
            # let's see the pdf
            x = np.linspace(t.ppf(0.01, df), t.ppf(0.99, df), 100) # ppf - inverse cdf at q of given r.v. (df)
            x_1 = np.linspace(t.ppf(0.01, 1), t.ppf(0.99, 1), 100)
            # plotting
            fig = plt.figure(figsize=(8, 5))
            ax = fig.add_subplot(1,1,1)
            ax.plot(x, t.pdf(x, df), 'r-', lw=3, alpha=0.6, label='t pdf, df=2.74')
            ax.plot(x_1, t.pdf(x_1, df_1), 'b-', lw=3, alpha=0.6, label='t pdf, df=1')
            ax.set_title("student-t pdf")
            ax.grid()
            ax.legend()
```

#### In [3]: plot\_student\_t\_example()



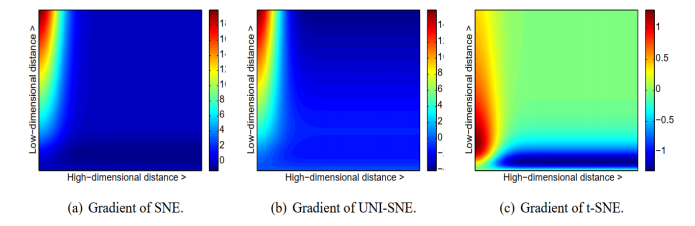
# The Gradient of t-SNE

• The gradient of the KL-divergence between P and and the Student-t joint-probablity distribution Q is given by

$$rac{\partial C}{\partial y_i} = 4 \sum_{j 
eq i} (p_{ij} - q_{ij}) (1 + ||y_i - y_j||^2)^{-1} (y_i - y_j)$$

• The proof is left for HW, but it uses the same tricks as the gradient for SNE.

• The following shows the gradients between two low-dimensional datapoints  $y_i$  and  $y_j$  as a function of their pairwise Euclidean distances in the high-dimensional ( $||x_i - x_j||$ ) and low-dimensional ( $y_i - y_j$ ) space:



- **Positive** values of the gradient represent an *attraction* between the low-dimensional datapoints  $y_i$  and  $y_j$ .
- **Negative** values of the gradient represent a *repulsion* between the low-dimensional datapoints  $y_i$  and  $y_j$ .
- The advantages of **t-SNE** gradient as seen above:
  - t-SNE's gradient **strongly repels** dissimilar datapoints that are modeled by a small pairwise distance in the low-dimensional represntation. SNE has similar behaviour, but its effect is minimal compared to the strong attractions elsewhere in the gradient.
  - Although t-SNE introduces strong repulsions between dissimilar points that are modeled by small pairwise distances, these repulsions don't go to infinity, which makes t-SNE different than UNI-SNE where the strength of the repulsion between very dissimilar datapoints is proportional to their pairwise distance in the low-dimensional map (ehich may cause dissimilar points to move too far away from each other).



#### t-SNE Optimization & Algorithm

- The optimization of the t-SNE cost function is much easier than the optimization of the cost functions of SNE and UNI-SNE.
- t-SNE introduces long-range forces in the low-dimensional map that can pull back together two clusters of similar points that get separated early on the optimization.

· The algorithm:

# Algorithm 1: Simple version of t-Distributed Stochastic Neighbor Embedding. Data: data set $X = \{x_1, x_2, ..., x_n\}$ , cost function parameters: perplexity Perp, optimization parameters: number of iterations T, learning rate $\eta$ , momentum $\alpha(t)$ . Result: low-dimensional data representation $\mathcal{Y}^{(T)} = \{y_1, y_2, ..., y_n\}$ . begin compute pairwise affinities $p_{j|i}$ with perplexity Perp (using Equation 1) set $p_{ij} = \frac{p_{j|i} + p_{i|j}}{2n}$ sample initial solution $\mathcal{Y}^{(0)} = \{y_1, y_2, ..., y_n\}$ from $\mathcal{N}(0, 10^{-4}I)$ for t=I to T do compute low-dimensional affinities $q_{ij}$ (using Equation 4) compute gradient $\frac{\delta C}{\delta \mathcal{Y}}$ (using Equation 5) set $\mathcal{Y}^{(t)} = \mathcal{Y}^{(t-1)} + \eta \frac{\delta C}{\delta \mathcal{Y}} + \alpha(t) \left(\mathcal{Y}^{(t-1)} - \mathcal{Y}^{(t-2)}\right)$ end end

- The **momentum** term,  $\alpha(t)$ , reduces the number of iterations requierd (and works best if it is small until the map points have become moderately well organized).
- The above version can be sped up using *adapative learning scheme*, which gradually increases the learning rate in directions in which the gradient is stable.
- Tricks to improve the algorithm:
  - "Early compression" forcing the map points to stay close together at the start of the optimization. When the distances between map points are small, it is easy for clusters to move through one another, which makes it easier to explore the space of possible global organizations of the data.
    - Implementation adding L2-penalty to the cost function that is proportional to the sum of squared distances of map points from the origin.
  - **"Early exaggeration"** multiplying all of the  $p_{ij}$ 's by, for example, 4, in the initial stages of the optimization. This means that almost all of the  $q_{ij}$ 's (which still add up to 1) are still too small to model their corresponding  $p_{ij}$ 's. This encourages the values of  $q_{ij}$ 's to become more focused on larger  $p_{ij}$ 's (i.e., closer points) which as a result causes natural clusters in the data to form tight widely separated clusters in the map (and in the process creating a lot of empty space in the map and allows cluster to move around more easily and find a good global optimization).



#### · Applying t-SNE to large data sets

- t-SNE has a computational and memory complexity that is *quadratic* in the number of datapoints (like many other visualizations techniques).
- This makes it almost infeasible to apply the standard version of t-SNE to data sets that contain more than 10,000 points.
  - The trivial soultion would be to just use a subset of the data, but this approach does not make use of all the information available.
  - There is a modification to t-SNE that uses random walk on a graph, you can read about it in the official paper (look in the "Credits" section).

- Dimensionality reduction for other purposes (other than visualization) it is unclear how t-SNE performs on the more general task of dimensionality reduction (using t-SNE to reduce to d > 3 dimensions).
  - Because of the heavy tails of the Student-t distribution, it is not enough to look at the behaviour of t-SNE on  $d \leq 3$  dimensions. In high-dimensional spaces, the heavy tails comprise a relatively large portion of the probability mass under the Student-t distribution, which may lead to d-dimensional data representations that **don't preserve the local structure of the data as well**. In the case of d > 3, it would be more approriate to use Student-t with more the one degree of freedom.
- Curse of intrinsic dimensionality t-SNE reduces the dimensionality of data mainly based on local properties of the data, which makes t-SNE sensitive to the curse of intrinsic dimensionality of data. In data sets with high intrinsic dimensionality and an underlying manifold that is highly varying, the local linearity assumption on the manifold that t-SNE implicitly makes (by using Euclidean distances between near neighbors) may be violated.
  - In other words, in complex manifolds, where even the local structure is complex, one cannot assume local linearity. In such cases, deep architectures, like autoencoders, may perform better.
- Non-convexity of the t-SNE cost function unlike other dimensionality reduction techniques such as PCA, Isomap and LLE, t-SNE's cost function is **not convex**, which requiers tuning of the optimization parameters.
  - This fact has important consequence: **t-SNE is non-deterministic**, that is, you can run it multiple times and get a different result each time!



#### **Example - The Wisconsin Breast Cancer Dataset**

We will now see how t-SNE works on the breast cancer dataset, both in 2D and 3D.

```
In [2]: # time to see it in action, let's load the breat cancer dataset
    dataset = pd.read_csv('./datasets/cancer_dataset.csv', index_col=-1)
    # print the number of rows in the data set
    number_of_rows = len(dataset)
    print('Number of rows in the dataset: {}'.format(number_of_rows))
    # separate labels from data
    X = dataset.drop(columns=['id', 'diagnosis'])
    y = dataset['diagnosis'].values == 'M'
    ## Show a sample 10 rows
    dataset.sample(10)
```

Number of rows in the dataset: 569

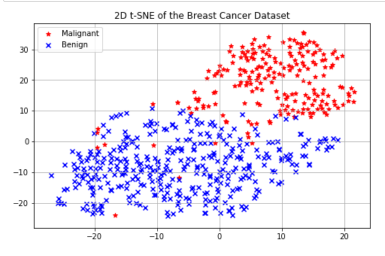
#### Out[2]:

	id	diagnosis	radius_mean	texture_mean	perimeter_mean	area_mean	smoothness_mean	compactness_mean	conca
NaN	871201	М	19.59	18.15	130.70	1214.0	0.11200	0.16660	
NaN	901549	В	11.27	12.96	73.16	386.3	0.12370	0.11110	
NaN	859465	В	11.31	19.04	71.80	394.1	0.08139	0.04701	
NaN	89511502	В	12.67	17.30	81.25	489.9	0.10280	0.07664	
NaN	8913	В	12.89	13.12	81.89	515.9	0.06955	0.03729	
NaN	906616	В	11.61	16.02	75.46	408.2	0.10880	0.11680	
NaN	8911230	В	11.33	14.16	71.79	396.6	0.09379	0.03872	
NaN	85638502	М	13.17	21.81	85.42	531.5	0.09714	0.10470	
NaN	8912284	В	12.89	15.70	84.08	516.6	0.07818	0.09580	
NaN	884689	В	11.52	14.93	73.87	406.3	0.10130	0.07808	

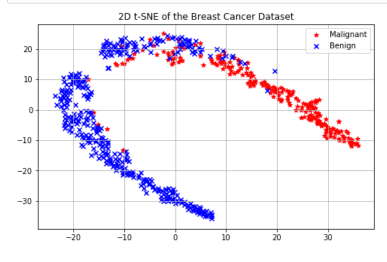
10 rows × 32 columns

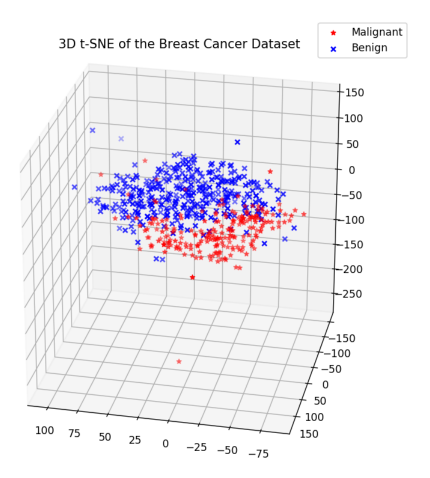
```
In [3]: def plot_tsne(X, dim=2, perplexity=30.0, scale_data=False):
                                      if dim < 2 or dim > 3:
                                                 print("OH NO :(")
                                                raise SystemError("2 <= dim <= 3")</pre>
                                     t_sne = TSNE(n_components=dim, perplexity=perplexity)
                                     if scale_data:
                                                scaler = StandardScaler()
                                                X = scaler.fit_transform(X)
                                     X_embedded = t_sne.fit_transform(X)
                                     if dim == 2:
                                                 fig = plt.figure(figsize=(8,5))
                                                 ax = fig.add_subplot(1,1,1)
                                                 ax.scatter(X\_embedded[y,\emptyset], X\_embedded[y,1], color='r', marker='*', label='Malignant')\\
                                                 ax.scatter(X_{embedded}[~y,0],~X_{embedded}[~y,~1],~color='b',~marker='x',~label='Benign')
                                                 ax.grid()
                                                 ax.legend()
                                                ax.set_title("2D t-SNE of the Breast Cancer Dataset")
                                     else:
                                                fig = plt.figure(figsize=(8, 8))
                                                 ax = fig.add_subplot(1, 1, 1, projection='3d')
                                                    ax.axis('equal')
                                                 ax.scatter(X\_embedded[y,\ 0],\ X\_embedded\ [y,\ 1],\ X\_embedded\ [y,\ 2],\ color='r',\ marker='*',\ label='Marker-'*',\ labe
                         alignant')
                                                ax.scatter(X\_embedded[\sim y,\ 0],\ X\_embedded\ [\sim y,\ 1],\ X\_embedded\ [\sim y,\ 2],\ color='b',\ marker='x',\ label
                         ='Benign')
                                                 ax.grid()
                                                 ax.legend()
                                                 ax.set_title("3D t-SNE of the Breast Cancer Dataset")
```

# In [4]: # let's try to reduce to 2 dimensions, with scaling plot\_tsne(X, dim=2, perplexity=30.0, scale\_data=True)



In [5]: # let's try to reduce to 2 dimensions, without scaling
plot\_tsne(X, dim=2, perplexity=30.0, scale\_data=False)







#### **Recommended Videos**



- These videos do not replace the lectures and tutorials.
- Please use these to get a better understanding of the material, and not as an alternative to the written material.

#### Video By Subject

- Student's T Distribution 365 Data Science Student's T Distribution (https://www.youtube.com/watch?v=32CuxWdOlow)
- t-SNE GoogleTechTalks Visualizing Data Using t-SNE (https://www.youtube.com/watch?v=RJVL80Gg3IA)



- Visualizing Data using t-SNE, der Maaten, Hinton and Bengio <a href="https://lvdmaaten.github.io/tsne/">https://lvdmaaten.github.io/tsne/</a> (<a href="https://lvdmaaten.github.io/">https://lvdmaaten.github.io/tsne/</a> (<a href="https://lvdmaaten.github.io/">https://lvdmaaten.github.io/</a> (<a href="https://lvdmaaten.github.io/">https://lvdmaaten.github.i
- Icons from Icon8.com (https://icons8.com/) https://icons8.com (https://icons8.com)
- Datasets from Kaggle (https://www.kaggle.com/) https://www.kaggle.com/ (https://www.kaggle.com/)