









# We're going to first learn about clustering algorithms & classifiers

**Clustering** = task of <u>grouping</u> a set of objects in such a way that objects in the same group (a **cluster**) are more similar (in some sense) to each other than to those in other groups (clusters).

Adapted from Wikipedia



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Let's motivate this with an important historical example:

## Distinct types of diffuse large B-cell lymphoma identified by gene expression profiling

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Nature 2000

"Diffuse large B-cell lymphoma (DLBCL), the most common subtype of non-Hodgkin's lymphoma ... is <u>one disease in which attempts to define subgroups on</u> <u>the basis of morphology have largely failed</u>..."

"DLBCL ... is clinically heterogeneous:

<u>40% of patients respond well</u> to current therapy and have prolonged survival, whereas <u>the remainder</u> <u>succumb</u> to the disease.

We proposed that this variability in natural history reflects unrecognized molecular heterogeneity in the tumours."

Nature 2000































	So, how does clustering work?
	Msamples
Ναρμος	The first clustering method we'll learn about simply groups the objects (samples or genes) in a hierarchy by the similarity of their feature vectors.
	Gene N, sample 1Gene N, sample jGene N, sample M













#### K-means clustering is a common alternative clustering approach \*mainly because it's easy and can be quite fast!\*

The basic algorithm:

- 1. Pick a number (k) of cluster centers
- 2. Assign each gene to its nearest cluster center
- 3. Move each cluster center to the mean of its assigned genes
- 4. Repeat steps 2 & 3 until convergence

See the K-means example posted on the web site











### Some features of K-means clustering

- Depending on how you seed the clusters, it may be stochastic. You may not get the same answer every time you run it.
- Every data point ends up in exactly 1 cluster (so-called *hard* clustering)
- Not necessarily obvious how to choose k
- Great example of something we've seen already: <u>Expectation-Maximization (E-M) algorithms</u>

EM algorithms alternate between assigning data to models (here, assigning points to clusters) and updating the models (calculating new centroids)



#### *k*-means

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### A fun clustering strategy that builds on these ideas: Self-organizing maps (SOMs)

SOMs have:

your data (points in some high-dimensional space) a grid of nodes, each node also linked to a point someplace in data space

1. First, SOM nodes are arbitrarily positioned in data space. Then:

2. Choose a training data point. Find the node closest to that point.

3. Move its position closer to the training data point.

4. Move its grid neighbors closer too, to a lesser extent.

Repeat 2-4. After many iterations, the grid approximates the data distribution.



























# BUT...

tSNE & the related technique UMAP lend themselves to misinterpretation, so <u>use caution in interpreting them!</u>

I recommend that you read "The specious art of single-cell genomics", by Tara Chari & Lior Pachter

https://journals.plos.org/ploscompbiol/article?id=10.1371/journal.pcbi.1011288

"In biology, single-cell expression studies almost always begin with reduction to two or three dimensions to produce 'all-in-one' visuals of the data that are amenable to the human eye, and these are subsequently used for qualitative and quantitative analysis of cell relationships. However, there is little theoretical support for this practice."



#### Some take aways

Data clustering and visualization are great to build some intuition for your data & ask questions like: *Are my data obviously clustered? What's that set of outliers over there?* ...etc...

But! High-dimensional data usually can't be perfectly represented in just 2- or 3-dimensions.

So, remember that most data visualization approaches (including SOMs, tSNE, and UMAP) <u>distort the true data relationships</u>. Try more than one approach and use caution in interpreting.