# DIMENSIONALITY REDUCTION AND VISUALIZATION

#### Loose ends from HW2

- Hyperparameters, bin size = 5, 10, 40, 100...?
  - Tune on test set error rate
- Variance of a recognizer
  - Accuracy 100%? 98? 90? 80?
  - What's the mean and variance of the accuracy?
- A majority class baseline
  - Powerful if one class dominates
  - Recognizer becomes biased towards the majority class (the prior term)
  - Often happens in real life
  - How to deal with this?

#### Loose ends from HW2

- What happens to P(x | leave), if there's no hk in the bin?
  - MLE estimates says P(a < x < b | leave) = 0</li>
  - 0 probability for the entire term
  - Is this due to a bad sampling of the training set?
  - Can solve with MAP

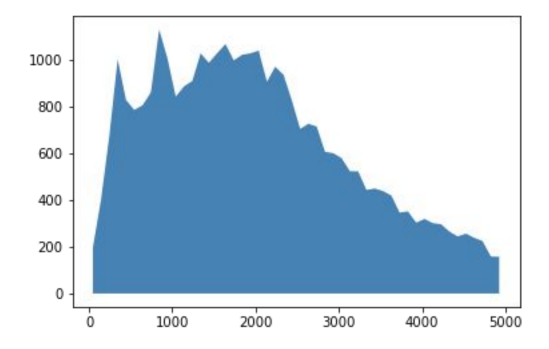
Map of a coin toss β, α are prior hyperparameters

$$\theta = \frac{k + \alpha - 1}{n + \alpha + \beta - 2}$$

Use unsupervised data for the priors?

#### Loose ends from HW2

- Another method to combat zero counts is to use Gaussian mixture models
  - How to select the number of mixtures?
    - Maybe all these can be a course project



# ML Methodology

 Re-train using the full set for deployment (using the hyperparameters tuned on test)

- But you should have a real test set to evaluate this
  - Kaggle submission
  - Held out data

Train set: learn parameters

Validation/dev set: learn hyper parameters

Test set: evaluate

## Evaluating a detection problem

4 possible scenarios

Actual	Yes
	No

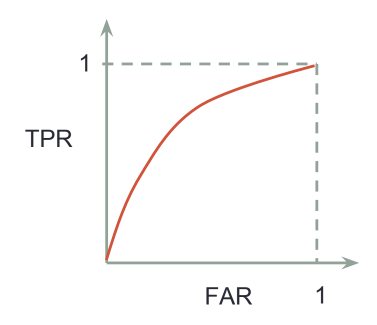
Detector	
Yes	No
True positive	False negative (Type II error)
False Alarm (Type I error)	True negative

True positive + False negative = # of actual yes False alarm + True negative = # of actual no

 False alarm and True positive carries all the information of the performance.

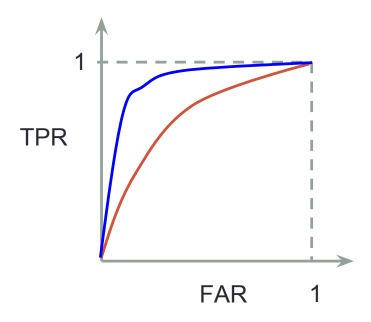
# Receiver operation Characteristic (RoC) curve

- What if we change the threshold
- FA TP is a tradeoff
- Plot FA rate and TP rate as threshold changes



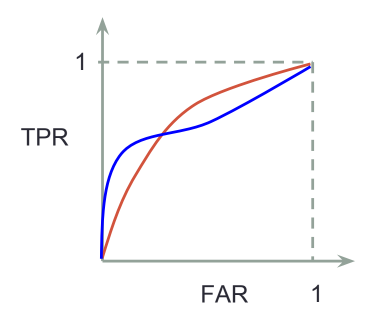
# Comparing detectors

• Which is better?



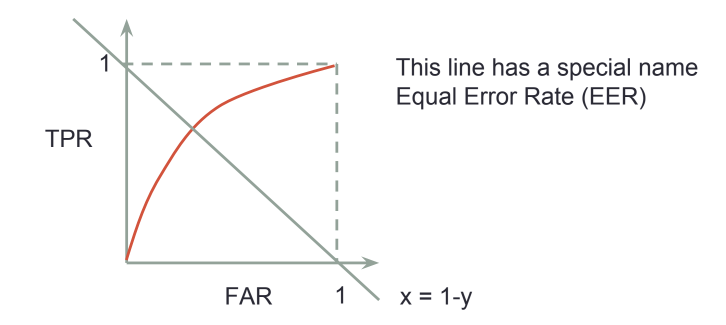
# Comparing detectors

• Which is better?



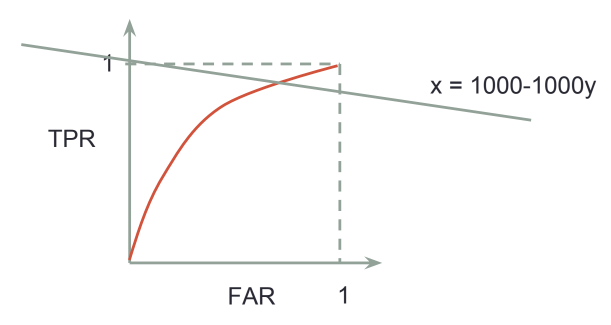
# Selecting the threshold

- Select based on the application
- Trade off between TP and FA. Know your application, know your users.
  - A miss is as bad as a false alarm
     FAR = 1-TPR => x = 1-y



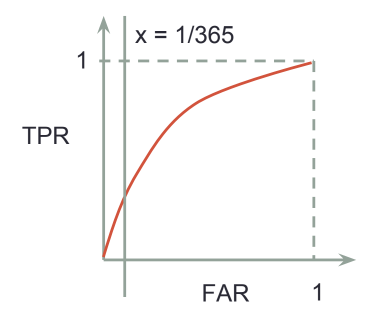
# Selecting the threshold

- Select based on the application
- Trade off between TP and FA. Know your application, know your users. Is the application about safety?
  - A miss is 1000 times more costly than false alarm.
    - FAR = 1000(1-TPR) => x = 1000-1000y



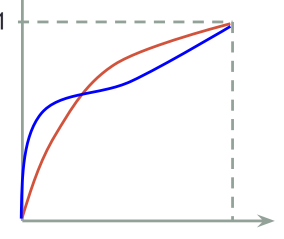
# Selecting the threshold

- Select based on the application
- Trade off between TP and FA.
  - Regulation or hard threshold
  - Cannot exceed 1 False alarm per year
    - If 1 decision is made everyday, FAR = 1/365



# Comparing detectors

**TPR** 



Which is better?

 You want to give your findings to a HR person FAR so that he will conduct an interview with the person which detector to pick?

# Interpretibility

HR asks what can they do better.

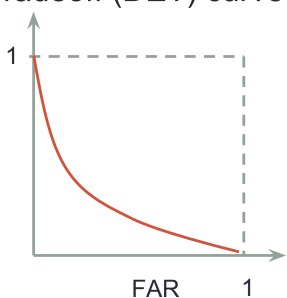
What should you tell them based on your model?

#### Notes about RoC

- Ways to compress RoC to just a number for easier comparison -- use with care!!
  - EER
  - Area under the curve
  - F score
- Other similar curve Detection Error Tradeoff (DET) curve

MR

- Plot False alarm vs Miss rate
- Can plot on log scale for clarity



# DIMENSIONALITY REDUCTION AND VISUALIZATION

#### Mixture models

$$p(x) = \sum_{k} p(k)p_k(x)$$

- A mixture of models from the same distributions (but with different parameters)
- Different mixtures can come from different sub-class
  - Cat class
    - Siamese cats
    - Persian cats
- p(k) is usually categorical (discrete classes)
- Usually the exact class for a sample point is unknown.
  - Latent variable

#### EM on GMM

- E-step
  - Set soft labels:  $w_{n,j}$  = probability that nth sample comes from jth mixture p
  - Using Bayes rule
    - $p(k|x; \mu, \sigma, \phi) = p(x|k; \mu, \sigma, \phi) p(k; \mu, \sigma, \phi) / p(x; \mu, \sigma, \phi)$
    - p(k|x; μ, σ, φ) α p(x|k; μ, σ, φ) p(k; φ)

$$p(k_n = j | x_n; \phi, \mu, \Sigma) = \frac{p(x_n; \mu_j, \sigma_j) p(k_n = j; \phi)}{\sum_l p(x_n; \mu_l, \sigma_l) p(k_n = l; \phi)}$$

#### EM on GMM

M-step (soft labels)

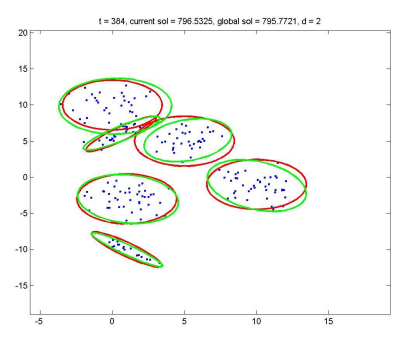
$$\phi_{j} = \frac{1}{N} \sum_{n=1}^{N} w_{n,j}$$

$$\mu_{j} = \frac{\sum_{n=1}^{N} w_{n,j} x_{n}}{\sum_{n=1}^{N} w_{n,j}}$$

$$\sigma_{j}^{2} = \frac{\sum_{n=1}^{N} w_{n,j} (x_{n} - \mu_{j})^{2}}{\sum_{n=1}^{N} w_{n,j}}$$

#### EM/GMM notes

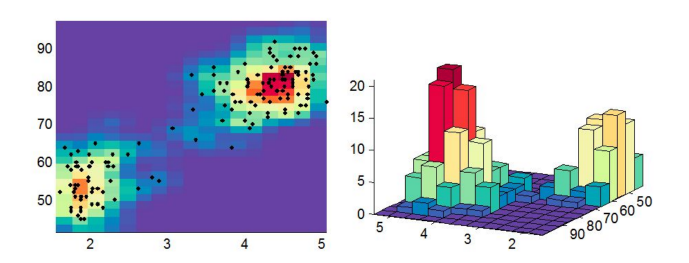
- Converges to local maxima (maximizing likelihood)
  - Just like k-means, need to try different initialization points
- What if it's a multivariate Gaussian?
  - The grid search gets harder as the number of number of dimension grows



https://www.mathworks.com/matlahcentral/fileeychange/7055-multivariate-gaussian-mixture-model-ontimization-hy-cross-entropy

# Histogram estimation in N-dimension

- Cut the space into N-dimensional cube
  - How many cubes are there?
  - Assume I want around 10 samples per cube to be able to estimate a nice distribution without overfitting. How many more samples do I need per one additional dimension?

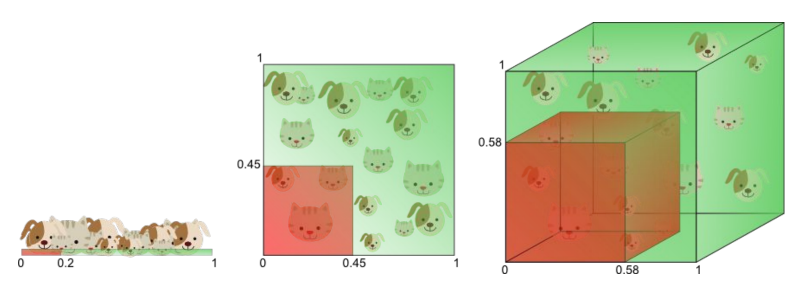


# The curse of dimensionality



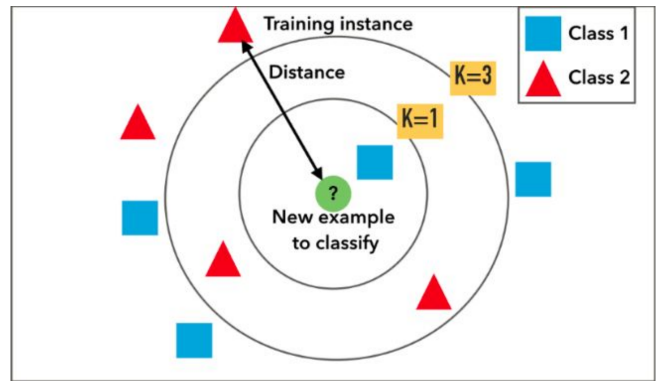
# The Curse of Dimensionality

- Harder to visualize or see structure of
  - Verifying that data come from a straight line/plane needs n+1 data points
- Hard to search in high dimension More runtime
- Need more data to get a get a good estimation of the data



# Nearest Neighbor Classifier

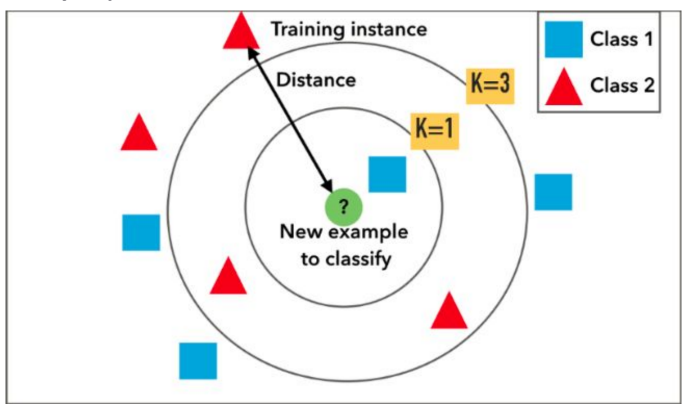
- The thing most similar to the test data must be of the same class Find the nearest training data, and use that label
- Use "distance" as a measure of closeness.
- Can use other kind of distance besides Euclidean



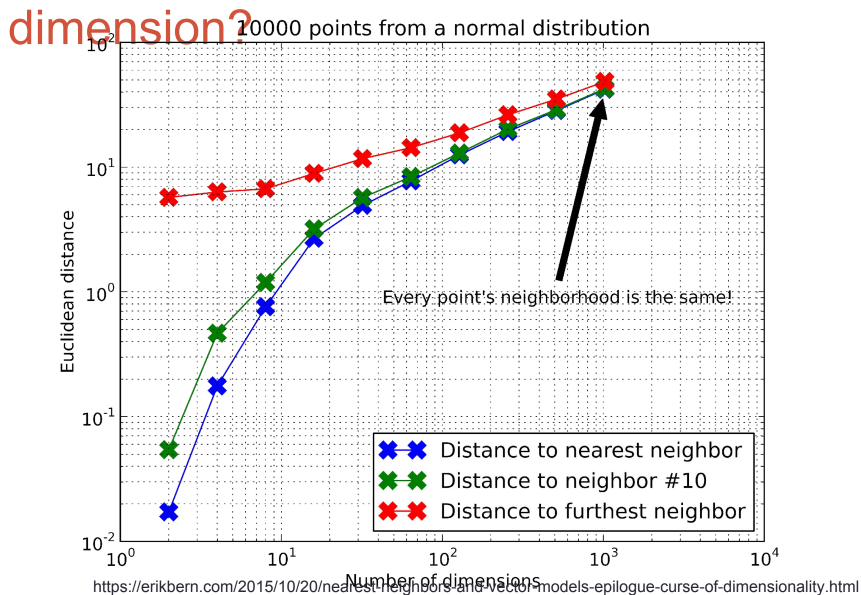
https://arifuzzamanfaisal.com/k-nearest-neighbor-regression/

## K-Nearest Neighbor Classifier

- Nearest neighbor is susceptible to label noise
- Use the k-nearest neighbors as the classification decision
  - Use majority vote



# What's wrong with knn in high



### Combating the curse of dimensionality

- Feature selection
  - Keep only "Good" features
- Feature transformation (Feature extraction)
  - Transform the original features into a smaller set of features

#### Feature selection vs Feature transform

- Keep original features
  - Useful for when the user wants to know which feature matters
    - But, correlation does not imply causation...

- New features (a combination of old features)
- Usually more powerful
  - Captures correlation between features

#### Feature selection

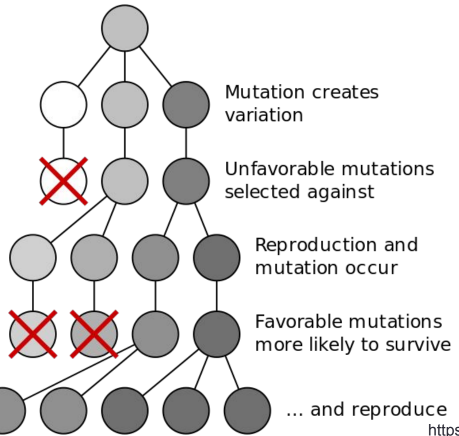
- Hackathon level (time limit days-a week)
  - Drop missing features
  - Low variance rows
    - A feature that is a constant is useless. Tricky in practice
  - Forward or backward feature elimination
    - Greedy algorithm: create a simple classifier with n-1 features, n times.
       Find which one has the best accuracy, drop that feature. Repeat.

#### Feature selection

- Proper methods
  - Algorithm that handles high dimension well and do selection as a by product
  - Tree-based classifiers
    - Random forest
  - Adaboost
  - Genetic Algorithm

# Genetic Algorithm

- A method based inspired by natural selection
  - No theoretical guarantees but often work



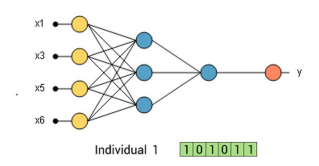
https://elitedatascience.com/dimensionality-reduction-algorithms

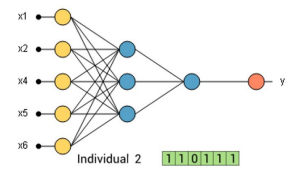
# Genetic Algorithm

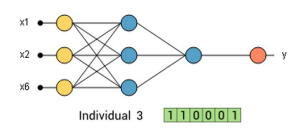
- Initialization
  - Create N classifiers, each using different subset of features
- Selection process
  - Rank the N classifiers according to some criterion, kill the lower half
- Crossover
  - The remaining classifier breeds offsprings by selecting traits from the parents
- Mutation
  - The offsprings can have mutations by random in order to generate diversity
- Repeat till satisfied

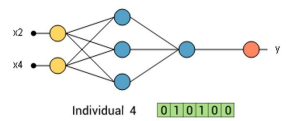
#### Initialization

- Create N classifiers
- Randomly select a subset of features to use









# Selection process

 Score the classifiers and kill the lower half (the amount to kill is also a parameter)

	Selection error	Rank
Individual 1	0.9	1
Individual 2	0.6	3
Individual 3	0.7	2
Individual 4	0.5	4

#### Crossover

Breed offsprings by randomly select genes from parents

Individual 3	1 1 0 0 0 1
Individual 4	0 1 0 1 0 0
Offspring 1	0 1 0 1 0 1
Offspring 2	1 1 0 1 0 1
Offspring 3	0 1 0 1 0 1
Offspring 4	1 1 0 0 0 0

#### **Mutation**

- Offspring can mutate with some probability to introduce diversity
- Mutation rate is usually 1/k where k is the number of features.
  - On average you mutate once per individual

Offspring1: Original 0 1 0 1 0 1

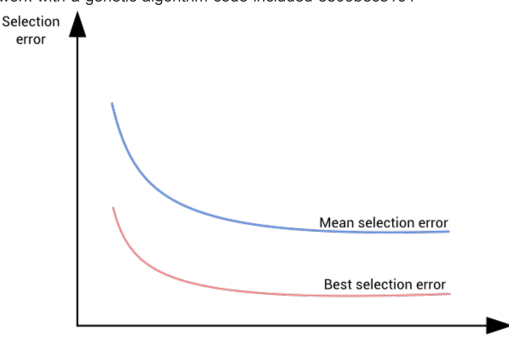
Offspring1: Mutated 0 1 0 0 0

#### Performance

- Usually performs well. The general population usually gets better (mean). The best performing (individual) also gets better after each generation
- Can be use to tune neural networks!

https://blog.coast.ai/lets-evolve-a-neural-network-with-a-genetic-algorithm-code-included-8809bece164

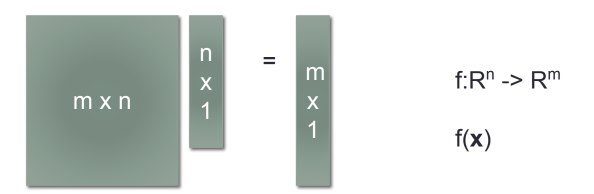
error



#### Feature transformation

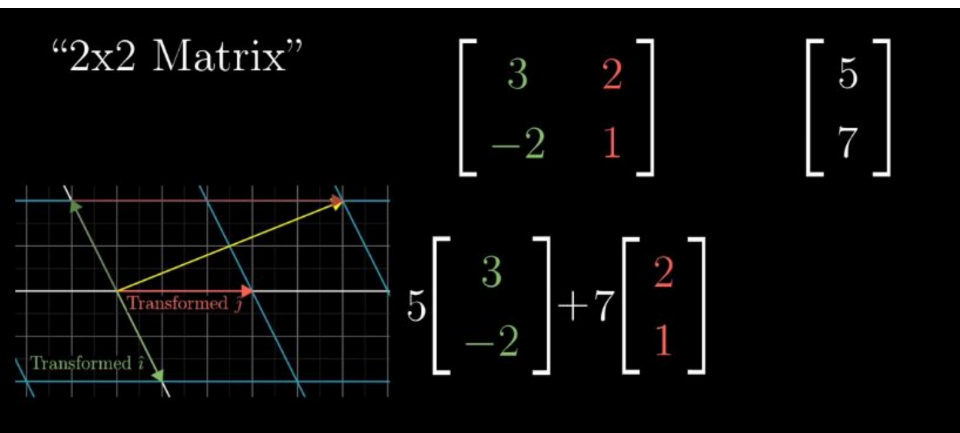
- Principle Component Analysis
- Linear Discriminant Analysis (NOT Latent Dirichlet Allocation)
- Random Projections

 Think Sets and Functions, rather than manipulation of number arrays/rectangles



https://www.linkedin.com/pulse/key-machine-learning-prereq-viewing-linear-algebra-through-ashwin-rao

Matrix as a sequence of column vectors

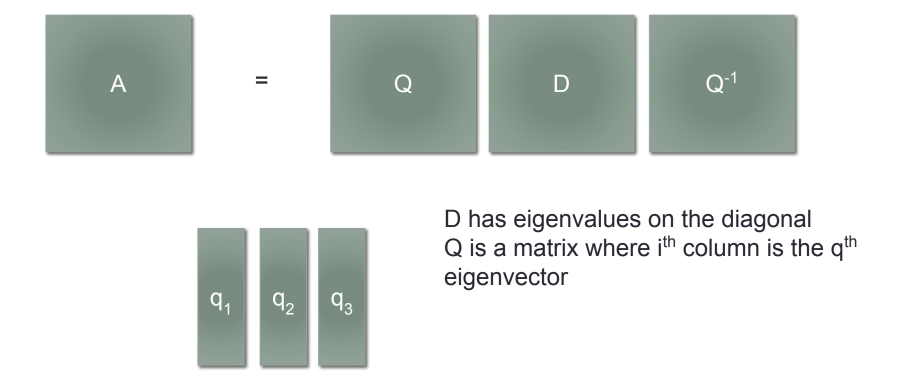


 Understand Matrix Factorizations as Compositions of "Simple" Functions:

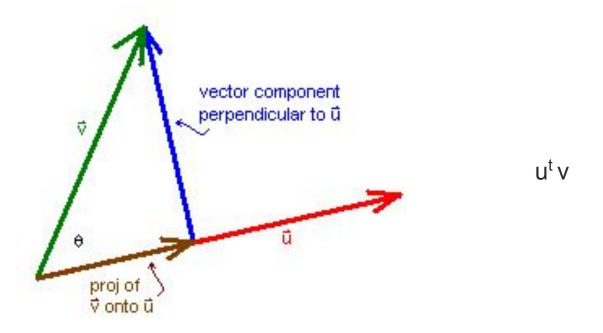


$$h(\mathbf{x}) = k(d(\mathbf{x}))$$

 View Eigendecomposition (ED) and Singular Value Decomposition (SVD) as rotations and stretches



- Projection as a change of basis
- Change basis from x,y coordinates to be on u



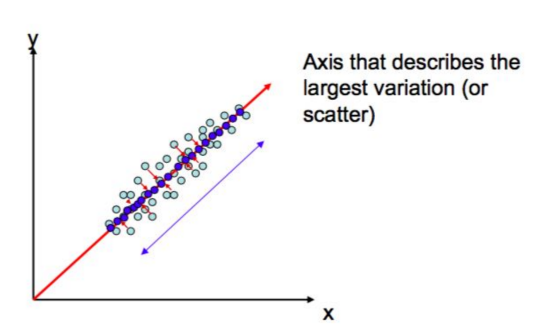
# Positive semi-definite and eigendecomposition

- Covariance matrix is positive semi-definite and symmetric.
- Eigenvalues are nonnegative, eigenvalues and vectors are real values, eigenvectors are mutually orthogonal.

$$\cdot q_i^t q_j = 0 \text{ for } i != j$$

#### What is PCA?

- We want to reduce the dimensionality but keep useful information
  - What is useful information? Variation
- We want to find a projection (a transformation) that describe maximum variation



#### **Formulation**

- Maximize the variance after projection ie
  - argmax Var(w<sup>t</sup> x)
- Subject to w is a unit vector

•  $\Sigma w = \lambda w <$  - eigenvector

## Trace properties

```
_{1} • tr (a) = a
2 • trA = trA^T
3 \cdot tr(A+B) = trA+trB
4 • tr(aA) = atr(A)
5 \nabla_A trAB = B^T
6 \nabla_{A^T} f(A) = (\nabla_A f(A))^T
\nabla_A tr A B A^T C = C A B + C^T A B^T
 \nabla_{A^T} tr A B A^T C = B^T A^T C^T + B A^T C
```

## So we got to eigenvectors

- A dxd covariance matrix has d eigen vectors/values pair.
   Do we use all of them?
- Which pair to use?

## Selecting eigenvectors

Remember the variance of projected data is

$$\omega^{\mathsf{T}} \Sigma \omega$$
. (1)

And our solution yielded

$$\Sigma \omega = \lambda \omega \tag{2}$$

Plug (2) in (1) and we get

projected variance = 
$$\omega^T \Sigma \omega = \omega^T \lambda \omega$$
  
=  $\lambda \omega^T \omega$  (remember ||  $\omega$ ||=1)  
=  $\lambda$ 

#### PCA

- The direction vector captures the variance corresponding to the eigenvalue
- So we want the higher eigenvalues
  - How many?

#### Matrix rank

- A square dxd matrix has full rank (e.g. rank d) if the columns are linearly independent.
- The number of linearly independent columns is the rank of the matrix
- A covariance matrix of size dxd will have have at most
   N-1 rank where N is the number of training samples
  - 640x640 images =  $\sim 400000$  dimensions
  - 1000 training images
  - The covariance matrix will be at most rank 999. The missing rank is because of the mean.

#### PCA

- The direction vector captures the variance corresponding to the eigenvalue
- So we want the higher eigenvalues
- Take the eigenvalues with non-zero eigenvalues (at most N-1 non-zero eigenvalues)

## Basis decomposition

- Let's consider our projection w<sub>i</sub> which is the eigenvectors to be a basis vector v<sub>i</sub>
- We can represent any vector as a sum of basis vectors as follows:

$$\mathbf{x} = \sum_{i=1}^{N} p_i \mathbf{v}_i = p_1 \begin{bmatrix} | \\ \mathbf{v}_1 \\ | \end{bmatrix} + p_2 \begin{bmatrix} | \\ \mathbf{v}_2 \\ | \end{bmatrix} + .. + p_n \begin{bmatrix} | \\ \mathbf{v}_n \\ | \end{bmatrix} = \mathbf{V} \mathbf{p}$$

## Finding the weights

$$\mathbf{x} = \sum_{i=1}^{N} p_i \mathbf{v}_i = p_1 \begin{bmatrix} | \\ \mathbf{v}_1 \\ | \end{bmatrix} + p_2 \begin{bmatrix} | \\ \mathbf{v}_2 \\ | \end{bmatrix} + .. + p_n \begin{bmatrix} | \\ \mathbf{v}_n \\ | \end{bmatrix} = \mathbf{V} \mathbf{p}$$

If v<sub>i</sub> are orthogonal, the projection of x onto v<sub>i</sub> gives p<sub>i</sub>

$$\mathbf{V}^{\mathsf{T}}\mathbf{x} = \begin{bmatrix} - & \mathbf{v}_1 & - \\ - & \mathbf{v}_2 & - \\ - & \mathbf{v}_3 & - \end{bmatrix} \begin{bmatrix} | \\ \mathbf{x} \\ | \end{bmatrix} = \begin{bmatrix} p_1 \\ p_2 \\ p_3 \end{bmatrix}$$

#### Means

- In PCA, we model variance. (Variation around the mean)
- In our projection we need to remove the mean

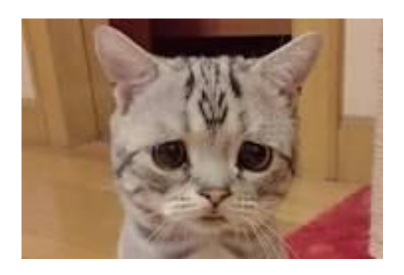
$$\mathbf{p} = \mathbf{V}^{\mathrm{T}}(\mathbf{x} - \mathbf{m})$$

- The mean is the mean of all your training data
- If we want to reconstruct the data we need to add back the mean

$$\mathbf{x} = \sum_{i=1}^{N} p_i \mathbf{v}_i + \mathbf{m} = p_1 \begin{bmatrix} 1 \\ \mathbf{v}_1 \\ 1 \end{bmatrix} + p_2 \begin{bmatrix} 1 \\ \mathbf{v}_2 \\ 1 \end{bmatrix} + ... + p_n \begin{bmatrix} 1 \\ \mathbf{v}_n \\ 1 \end{bmatrix} + \mathbf{m} = \mathbf{V}\mathbf{p} + \mathbf{m}$$

#### Practical issues

- If your data has different magnitudes in different dimensions, normalize each dimension before PCA
- If we have 640x640 images =  $\sim 400000$  dimensions.
- What is the size of the covariance matrix?



#### Practical issues

- You have N training examples.
- For the case where N << 400000, we only have N-1 eigen values we care about anyway

#### **Gram Matrix**

$$\Sigma = E(\mathbf{x} - \mu)(\mathbf{x} - \mu)^T = \mathbf{X}\mathbf{X}^T$$

Covariance matrix is the outer-product of the input matrix

Must solve 
$$\Sigma v = \lambda v$$
 
$$XX^Tv^=\lambda v \text{ (pre-mult by } X^T) \text{ (1)}$$
 
$$X^TXX^Tv^=\lambda X^Tv \text{ (} v'=X^Tv) \text{ (2)}$$
 Solve eigenvalue problem  $X^TXv'=\lambda v'$ 

X<sup>T</sup>X is a gram of inner-product matrix. Its size is NxN where N is the number of data samples.

## But how to get v from v'?

- From previous slide, equation (1) and (2)
  - $XX^Tv = \lambda v$  (1)
  - $v' = X^T v (2)$
- Substitute (2) into (1)
  - $Xv' = \lambda v$
- Thus, v = Xv'. We don't care about the scaling term because we will always scale the eigenvector so that it is orthonormal i.e. ||v|| = 1.

## How many eigenvectors?

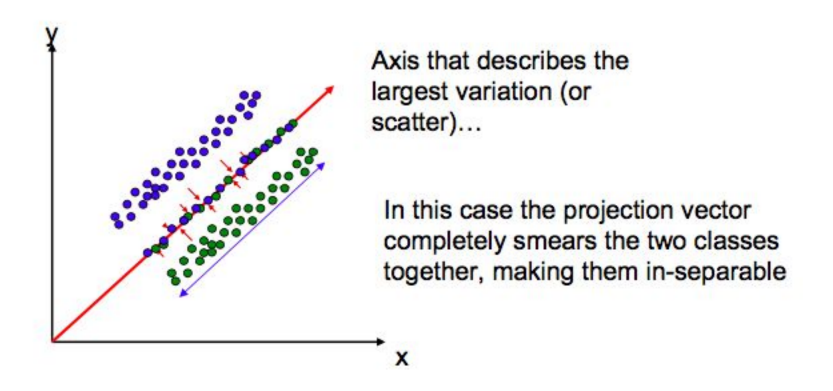
- Select based on amount of variance explained
  - Sum of eigenvalues exceeds some percent of total
- Reconstruction error

$$\mathbf{x} = \sum_{i=1}^{N} p_i \mathbf{v}_i = p_1 \begin{bmatrix} 1 \\ \mathbf{v}_1 \\ 1 \end{bmatrix} + p_2 \begin{bmatrix} 1 \\ \mathbf{v}_2 \\ 1 \end{bmatrix} + ... + p_n \begin{bmatrix} 1 \\ \mathbf{v}_n \\ 1 \end{bmatrix} = \mathbf{V} \mathbf{p}$$

 Select enough v so that the difference between original x and reconstructed x is small

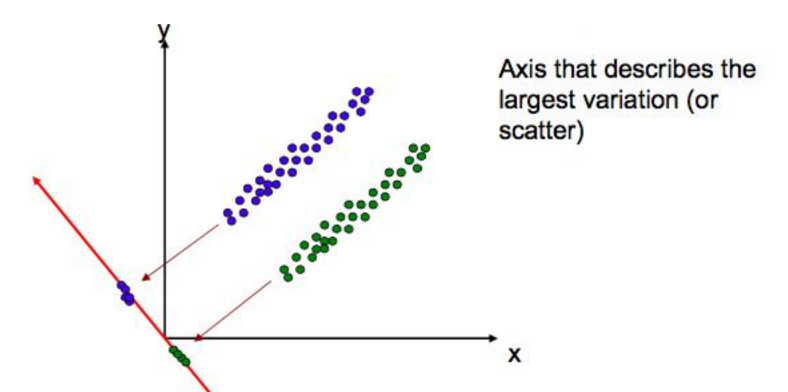
#### PCA for classification

PCA does not cares about the class labels



#### What is LDA

- Find the projections that separate the classes.
- Assumes unimodal Gaussian model for each class
  - Maximize the distance between the means and minimize the variance of each class -> best classification performance



## Simple 2 class case

 We want to maximize the distance between the projected means:

e.g. maximize 
$$|(\tilde{\mu}_1 - \tilde{\mu}_2)|^2$$

Where  $\tilde{\mu}_1$  is the projected mean  $\mu_1$  of class onto LDA direction vector  $\mathbf{w}$ , i.e.

$$\tilde{\boldsymbol{\mu}}_{1} = \mathbf{w}^{T} \boldsymbol{\mu}_{1}$$
and for class 2: 
$$\tilde{\boldsymbol{\mu}}_{2} = \mathbf{w}^{T} \boldsymbol{\mu}_{2} \text{ thus}$$

$$|(\tilde{\boldsymbol{\mu}}_{1} - \tilde{\boldsymbol{\mu}}_{2})|^{2} = |(\mathbf{w}^{T} \boldsymbol{\mu}_{1} - \mathbf{w}^{T} \boldsymbol{\mu}_{2})|^{2}$$

$$= \mathbf{w}^{T} (\boldsymbol{\mu}_{1} - \boldsymbol{\mu}_{2})^{T} (\boldsymbol{\mu}_{1} - \boldsymbol{\mu}_{2})^{T} \mathbf{w}$$

$$= \mathbf{w}^{T} \mathbf{S}_{R} \mathbf{w}$$

## Between class scatter matrix S<sub>B</sub>

$$(\tilde{\mu}_1 - \tilde{\mu}_2)^2 = (\mathbf{w}^T \boldsymbol{\mu}_1 - \mathbf{w}^T \boldsymbol{\mu}_2)^2$$

$$= \mathbf{w}^T (\boldsymbol{\mu}_1 - \boldsymbol{\mu}_2)^T (\boldsymbol{\mu}_1 - \boldsymbol{\mu}_2)^T \mathbf{w}$$

$$= \mathbf{w}^T \mathbf{S}_{\mathbf{B}} \mathbf{w}$$

We want to maximize w<sup>T</sup>S<sub>B</sub>w where S<sub>B</sub> is the between class scatter matrix defined as:

$$S_B = (\mu_1 - \mu_2)(\mu_1 - \mu_2)^T$$

## We also want to minimize within class scatter

 The variance or scatter of each class. We also want to minimize them.

$$\tilde{s}_1^2 = \sum_{i=1}^{N_1} (\tilde{x}_i - \tilde{\mu}_1)^2$$

Minimize the total scatter

$$\tilde{s}_{1}^{2} + \tilde{s}_{2}^{2}$$

#### Within class scatter

Lets expand on scatter s<sub>1</sub>,s<sub>2</sub>

$$\tilde{\mathbf{s}}_{1}^{2} = \sum_{i=1}^{N_{1}} (\tilde{\mathbf{x}}_{i} - \tilde{\boldsymbol{\mu}}_{1})^{2}$$

$$= \sum_{i=1}^{N_{1}} (\mathbf{w}^{T} \mathbf{x}_{i} - \mathbf{w}^{T} \boldsymbol{\mu}_{1})^{2}$$

$$= \sum_{i=1}^{N_{1}} \mathbf{w}^{T} (\mathbf{x}_{i} - \boldsymbol{\mu}_{1}) (\mathbf{x}_{i} - \boldsymbol{\mu}_{1})^{T} \mathbf{w}$$

$$= \mathbf{w}^{T} \mathbf{S}_{1} \mathbf{w}$$

#### Total within class scatter

We want to minimize

$$\tilde{s}_{1}^{2} + \tilde{s}_{2}^{2}$$

This is the same as

$$\mathbf{S}_{\mathbf{w}} = \sum_{i=1}^{C} \sum_{j=1}^{Ni} (\mathbf{x}_{j} - \boldsymbol{\mu}_{i}) (\mathbf{x}_{j} - \boldsymbol{\mu}_{i})^{\mathrm{T}}$$

C number of classes, Ni number of images from class i

#### Fisher Linear Discriminant Criterion

- We want to maximize between class scatter
- We want to minimize within class scatter
- We have an objective function as a ratio so we can achieve both!

$$J(\mathbf{w}) = \frac{|(\tilde{\mu}_1 - \tilde{\mu}_2)|^2}{\tilde{s}_1^2 + \tilde{s}_2^2}$$
$$J(\mathbf{w}) = \frac{\mathbf{w}^T \mathbf{S}_B \mathbf{w}}{\mathbf{w}^T \mathbf{S}_W \mathbf{w}}$$

#### LDA solution

If you do calculus

$$S_B w = \lambda S_W w$$

$$\mathbf{S_{w}}^{-1}\mathbf{S_{B}w} = \lambda \mathbf{w}$$

If S<sub>w</sub> is non-singular and invertible.

- Generalized eigenvalue problem. The number of solutions is min(rankS<sub>R</sub>, rankS<sub>W</sub>) = C-1 or N-C
- For 2 class this simplifies to
  - Note this is only one projection direction

$$\mathbf{w} = \mathbf{S_w}^{-1} (\boldsymbol{\mu_1} - \boldsymbol{\mu_2})$$

#### LDA+PCA

- First do PCA to reduce dimension
- Then do LDA to maximize classification ability
- How many dimensions to PCA?
  - Do PCA to keep N-C eigenvectors -> Makes S<sub>w</sub> full rank and invertible
  - Then, do LDA and compute C-1 projections in this N-C subspace
- PCA+LDA = Fisher projection

## Random projection

- Original d-dimensional data is project to k-dimensional subspace
- Using a random k x d matrix R with unit norm columns
  - Johnson-Lindenstrauss lemma: If points in a vector space are projected onto a randomly selected subspace of suitably high dimension, then the distances between the points are approximately preserved
- Elements of R are usually selected from Gaussians.
  - Generally any zero mean unit variance distribution would satisfy Johnson-Lindenstrauss lemma.

## Random projection notes

- R is not generally orthogonal.
  - But in a substantially large subspace, random vectors might be close to orthogonal.
- Looks weird but works...

#### Visualization

#### Methods covered

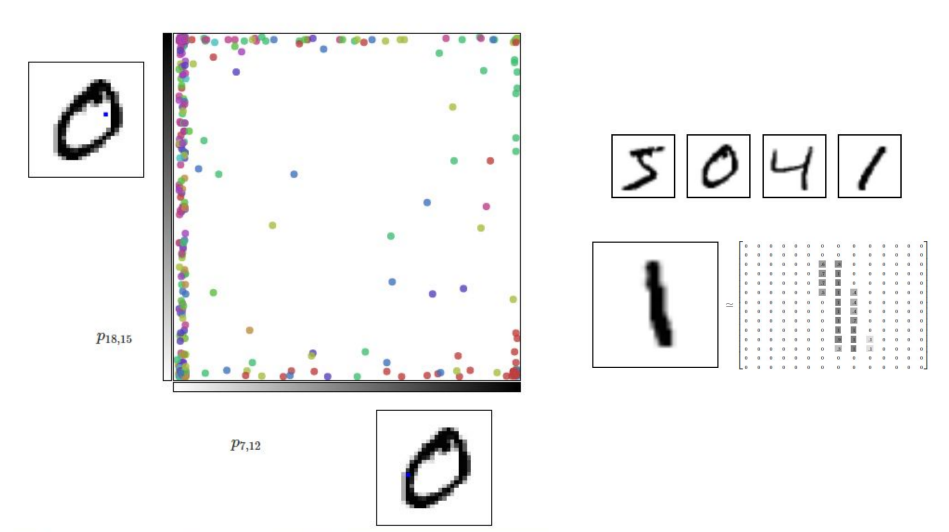
- 1. Reducing reconstruction error: PCA
- 2. Keeping direction of maximum seperability: LDA
- 3. Preserving distance (globally): RP

These are usually useful for downstream machine learning methods. (Classification/Regression/Clustering)

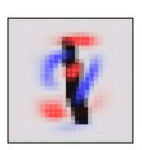
But what if we, as humans, want to get a sense of our data?

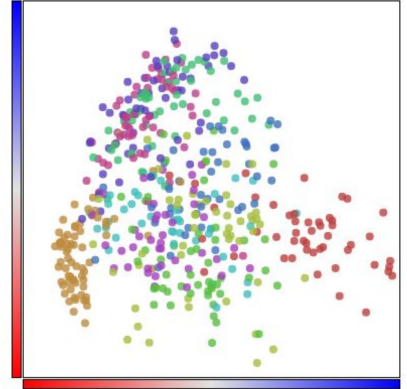
1. Interpretibility (in some sense): ????

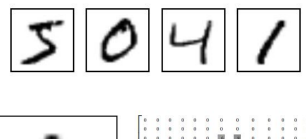
# Visualizing MNIST

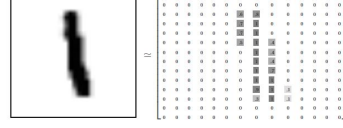


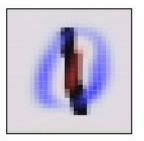
## PCA with MNIST











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

Preserves neighbor (preserves local distance).

- Things close together should be close together in the projected space
- Prefer using few projected dimensions (2-3)

## Defining neighbors

Define  $\mathsf{P}_{\mathsf{j}|\mathsf{i}}$  probability that i would pick j as its neighbor Assume i picks proportional to Gaussian centered at i  $exp(-||x_i-x_j||^2)/2\sigma_i^2$ 

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

 $P_{i|i}$  = 0 since we don't want to have it pick itself. The variance is fixed to some value.

## Defining neighbors

Define  $\mathbf{q}_{\mathbf{j}|\mathbf{i}}$  probability that i would pick j as its neighbor Assume i picks proportional to Gaussian centered at i  $exp(-||x_i-x_j||^2)/2\sigma_i^2$ 

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

When projected to set of points  $\{y_i\}$ , define  $q_{j|i}$  the probability that i would pick j in embedding/latent space

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

We set the variance in the y space to be 1/sqrt(2)

## Defining neighbors

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

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

We expect p and q to be the same -> small distance

How to measure distance between probability functions? Kullback-Leibler (KL) divergence

## KL divergence

Distance between two distributions

$$D_{KL}(P||Q) = \sum_{i} P(i)lograc{P(i)}{Q(i)} = -\sum_{i} P(i)lograc{Q(i)}{P(i)}$$

Note  $D_{KL}(P||Q) \neq D_{KL}(Q||P)$  (Not a real distance) Always positive. Equals 0 iff Q = P at every point.

$$P(head) = 0.5 P(tail) = 0.5$$

$$Q(head) = 0.7 Q(tail) = 0.3$$

$$D_{KI}(P||Q) = 0.5*In 0.5/0.7 + 0.5*In 0.5/0.3 = 0.087$$

$$D_{KI}(Q||P) = 0.7*In 0.7/0.5 + 0.3*In 0.3/0.5 = 0.082$$

#### Loss function

$$p_{j|i} = rac{exp(-||x_i - x_j||^2)/2\sigma_i^2}{\sum_{k 
eq i} exp(-||x_i - x_k||^2)/2\sigma_i^2} \hspace{0.5cm} q_{j|i} = rac{exp(-||y_i - y_j||^2)}{\sum_{k 
eq i} exp(-||y_i - y_k||^2)}$$

We expect p and q to be the same -> small distance

Loss function A

All points i KL computes over j

$$\sum_i D_{KL}(p_i||q_i)$$

$$D_{KL}(P||Q) = \sum_i P(i) log rac{P(i)}{Q(i)}$$

Note P can be considered as the weight for the distance Where p is large but q is small -> large penalty q is small but p is large -> small penalty

D(p||q) focuses on local structure in p



What are we minimizing wrt?
How to minimize loss?

## From SNE to t-SNE

Symmetric density t-distributed

## Symmetric with joint probability

$$\mathsf{p}_{\mathsf{i}\mathsf{l}\mathsf{i}}$$
  $eq \mathsf{p}_{\mathsf{i}\mathsf{l}\mathsf{i}}$   $eq D_{KL}(P||Q) = \sum_i P(i)lograc{P(i)}{Q(i)}$ 

A point i far away from everything will have small pi

Location of points in q no longer matter

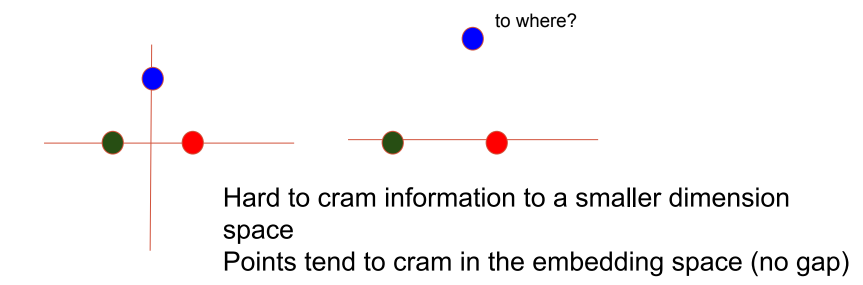
Create "joint probability"  $p_{ij} = p_{ji} = (p_{i|j} + p_{j|i})/2$ 

Each data point will contribute to the loss

Use instead of conditional probability in KL divergence

#### t-distributed

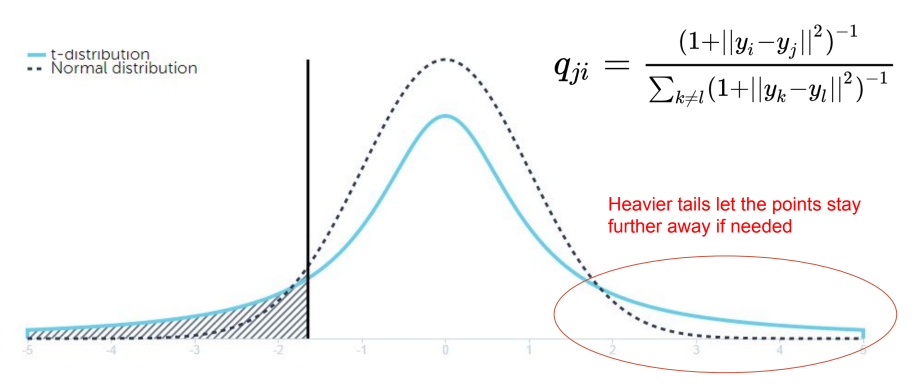
- "Crowding problem"
- In N dimension, you can have N+1 points at equal distance. But you cannot model this in smaller dimension



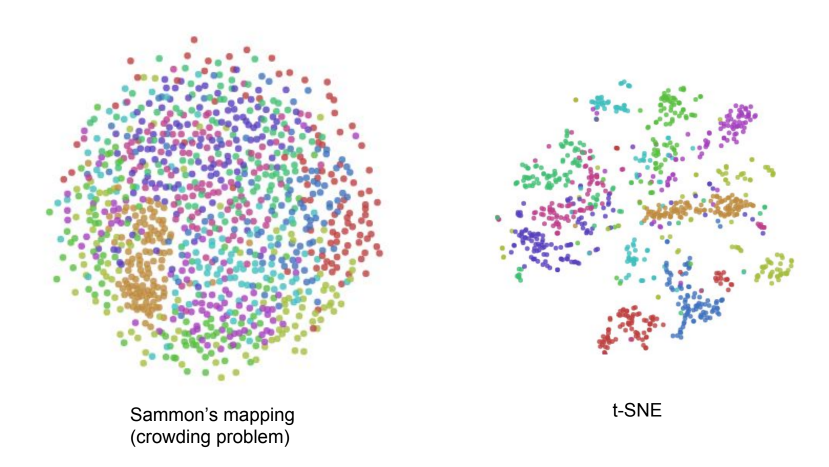
#### t-distributed

Instead of Gaussian for q we use student's t distribution

$$|q_{ji} \propto (1 + ||y_i - y_j||^2)^{-1}$$



## Crowding and t-SNE



## Variance

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

How to set the variance of our original space? A single variance for all points is not ideal.

- Want small variance for dense parts
- Want big variance for sparse parts

Set variance by amount of neighbors you want! How to quantify amount of neighbors?

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

$$Perp(P_i) = 2^{H(P_i)} \ H(P_i) = -\sum_j p_{j|i} log_2 p_{j|i}$$
 Entropy

Perplexity of P<sub>i</sub> represents effective amount of neighbors for the point i

Set Perp(P<sub>i</sub>) then t-SNE algorithm searches for the corresponding variance

Typical values for perplexity 5 to 50

## t-SNE summary

Goal: preserves local neighbors

Gradient-based -> need multiple runs to see the best

Two parameters: #iteration, perplexity

#### Does not learn a projection (unlike PCA, LDA, RP)

- If you have a new sample, you have to re-run the whole thing
- Might use a non-linear model to create learn t-SNE embeddings (deep learning)

## Summary

- PCA
- LDA
  - PCA+LDA
- Random projection
- tSNE
- Homework

Next time SVM