# STAT 215A Fall 2020 Week 12

James Duncan, OH: M, Th 2-4pm

Thanks to Tiffany Tang and past GSIs for sharing their slides

#### **Announcements**

- Lab 4 due in less than one week on November 19 at 11:59pm
  - Everyone in the group submits the **same** lab4 report / files but each person needs to push the files to their individual private repos
  - You may save results, but make sure it's clear what code generated them
- Next Friday's plan:
  - Guest talk: Tiffany Tang will introduce the Yu Group's COVID-19 data curation efforts and discuss the data itself
  - This is going to be super helpful for the **final lab** which I will introduce after Tiffany's talk

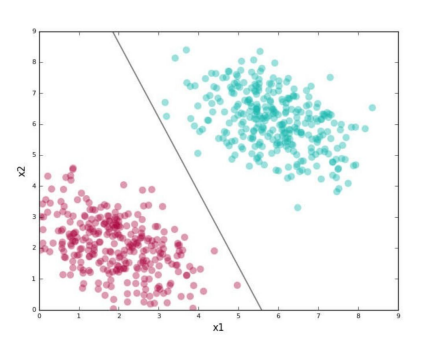
### Outline for today

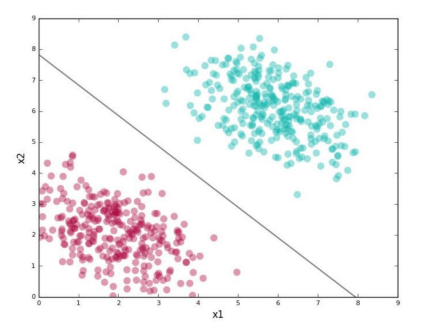
- More classification algorithms
  - SVM
  - Random forest
  - Ensembles
- Evaluation of classification performance

### Support vector machines

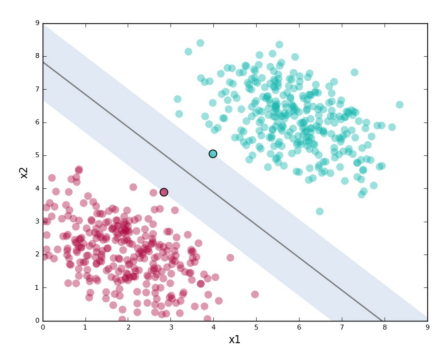
- Intuition: <a href="https://blog.statsbot.co/support-vector-machines-tutorial-c1618e635e93">https://blog.statsbot.co/support-vector-machines-tutorial-c1618e635e93</a>
- More in-depth discussion of the math:
  - https://towardsdatascience.com/understanding-support-vector-machine-part
     -1-lagrange-multipliers-5c24a52ffc5e
  - https://towardsdatascience.com/understanding-support-vector-machine-part
     -2-kernel-trick-mercers-theorem-e1e6848c6c4d
- Elements of Statistical Learning
  - Section 4.5 and Chapter 12
- We'll focus on intuition

#### Which is better?

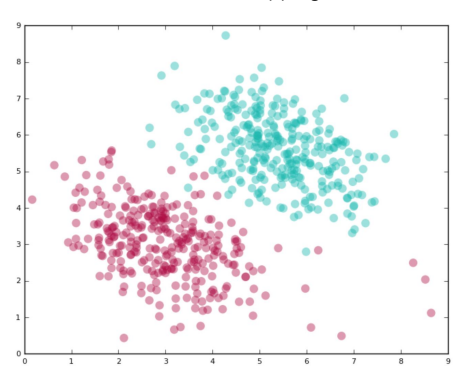




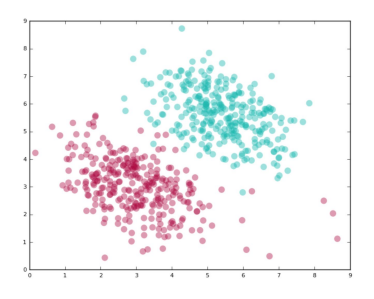
**An idea**: maximize space between two hyperplanes that separate the classes "Maximum margin" classifier

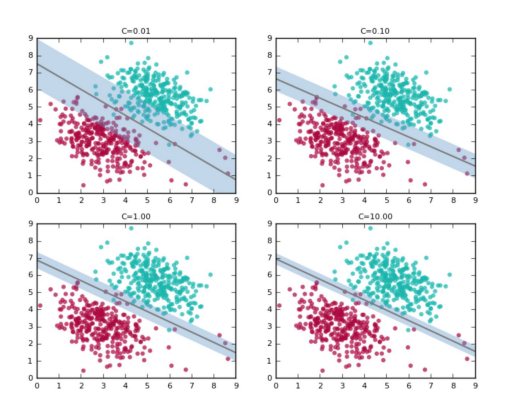


What about when the two classes are overlapping?

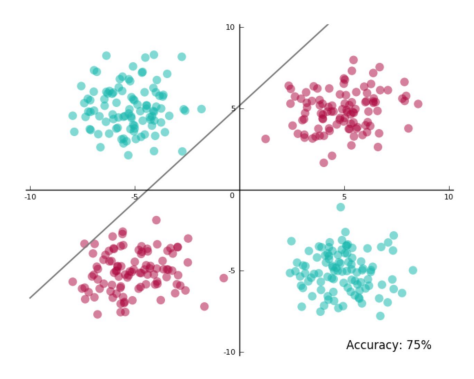


Another idea: Allow for some "slack"





What if there is no good separating hyperplane?

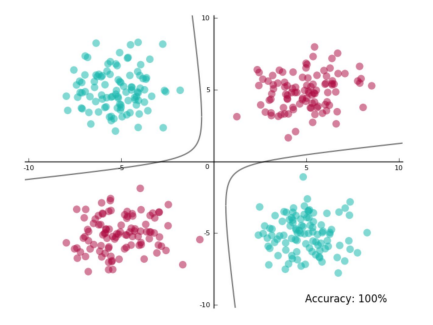


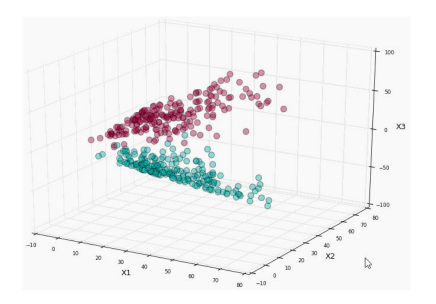
**Idea**: Find a higher-dimensional representation of the data where it becomes linearly separable

$$X_1 = x_1^2$$

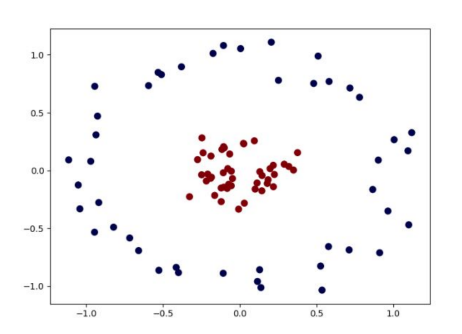
$$X_2 = x_2^2$$

$$X_3 = \sqrt{2}x_1x_2$$

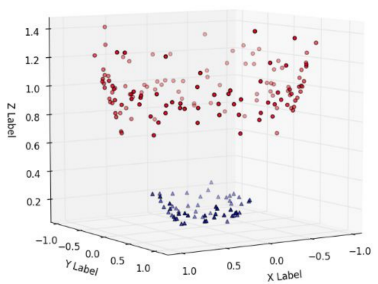




Another example of a higher dimensional representation that is linearly separable



Data in R^3 (separable)



So how do we perform this "lifting" to higher dimensions trick in a computationally feasible way? The answer: the **kernel trick**.

 Can show that by maximizing the margins while allows for slack, SVM solves the following optimization problem:

Inner product between two of the data points 
$$L_D = \sum_{i=1}^N \alpha_i - \frac{1}{2} \sum_{i=1}^N \sum_{k=1}^N \alpha_i \alpha_k y_i y_k x_i^T x_k$$
 subject to  $\alpha_i \geq 0$  and  $\sum_{i=1}^N \alpha_i y_i = 0$ 

Why not replace the usual inner product  $x_i^{ op} x_k$  with

$$\varphi(x_i)^{\top}\varphi(x_k)$$

where  $\varphi$  is some map from  $\mathbb{R}^p$  to a higher-dimensional space (possible even infinite dimensional).

- ullet The trick: don't need to know what arphi actually is.
  - Good news: we don't have to compute an infinite dimensional map.
- Instead, we find the kernel function:

$$K(x_i, x_k) = \varphi(x_i)^{\top} \varphi(x_k)$$

#### Some common kernel functions:

- Linear kernel:  $K(x_i, x_k) = x_i^{\top} x_k$
- Naive polynomial kernel:  $K(x_i,x_k)=(x_i^{ op}x_k)^d$
- Polynomial kernel:  $K(x_i, x_k) = (1 + x_i^{\top} x_k)^d$
- Gaussian kernel:  $K(x_i,x_k) = \exp\left\{-\frac{1}{2}\|x_i-x_k\|_2^2\right\}$
- Radial basis kernel:  $K(x_i, x_k) = \exp\left\{-\gamma \|x_i x_k\|_2^2\right\}$
- Sigmoid kernel:  $K(x_i, x_k) = \tanh(\eta x_i^{\top} x_j + \nu)$

An example: polynomial kernels for 2-dimensional data

$$k(\mathbf{x}, \mathbf{y}) = (1 + \mathbf{x}^T \mathbf{y})^2 = (1 + x_1 y_1 + x_2 y_2)^2 =$$

$$= 1 + x_1^2 y_1^2 + x_2^2 y_2^2 + 2x_1 y_1 + 2x_2 y_2 + 2x_1 x_2 y_1 y_2$$

This is an inner product between two 6-dimensional vectors:

$$\varphi(x) = \varphi(x_1, x_2) = (1, x_1^2, x_2^2, \sqrt{2}x_1, \sqrt{2}x_2, \sqrt{2}x_1x_2)$$

$$\varphi(y) = \varphi(y_1, y_2) = (1, y_1^2, y_2^2, \sqrt{2}y_1, \sqrt{2}y_2, \sqrt{2}y_1y_2)$$

$$\Rightarrow K(x, y) = \varphi(x)^T \varphi(y)$$

What happens if we use the naive polynomial kernel?  $K(x,y) = (x^Ty)^2$ 

Another example: write the Gaussian kernel as an inner product.

$$K(x,z) = e^{-\frac{1}{2\sigma^2}(x-z)^2} = e^{-\frac{x^2+z^2}{2\sigma^2}} e^{\frac{xz}{\sigma^2}}$$

$$= e^{-\frac{x^2+z^2}{2\sigma^2}} \left( \sum_{n=0}^{\infty} \frac{(xz)^n}{\sigma^{2n} n!} \right)$$

$$= e^{-\frac{x^2+z^2}{2\sigma^2}} \left( \sum_{n=0}^{\infty} \sqrt{\frac{1}{\sigma^{2n} n!}} x^n \cdot \sqrt{\frac{1}{\sigma^{2n} n!}} z^n \right)$$

$$= e^{-\frac{x^2}{2\sigma^2}} e^{-\frac{z^2}{2\sigma^2}} \left[ 1 \cdot 1 + \sqrt{\frac{1}{\sigma^2 1!}} x \cdot \sqrt{\frac{1}{\sigma^2 1!}} z + \sqrt{\frac{1}{\sigma^4 2!}} x^2 \cdot \sqrt{\frac{1}{\sigma^4 2!}} z^2 + \dots \right]$$

$$= \phi(x)^{\top} \phi(z)$$

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$$= \phi(x)^{\top} \phi(z)$$

**Takeaway:** by replacing the usual inner product with the Gaussian kernel it's as if we're projecting the data into an infinite dimensional space and finding a separating hyperplane there.

### Recap of SVMs + kernel trick

- Idea: find a separating hyperplane that maximizes margins (with some slack) between classes
- This becomes an optimization problem:

$$L_D = \sum_{i=1}^{N} \alpha_i - \frac{1}{2} \sum_{i=1}^{N} \sum_{k=1}^{N} \alpha_i \alpha_k y_i y_k K(x_i, x_k)$$
subject to  $\alpha_i \ge 0$  and  $\sum_{i=1}^{N} \alpha_i y_i = 0$ 

The optimization depends on the data only through the inner product, can
use the kernel trick to "lift" the data into a higher-dimensional space which
hopefully helps us to find a separating hyperplane

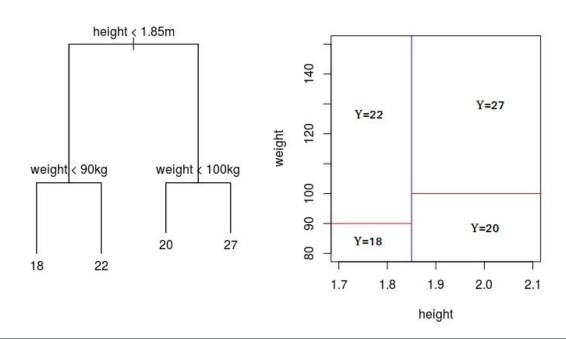
### SVM in practice

- Kernel trick allows for extreme flexibility
- However, with this greater flexibility comes a greater danger of overfitting,
   especially if p is large
- Lots of other methods based upon this kernel trick:
  - Kernel PCA
  - Kernel ridge regression
  - Spectral clustering
  - o etc.

# Trees and random forests



**Idea**: recursively partition the data space via binary splits and fit a simple model for each result region



- At each split in the tree, how to choose what variable (j) to split on and what threshold (t)?
- For now, assume we have a regression problem: at each split, we want to optimize  $L^2$  loss

$$\min_{j,t} \left\{ \min_{\mu_L} \sum_{i:x_{ij} \le t} (y_i - \mu_L)^2 + \min_{\mu_R} \sum_{i:x_{ij} > t} (y_i - \mu_R)^2 \right\}$$

 Can find global minimum for each split via a brute force search, but not necessarily for the entire tree

Brute force search algorithm:

- For each feature j:
  - Sort X:  $x_{1j} \leq \ldots \leq x_{nj} \Rightarrow O(n \log n)$
  - Scan from left to right and threshold  $t_j$  that minimizes  $L^2$  loss => O(n)
- Out of the p possible splits, take the best  $t_{j}$

Total complexity:  $O(pn \log n + pnK)$  where K is the number of splits

• For classification, can replace  $L^2$  loss with classification error, Gini index, etc.

#### **Advantages:**

- Can deal with continuous, categorical, binary, count features all at the same time
- Doesn't depend on scale of X
- Easily interpretable, fairly flexible, and fast

#### **Disadvantages:**

- Potentially too simple
- Not a great balance between bias-variance tradeoff
  - As depth of tree increases, overfits to the training data, resulting in high variance and no bias
  - If tree is too shallow, underfits and we have the opposite problem

#### Random forest

**Idea**: try to reduce both the bias and variance using decision trees

- To reduce the bias, grow deep trees (i.e., grow trees to purity so that in each of the leaf nodes, we have 1-3 observations left)
- To reduce the variance:
  - Grow many trees (e.g., 500 trees) using bootstrap samples of the data and average over this "forest".
  - $\circ$  Try to force these trees to be close to i.i.d.: at each split, select  $m_{\rm try}$  out of p variables randomly to search and potentially split on.

### Random forest algorithm

**Inputs**: number of trees to grow (B), number of variables to randomly select a each split ( $m_{\rm trv}$ ), number of leaf/terminal nodes (M)

For each tree, b = 1, ..., B:

- Bootstrap data:  $\mathbf{X}^{*b}$
- Grow decision (CART) tree T<sup>b</sup> such that:
  - $\circ$  At each split in the tree, randomly choose  $m_{\rm try}$  out of p variables to try and potentially split on
  - Grow until tree has M leaf / terminal nodes
- Make prediction:  $\hat{y}(x) = \frac{1}{B} \sum_{b=1}^{B} T^b(x)$

#### Random Forest in Practice

- Because we are bootstrapping the data before constructing each tree, we essentially have a "test set" for each tree that we can exploit
  - We call this left out data due to bootstrapping the out-of-bag (OOB)
     data, from which we can compute the OOB error
  - $\circ$   $\,$  OOB error can be used like CV error to tune parameters like  $m_{\rm try}$
- Can obtain marginal feature importances from RF

#### Random Forest in Practice

#### **Advantages:**

- Doesn't depend on scale of X
- Great prediction for lots of problems
- Reduces bias and variance simultaneously unlike CART

#### **Disadvantages:**

- May not be optimal with correlated features or p >> n
- No longer easily interpretable

In R: randomForest and ranger

ranger is much faster

#### Evaluation metrics for classification

How to evaluate your classification methods?

- Going beyond classification error
- What if we have class imbalance?
  - For example, if we take a sample of 100 people and only 10 have the disease, then always predicting healthy gives 90% classification accuracy!
  - We can do better.

### Confusion matrix

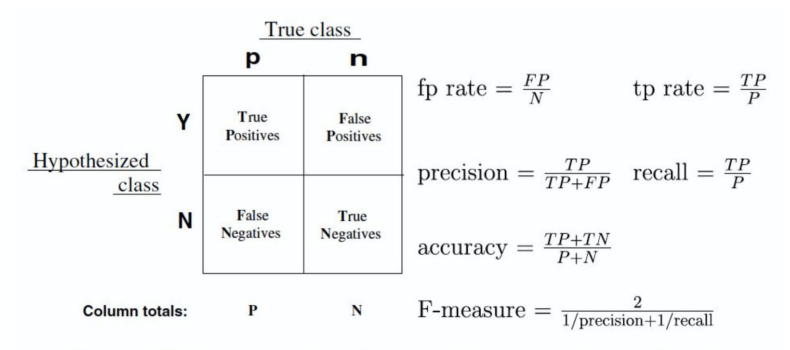


Fig. 1. Confusion matrix and common performance metrics calculated from it.

#### Confusion matrix

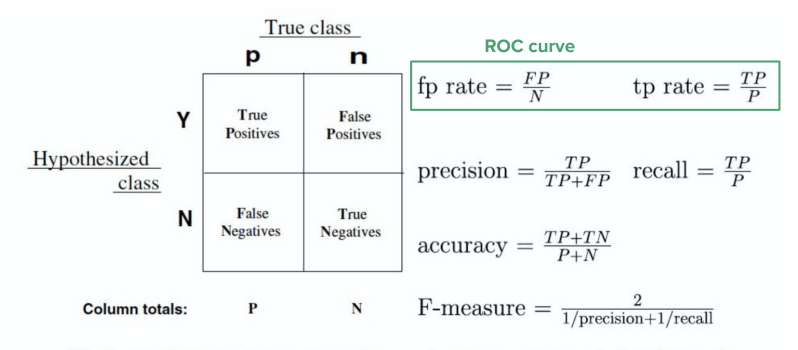


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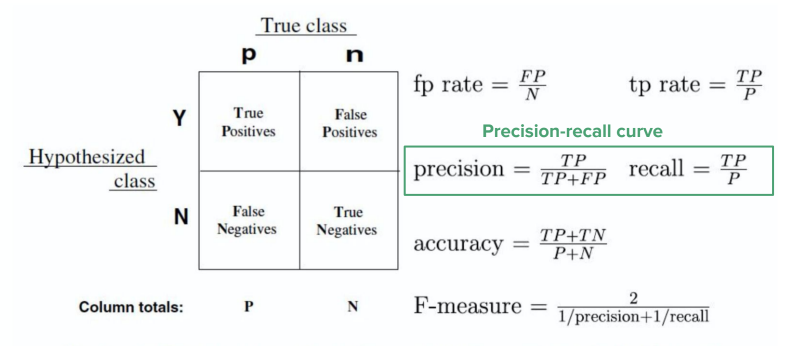
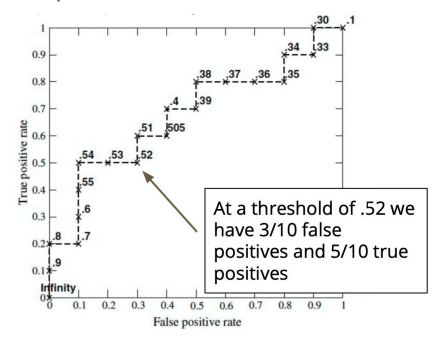


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### Receiver operating characteristics (ROC) curve

We can generate an ROC curve when the output of a classifier is a probability and we must choose a threshold for the final predicted class

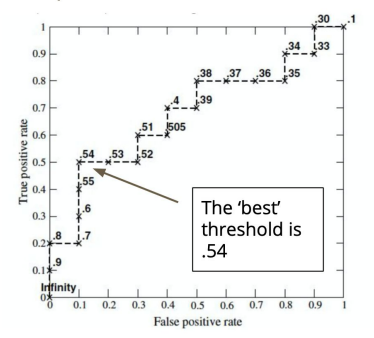
Inst#	Class	Score	Inst#	Class	Score
1	p	.9	11	p	.4
2	p	.8	12	n	.39
3	n	.7	13	p	.38
4	p	.6	14	n	.37
5	p	.55	15	n	.36
6	p	.54	16	n	.35
7	n	.53	17	p	.34
8	n	.52	18	n	.33
9	p	.51	19	p	.30
10	n	.505	20	n	.1



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#### Area under the curve

The area under the curve (AUC) is a method for comparing algorithms and evaluating classifiers.

The AUC has an important statistical property:

The AUC of a classifier is equivalent to the probability that the classifier will rank a randomly chosen positive instance higher than a randomly chosen negative instance

#### Area under the curve

Care should be taken when using ROC curves to compare classifiers

- The ROC graph is often used to select the best classifiers simply by graphing them in ROC space and seeing which one dominates.
- ☐ This is misleading: it is analogous to taking the maximum of a set of accuracy figures from a single test set.
- ☐ Without a measure of **variance** we cannot compare classifiers

It is a good idea to the average of multiple ROC curves (e.g. via cross validation)

See Fawcett (2005) for examples on how to average

### ROC vs Precision-Recall (PR) Curves

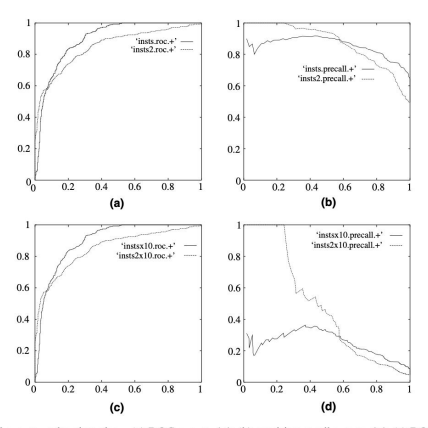


Fig. 5. ROC and precision-recall curves under class skew. (a) ROC curves, 1:1; (b) precision-recall curves, 1:1; (c) ROC curves, 1:10 and (d) precision-recall curves, 1:10.

#### ROC vs PR curves

- Generally, precision-recall curves are preferred when there is class imbalance
- ROC curves tend to paint an overly optimistic view of the model on datasets with class imbalance
- PR calculations do not involve the true negatives rate and hence do not typically present such an optimistic view

