## CSE/STAT 416

Precision/Recall k-Nearest Neighbors

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May 3, 2021



Pre-Lecture Video

## Detecting Spam

Imagine I made a "Dummy Classifier" for detecting spam

- The classifier ignores the input, and always predicts spam.
- This actually results in 90% accuracy! Why?
  - Most emails are spam...

This is called the majority class classifier.

A classifier as simple as the majority class classifier can have a high accuracy if there is a **class imbalance**.

A class imbalance is when one class appears much more frequently than another in the dataset

This might suggest that accuracy isn't enough to tell us if a model is a good model.



## Assessing Accuracy

Always digging in and ask critical questions of your accuracy.

- Is there a class imbalance?
- How does it compare to a baseline approach?
  - Random guessing
  - Majority class
  - ...
- Most important: What does my application need?
  - What's good enough for user experience?
  - What is the impact of a mistake we make?



## Confusion Matrix

For binary classification, there are only two types of mistakes

$$\hat{y} = +1, y = -1$$

$$\hat{y} = -1, y = +1$$

Generally we make a confusion matrix to understand mistakes.

#### **Predicted Label**

		4	
True Label	<b>♣</b>	True Positive (TP)	False Negative (FN)
		False Positive (FP)	True Negative (TN)



## Binary Classification Measures

#### Notation

$$C_{TP} = \text{\#TP}, \quad C_{FP} = \text{\#FP}, \quad C_{TN} = \text{\#TN}, \quad C_{FN} = \text{\#FN}$$

$$N = C_{TP} + C_{FP} + C_{TN} + C_{FN}$$

$$N_P = C_{TP} + C_{FP}, \quad N_N = C_{FP} + C_{TN}$$

#### **Error Rate**

$$\frac{C_{FP} + C_{FN}}{N}$$

#### **Accuracy Rate**

$$\frac{C_{TP} + C_{TN}}{N}$$

#### False Positive rate (FPR)

$$\frac{C_{FP}}{N_N}$$

#### False Negative Rate (FNR)

$$\frac{C_{FN}}{N_P}$$

#### True Positive Rate or Recall

$$\frac{T_P}{N_P}$$

#### Precision

$$\frac{T_P}{C_{TP} + C_{FP}}$$

#### F1-Score

$$2\frac{Precision \cdot Recall}{Precision + Recall}$$

#### See more!





## Change Threshold

What if I never want to make a false positive prediction?

What if I never want to make a false negative prediction?

One way to control for our application is to change the scoring threshold. (Could also change intercept!)

- If  $Score(x) > \alpha$ :
  - Predict  $\hat{y} = +1$
- Else:
  - Predict  $\hat{y} = -1$



## ROC Curve

What happens to our TPR and FPR as we increase the threshold?





# Assessing Accuracy

Often with binary classification, we treat the positive label as being the more important of the two. We then often then focus on these metrics:

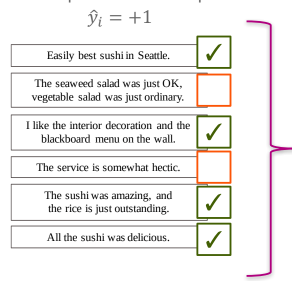
**Precision:** Of the ones I predicted positive, how many of them were actually positive?

**Recall:** Of all the things that are truly positive, how many of them did I correctly predict as positive?



### Precision

What fraction of the examples I predicted positive were correct? Sentences predicted to be positive:



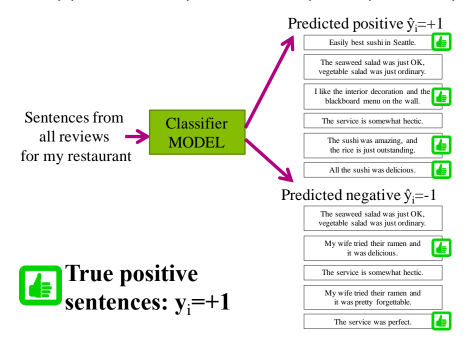
Only 4 out of 6
sentences
predicted to be
positive are
actually positive

$$precision = \frac{C_{TP}}{C_{TP} + C_{FP}}$$



### Recall

Of the truly positive examples, how many were predicted positive?



$$recall = \frac{C_{TP}}{N_P} = \frac{C_{TP}}{C_{TP} + C_{FN}}$$

# Precision & Recall

An optimistic model will predict almost everything as positive

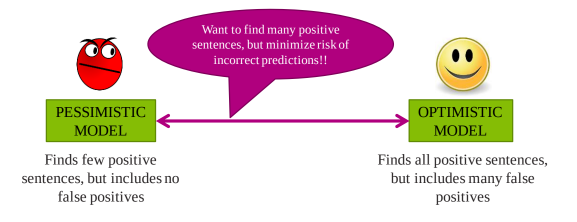


High recall, low precision

A pessimistic model will predict almost everything as negative



High precision, low recall



## Controlling Precision/Recall

Depending on your application, precision or recall might be more important

Ideally you will have high values for both, but generally increasing recall will decrease precision and vice versa.

For logistic regression, we can control for how optimistic the model is by changing the threshold for positive classification

#### **Before**

$$\hat{y}_i = +1 \text{ if } \hat{P}(y = +1|x_i) > 0.5 \text{ else } \hat{y}_i = -1$$

#### Now

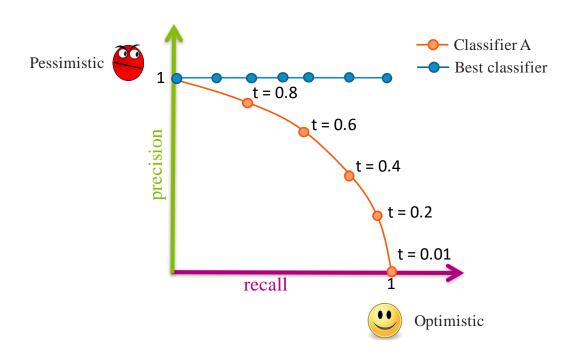
$$\hat{y}_i = +1$$
 if  $\hat{P}(y = +1|x_i) > t$  else  $\hat{y}_i = -1$ 





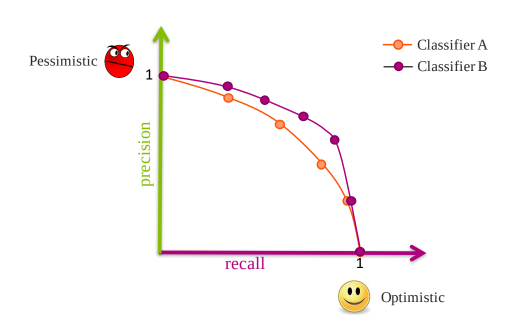


Can try every threshold to get a curve like below





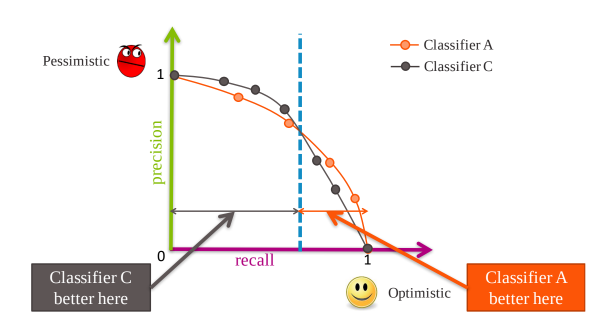
#### Sometimes, Classifier B is strictly better than Classifier A







#### Most times, the classifiers are incomparable





## Compare Classifiers

Often come up with a single number to describe it

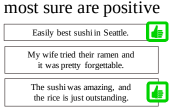
- F1-score, AUC, etc.
- Remember, what your application needs is most important

#### Also common to use precision at k

If you show the top **k** most likely positive examples, how many of them are true positives

Showing k=5 sentences on website





All the sushi was delicious.

The service was perfect.

Sentences model



precision at k = 0.8



## Class Session

## Roadmap

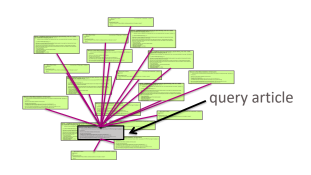
- 1. Housing Prices Regression
  - Regression Model
  - Assessing Performance
  - Ridge Regression
  - LASSO
- 2. Sentiment Analysis Classification
  - Classification Overview
  - Logistic Regression
  - Bias / Fairness
  - Decision Trees
  - Ensemble Methods
- 3. Document Retrieval Clustering and Similarity
  - Precision / Recall
  - k-Nearest Neighbor
  - Kernel Methods
  - Locality Sensitive Hashing
  - Clustering
  - Hierarchical Clustering



### Document Retrieval

- Consider you had some time to read a book and wanted to find other books similar to that one.
- If we wanted to write an system to recommend books
  - How do we measure similarity?
  - How do we search over books?
  - How do we measure accuracy?

Big Idea: Define an **embedding** and a **similarity metric** for the books, and find the **"nearest neighbor"** to some query book.





Nearest Neighbors

## 1-Nearest Neighbor

#### Input

- $x_q$ : Query example (e.g. my book)
- $x_1, ..., x_n$ : Corpus of documents (e.g. Amazon books)

#### Output

The document in corpus that is most similar to  $x_q$ 

$$x^{NN} = \underset{x_i \in [x_1, \dots, x_n]}{\operatorname{arg \, min}} \, distance(x_q, x_i)$$

It's very critical to properly define how we represent each document  $x_i$  and the similarity metric distance! Different definitions will lead to very different results.



## 1-Nearest Neighbor

How long does it take to find the 1-NN? About n operations

```
Input: x_q
x^{NN} = \emptyset
nn_dist = \infty
for x_i \in [x_1, ..., x_n]:
   dist = distance(x_q, x_i)
   if dist < nn_dist:
       x^{NN} = x_i
       nn_dist = dist
Output: x^{NN}
```

## k-Nearest Neighbors

#### Input

- $x_q$ : Query example (e.g. my book)
- $x_1, ..., x_n$ : Corpus of documents (e.g. Amazon books)

#### Output

List of k documents most similar to  $x_q$ 

Formally



## k-Nearest Neighbors

Same idea as 1-NN algorithm, but maintain list of k-NN

```
Input: x_a
X^{k-NN} = [x_1, \dots, x_k]
nn\_dists = [dist(x_1, x_q), dist(x_2, x_q), ..., dist(x_k, x_q)]
for x_i \in [x_{k+1}, ..., x_n]:
   dist = distance(x_q, x_i)
   if dist < max(nn\_dists):
        remove largest dist from X^{k-NN} and nn_dists
       add x_i to X^{k-NN} and distance (x_q, x_i) to nn_dists
Output: X^{k-NN}
```

## k-Nearest Neighbors

Can be used in many circumstances!

Retrieval

Return  $X^{k-NN}$ 

Regression

$$\hat{y}_i = \frac{1}{k} \sum_{j=1}^k x^{NNj}$$

Classification

$$\hat{y}_i = majority\_class(X^{k-NN})$$





## Important Points

While the formalization of these algorithms are fairly tedious, the intuition is fairly simple. Find the 1 or k nearest neighbors to a given document and return those as the answer.

This intuition relies on answering two important questions

- How do we represent the documents  $x_i$ ?
- How do we measure the distance  $distance(x_a, x_i)$ ?



### Document Representation

Like our previous ML algorithms, we will want to make a vector out of the document to represent it as a point in space.

Simplest representation is the **bag-of-words** representation.

- Each document will become a *W* dimension vector where *W* is the number of words in the entire corpus of documents
- The value of  $x_i[j]$  will be the number of times word j appears in document i.
- This ignores order of words in the document, just the counts.



### Bag of Words

#### Pros

- Very simple to describe
- Very simple to compute

#### Cons

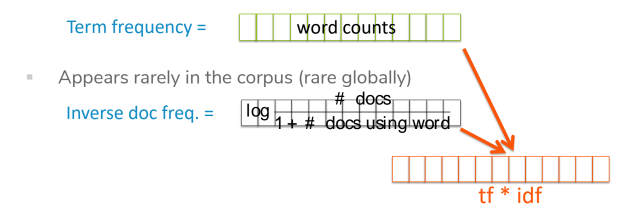
- Common words like "the" and "a" dominate counts of uncommon words
- Often it's the uncommon words that uniquely define a doc.



### TF-IDF

#### **Goal**: Emphasize important words

Appear frequently in the document (common locally)



Do a pair-wise multiplication to compute the TF-IDF for each word

Words that appear in every document will have a small IDF making the TF-IDF small!



Distance

# Euclidian Distance

Now we will define what similarity/distance means

Want to define how "close" two vectors are. A smaller value for distance means they are closer, a large value for distance means they are farther away.

The simplest way to define distance between vectors is the **Euclidean distance** 

$$distance(x_i, x_q) = ||x_i - x_q||_2$$
$$= \sqrt{\sum_{j=1}^{D} (x_i[j] - x_q[j])^2}$$



## Manhattan Distance

#### Another common choice of distance is the Manhattan Distance

$$distance(x_i, x_q) = ||x_i - x_q||_1$$
$$= \sum_{j=1}^{D} |x_i[j] - x_q[j]|$$



# Weighted Distances

Some features vary more than others or are measured in different units. We can weight different dimensions differently to make the distance metric more reasonable.



#### Weighted Educlidean distance

$$distance(x_i, x_q) = \sqrt{\sum_{j=1}^{D} a_j^2(x_i[j] - x_q[j])^2}$$



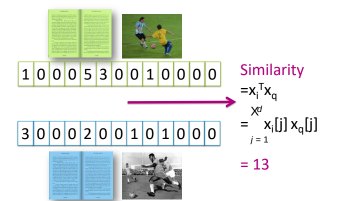
## Similarity

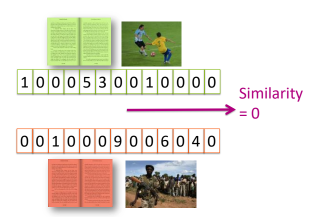
Another natural similarity measure would use

$$x_i^T x_q = \sum_{j=1}^D x_i[j] x_q[j]$$

Notice this is a measure of similarity, not distance

This means a bigger number is better





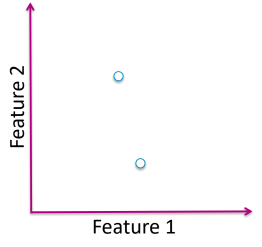
## Cosine Similarity

Should we normalize the vectors before finding the similarity?

$$similarity = \frac{x_i^T x_q}{\left| |x_i| \right|_2 \left| |x_q| \right|_2} = \cos(\theta)$$

#### Note:

- Not a true distance metric
- Efficient for sparse vectors!





## Cosine Similarity

In general

 $-1 \le cosine \ similarity \le 1$ 

For positive features (like TF-IDF)

 $0 \le cosine \ similarity \le 1$ 

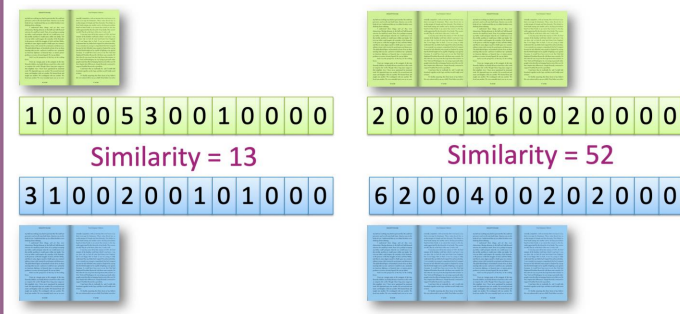
Define

distance = 1 - similarity



## To Normalize or Not To Normalize?

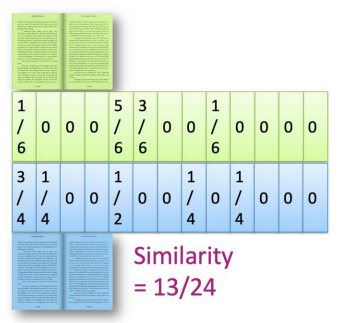
#### Not normalized

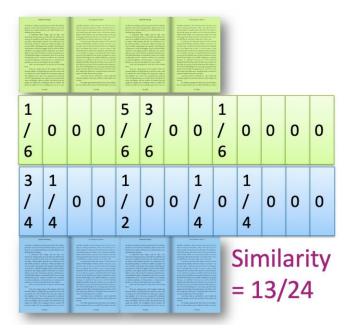




## To Normalize or Not To Normalize?

#### Normalized







## To Normalize or Not To Normalize?

Normalization is not desired when comparing documents of different sizes since it ignores length.





Normalizing can make dissimilar objects appear more similar





Common compromise: Just cap maximum word counts



long document

In practice, can use multiple distance metrics and combine them using some defined weights





Think &

3 min



For the given documents, what are their Euclidean Distance and Cosine Similarity?

Assume we are using a bag of words representation

Document 1: "I really like dogs"

Document 2: "dogs are really really awesome"

#### Steps:

- Write out bag of words vectors
- Compute Euclidean distance
- Compute Cosine similarity



## Poll Everywhere

Think &



### Recap

**Theme**: Use nearest neighbors to recommend documents.

#### Ideas:

- Precision and Recall Curves
- Implement a nearest neighbor algorithm
- Compare and contrast different document representations
  - Emphasize important words with TF-IDF
- Compare and contrast different measurements of similarity
  - Euclidean and weighted Euclidean
  - Cosine similarity and inner-product similarity

