Introduction to locality sensitive hashing (LSH)

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STA 325, Supplemental Material

October 11, 2018

Mid-term Evaluations

Thank you for completing the midterm evaluations!

Things that students liked:

- Real examples
- 2 The fact that the lectures are all reproducible
- 3 Interactive parts of the lectures
- 4 Course materials, homeworks, labs
- **5** Pace of the course

12/13 students very much liked the course, content, material (4-5 range overall).

Things you would like to improve:

- Posted readings for course material
- More real examples
- 3 The homeworks to be more clear in the wording
- 4 More interactive lectures



Homework Reminders

- Please remember to include all files when you submit (otherwise they will not be graded). A big part of the course is reproducibility.
- 2 Please comment your code. (There will be a 10 point deduction for code that is not commented. Please also make sure that your comments are clear).

Questions?

Agenda

- Defining similarity
- 2 Representing data as sets (shingling)
- 3 Hashing
- 4 Hashing with compression (minhashing)
- 5 Too many pairs to compare! (LSH)
- 6 Evaluation
- Teven faster?

Reading

The reading for this module can be found in Mining Massive Datasets, Chapter 3.

http://infolab.stanford.edu/~ullman/mmds/book.pdf

There are no applied examples in the book, however, we will be covering these in class and homework.

Motivations

In information retrieval, recall that one main goal is understanding how similar items are in our task (application) at hand.

Documents

Suppose that we have a large number of documents (articles, songs, etc). We may wish to know which documents are the most similar.

Entity resolution

Entity resolution (record linkage or de-duplication) is the process of removing duplicate information from large noisy databases.

Often times we perform entity resolution and then perform a regression task (or some other inference/prediction task).

Information retrieval for comparing documents, records, etc

- 1 If we're looking to remove duplicate entities from a database, this requires comparing all *n* records in the database. (This is a quadratic operation.)
- 2 Typically, before performing entity resolution, we want to perform dimension reduction to reduce down the space of records to something more manageable.

(Suppose our database has 1000 records, then then without dimension reduction we're having to do $(1000)^2$ comparisons). Now think larger and more realistic (1 million records).

For further reading on how to do this in practice, see Chen, Shrivastava, Steorts (2018), Steorts et al. (2018), Steorts et al. (2014).

Overall questions

- How can we use take similar data points (records) and put them in clusters (buckets or bins)?
- How can we evaluate how well our method is doing? (Look at the precision and recall).

Blocking (partitioning)

Blocking partitions of data points so that we do not have to make all-to-all data point comparisons.

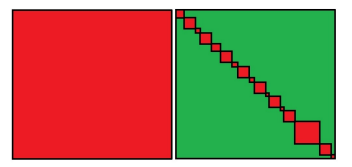


Figure: All-to-all data point comparisons (left) versus partitioning data points into blocks/bins (right).

Entity resolution

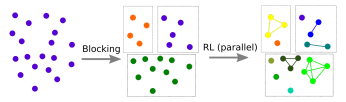


Figure: Entire process of the blocking step (via LSH) and then the record linkage step. Step 1: Dimension reduction via blocking. Step 2: Removing duplicate entities via record linkage.

Remark: In this module, we will focus on how to perform dimension reduction methods using LSH. For reading on how to perform the record linkage step, see Christen (2012).

Goal

Our overall goal in this lecture will be able to quickly compare the similarity of the following:

- documents
- songs
- data points
- other examples

We will work with two data sets. The first is a synthetic dataset from the RecordLinkage package in R called RLdata500. The second is a real data set on github (RLdata) called cora¹

¹This can be found at https://github.com/resteorts/RLdata. ← ≥ ▶

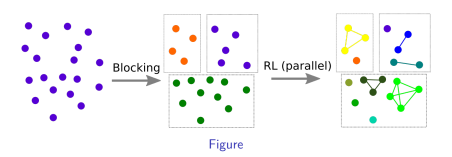
Install instructions

To install the RLdata package use the following commands:

```
install.packages("devtools")
devtools::install_github("resteorts/RLdata")
```

Goal

Goal: Introduce locality sensitive hashing, a fast method of blocking for record linkage, and get some experience doing LSH in R



Finding similar items

- We want to find similar items
 - Maybe we are looking for near duplicate documents (plagiarism)
 - More likely, we are trying to block our data which we can later pass to a record linkage process
- How do we define similar?

Jaccard similarity

There are many ways to define similarity, we will use *Jaccard* similarity for this task

$$Jac(S,T) = \frac{\mid S \cap T \mid}{\mid S \cup T \mid}$$

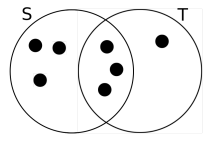


Figure: Two sets S and T with Jaccard similarity 3/7. The two sets share 3 elements in common, and there are 7 elements in total.

How to represent data as sets

We want to talk about similarity of data \Rightarrow we need sets to compare!

- One way is to construct from the data the set of short strings that appear within it
- Similar documents/datasets will have many common elements,
 i.e. many commong short strings
- We can do construct these short strings using shingling

k-shingling (how-to)

1 Think of a document or record as a string of characters

A k-shingle (k-gram) is any sub-string (word) of length k found within the document or record

3 Associate with each document or record the set of k-shingles that appear one or more times within it

Let's try

Suppose our document is the string "Hello world", then

• the set of 2-shingles is

```
{he, el, II, lo, ow, wo, or, rl, ld}
```

the set of 3-shingles is

```
{hel, ell, llo, low, owo, wor, orl, rld}
```

Your turn

We have the following two records:

	First name	Last name
129	MICHAEL	VOGEL
130	MICHAEL	MEYER

- 1 Compute the 2-shingles for each record
- Using Jaccard similarity, how similar are they?

Your turn solution

1. The 2-shingles for the first record are

and for the second are

2. There are 6 items in common

and 15 items total

 ${mi, ic, ch, ha, ae, el, lv, vo, og, ge, lm, me, ey, ye, er},$

so the Jaccard similarity is

$$\frac{6}{15} = \frac{2}{5} = 0.4$$



Useful packages/functions in R

It would be better to automate this. (And we can do this in R.)

```
library(textreuse) # text reuse/document similarity
library(tokenizers) # shingles
```

We can use the following functions to create k-shingles and calculate Jaccard similarity for our data

```
# get k-shingles
tokenize_character_shingles(x, n)

# calculate jaccard similarity for two sets
jaccard_similarity(a, b)
```

How to automate this?

```
# load packages
library(textreuse)
library(tokenizers)
# list strings
string1 <- "michael vogel"
string2 <- "michael meyer"
# create shingles
shingle1<- tokenize_character_shingles(string1, n=2)[[1]]
shingle2 <- tokenize_character_shingles(string2, n=2)[[1]]</pre>
# compute jaccard similarity
jaccard_similarity(shingle1,shingle2)
```

Cora data

Research paper headers and citations, with information on authors, title, institutions, venue, date, page numbers and several other fields

```
library(RLdata) # data library
data(cora) # load the cora data set
str(cora) # structure of cora
```

```
## 'data.frame': 1879 obs. of 16 variables:
## $ id
              : int 1 2 3 4 5 6 7 8 9 10 ...
## $ title
               :Class 'noquote' chr [1:1879] "Inganas and M.R" NA NA NA ...
## $ book title :Class 'noquote' chr [1:1879] NA NA NA NA ...
## $ authors :Class 'noquote' chr [1:1879] "M. Ahlskog, J. Paloheimo, H. Stubb, P. Dyrekley, M. Fahl
## $ address :Class 'noquote' chr [1:1879] NA NA NA NA ...
## $ date :Class 'noquote' chr [1:1879] "1994" "1994" "1994" "1994" ...
## $ year :Class 'noquote' chr [1:1879] NA NA NA NA ...
## $ editor :Class 'noquote' chr [1:1879] NA NA NA NA ...
## $ journal :Class 'noquote' chr [1:1879] "Andersson, J Appl. Phys." "JAppl. Phys." "J Appl. Phys."
## $ volume
              :Class 'noquote' chr [1:1879] "76" "76" "76" "76" ...
## $ pages
             :Class 'noquote' chr [1:1879] "893" "893" "893" "893" ...
## $ publisher :Class 'noquote' chr [1:1879] NA NA NA NA ...
## $ institution:Class 'noquote' chr [1:1879] NA NA NA NA ...
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## $ tech
              :Class 'noquote' chr [1:1879] NA NA NA NA ...
               :Class 'noquote' chr [1:1879] NA NA NA NA ...
## $ note
```

Your turn

Using the title, authors, and journal fields in the cora dataset,

1. Get the 3-shingles for each record (hint: use tokenize_character_shingles)

Obtain the Jaccard similarity between each pair of records (hint: use jaccard_similarity)

Your turn (solution)

```
# get only the columns we want
n <- nrow(cora) # number of records
dat <- data.frame(id = seq len(n)) # create id column
dat <- cbind(dat, cora[, c("title", "authors", "journal")]) # get columnds we want
# 1. paste the columns together and tokenize for each record
shingles <- apply(dat, 1, function(x) {
  # tokenize strings
 tokenize character shingles(paste(x[-1], collapse=" "), n = 3)[[1]]
1)
# 2. Jaccard similarity between pairs
jaccard <- expand.grid(record1 = seq_len(n), # empty holder for similarities
                       record2 = seq len(n)
# don't need to compare the same things twice
iaccard <- iaccard[iaccard$record1 < iaccard$record2.]</pre>
time <- Sys.time() # for timing comparison
iaccard$similarity <- apply(jaccard, 1, function(pair) {</pre>
 jaccard_similarity(shingles[[pair[1]]], shingles[[pair[2]]]) # qet jaccard for each pair
1)
time <- difftime(Sys.time(), time, units = "secs") # timing
```

This took took 164.95 seconds ≈ 2.75 minutes

Your turn (solution, cont'd)

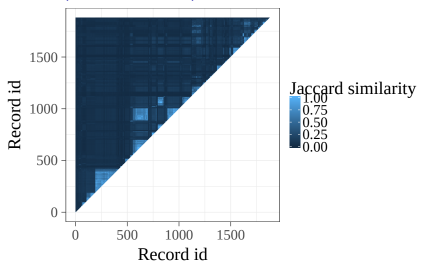


Figure: Jaccard similarity for each pair of records. Light blue indicates the two records are more similar and dark blue indicates less similar.

Hashing

For a dataset of size n, the number of comparisons we must compute is

$$\frac{n(n-1)}{2}$$

- For our set of records, we needed to compute 1,764,381 comparisons

- A better approach for datasets of any realistic size is to use hashing

Hash functions

- Traditionally, a hash function maps objects to integers such that similar objects are far apart
- Instead, we want special hash functions that do the opposite of this, i.e. similar objects are placed closed together!

Definition: Hash function

Hash functions h() are defined such that

If records A and B have high similarity, then the probability that h(A) = h(B) is **high** and if records A and B have low similarity, then the probability that $h(A) \neq h(B)$ is **high**.

Hashing shingles

Instead of storing the strings (shingles), we can just store the hashed values

These are integers, they will take less space

```
# instead store hash values (less memory)
hashed_shingles <- apply(dat, 1, function(x) {
   string <- paste(x[-1], collapse=" ") # get the string
   shingles <- tokenize_character_shingles(string, n = 3)[[1]] # 3-shing
   hash_string(shingles) # return hashed shingles
})</pre>
```

This took up 6.38256×10^5 bytes, while storing the shingles took 7.36544×10^6 bytes; the whole pairwise comparison still took the same amount of time (≈ 2.35 minutes)

Similarity preserving summaries of sets

- Sets of shingles are large (larger than the original document)
- If we have millions of documents, it may not be possible to store all the shingle-sets in memory
- We can replace large sets by smaller representations, called signatures
- And use these signatures to approximate the Jaccard similarity

Characteristic matrix

In order to get a signature of our data set, we first build a characteristic matrix

Columns correspond to records and the rows correspond to all hashed shingles

	Record 1	Record 2	Record 3	Record 4	Record 5
-78464425	1	1	1	1	1
-78234440	1	0	0	0	0
-78221717	1	0	0	0	0
-78235289	1	1	1	1	1
-78555255	1	1	1	1	1
-78132973	1	1	1	1	1

The result is a 3551×1879 matrix

Question: Why would we not store the data as a characteristic matrix?

Minhashing

Want create the signature matrix through minhashing

- \bigcirc Permute the rows of the characteristic matrix m times
- 2 Iterate over each column of the permuted matrix
- 3 Populate the signature matrix, row-wise, with the row index from the first 1 value found in the column

The signature matrix is a hashing of values from the permuted characteristic matrix and has one row for the number of permutations calculated (m), and a column for each record

Minhashing (cont'd)

Record 1	Record 2	Record 3	Record 4	Record 5
30	30	30	30	30
8	8	8	8	8
6	1	1	1	1
2	2	2	2	2
102	102	102	102	102
16	16	16	16	16
8	8	8	8	8
112	161	161	161	161
1	1	1	1	1
76	27	27	27	27

Signature matrix and Jaccard similarity

The relationship between the random permutations of the characteristic matrix and the Jaccard Similarity is

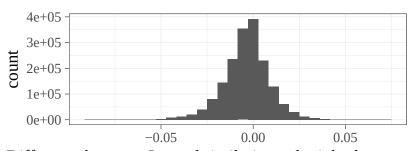
$$Pr\{\min[h(A)] = \min[h(B)]\} = \frac{|A \cap B|}{|A \cup B|}$$

We use this relationship to **approximate** the similarity between any two records

We look down each column of the signature matrix, and compare it to any other column

The number of agreements over the total number of combinations is an approximation to Jaccard measure

Jaccard similarity approximation



Difference between Jaccard similarity and minhash approx

Used minhashing to get an approximation to the Jaccard similarity, which helps by allowing us to store less data (hashing) and avoid storing sparse data (signature matrix)

We still haven't addressed the issue of pairwise comparisons

Minhashing (review of the entire process)

Let's take a look at the entire minhash process as a quick review.

Now, we permute the rows of the characteristic matrix to form a permuted matrix.

The permuted matrix is simply a reordering of the original characteristic matrix, with the rows swapped in some arrangement.

Figure 2 shows the characteristic matrix converted to a permuted matrix by a given permutation. We repeat the permutation step for several iterations to obtain multiple permuted matrices.

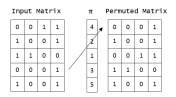


Figure: Permuted matrix from the characteristic one. The π vector is the specified permutation.

The Signature Matrix

Now, we compute the signature matrix.

The signature matrix is a hashing of values from the permuted one.

The signature has a row for the number of permutations calculated, and a column corresponding with the columns of the permuted matrix.

We iterate over each column of the permuted matrix, and populate the signature matrix, row-wise, with the row index from the first 1 value found in the column.

Signature Matrix

Avoiding pairwise comparisons

- Performing pairwise comparisons is time-consuming because the number of comparisons grows at $O(n^2)$
- Most of those comparisons are unnecessary because they do not result in matches due to sparsity
- We will use the combination of minhash and locality-sensitive hashing (LSH) to compute possible matches only once for each document, so that the cost of computation grows linearly

Locality Sensitive Hashing (LSH)

Idea: We want to hash items several times such that similar items are more likely to be hashed into the same bucket

- Divide signature matrix into b bands with r rows each so m = b * r where m is the number of times that we drew a permutation of the characteristic matrix in the process of minhashing
- 2 Each band is hashed to a bucket by comparing the minhash for those permutations
 - If they match within the band, then they will be hashed to the same bucket
- If two documents are hashed to the same bucket they will be considered candidate pairs

We only check candidate pairs for similarity

Banding and buckets

	Record 1	Record 2	Record 3	Record 4	Record 5
1	30	30	30	30	30
2	8	8	8	8	8
3	6	1	1	1	1
4	2	2	2	2	2
5	102	102	102	102	102
6	16	16	16	16	16
7	8	8	8	8	8
8	112	161	161	161	161
9	1	1	1	1	1
_10	76	27	27	27	27

Tuning

How to choose k

How large k should be depends on how long our data strings are

The important thing is k should be picked large enough such that the probability of any given shingle is low

How to choose b

b must divide m evenly such that there are the same number of rows r in each band

What else?

Choosing b

 $P(\text{two documents w}/\text{ Jaccard similarity } s \text{ marked as potential match}) = 1 - (1 - s^{m/b})^b$

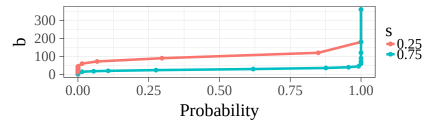


Figure: Probability that a pair of documents with a Jaccard similarity s will be marked as potential matches for various bin sizes b for s = .25, .75 for the number of permutations we did, m = 360.

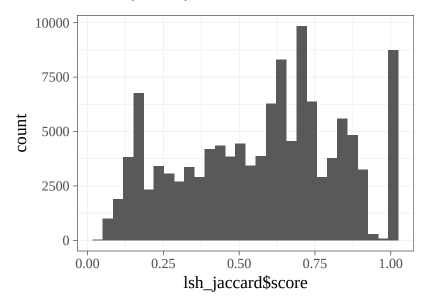
For b=90, a pair of records with Jaccard similarity .25 will have a 29.7% chance of being matched as candidates and a pair of records with Jaccard similarity .75 will have a 100.0% chance of being matched as candidates

"Easy" LSH in R

There an easy way to do LSH using the built in functions in the textreuse package via the functions minhash_generator and lsh (so we don't have to perform it by hand):

```
# choose appropriate num of bands
b <- 90
# create the minhash function
minhash <- minhash generator(n = m, seed = 02082018)
# build the corpus using textreuse
docs <- apply(dat, 1, function(x) paste(x[-1], collapse = " ")) # get strings</pre>
names(docs) <- dat$id # add id as names in vector
corpus <- TextReuseCorpus(text = docs, # dataset
                          tokenizer = tokenize character shingles, n = 3, simplify = TRUE, # shingles
                          progress = FALSE, # quietly
                          keep tokens = TRUE, # store shingles
                          minhash func = minhash) # use minhash
# perform lsh to get buckets
buckets <- lsh(corpus, bands = b, progress = FALSE)
# grab candidate pairs
candidates <- lsh candidates(buckets)
# get Jaccard similarities only for candidates
1sh jaccard <- 1sh compare(candidates, corpus, jaccard similarity, progress = FALSE)
```

"Easy" LSH in R (cont'd)



Putting it all together

The last thing we need is to go from candidate pairs to blocks

```
library(igraph) #graph package
# think of each record as a node
# there is an edge between nodes if they are candidates
g <- make_empty_graph(n, directed = FALSE) # empty graph
g <- add edges(g, as.vector(t(candidates[, 1:2]))) # candidate edges
g <- set_vertex_attr(g, "id", value = dat$id) # add id
# get custers, these are the blocks
clust <- components(g, "weak") # get clusters</pre>
blocks <- data.frame(id = V(g)$id, # record id
                     block = clust$membership) # block number
head(blocks)
```

```
## id block
## 1 1 1
## 2 2 1
## 3 3 1
## 4 4 1
## 5 5 1
```

Your turn

Using the fname_c1 and lname_c1 columns in the RecordLinkage::RL500 dataset,

- 1 Use LSH to get candidate pairs for the dataset
- What k to use for shingling?
- What b to use for bucket size?
- 2 Append the blocks to the original dataset as a new column, block

Even faster?

(fast): In minhashing we have to perform m permutations to create multiple hashes

(faster): We would like to reduce the number of hashes we need to create – "Densified" One Permutation Hashing (DOPH)

- One permutation of the signature matrix is used
- The feature space is then binned into m evenly spaced bins
- The m minimums (for each bin separately) are the m different hash values

See Chen, Shrivastava, Steorts (2018) for applying this to a real problem. (Code available in C++ and Python).