Module 4: Probabilistic Blocking, Part I

Rebecca C. Steorts

Agenda

- Data Cleaning Pipeline
- Probabilistic Blocking
- Locality Sensitive Hashing (LSH)
- Hash functions
- Hashed shingles
- Signatures
- Characteristic Matrix
- Minhash (Jaccard Similarity Approximation)
- Back to LSH

Load R packages

```
## Loading required package: DBI
## Loading required package: RSQLite
## Loading required package: ff
## Loading required package: bit
##
## Attaching package: 'bit'
## The following object is masked from 'package:base':
##
##
       xor
## Attaching package ff
## - getOption("fftempdir")=="/var/folders/bv/xhclmwh90zg08
## - getOption("ffextension")=="ff"
## - getOption("ffdrop")==TRUE
```

Data Cleaning Pipeline

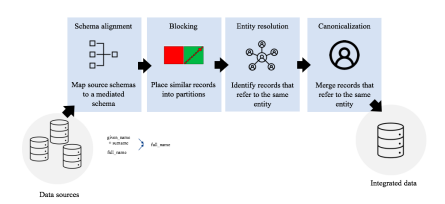


Figure 1: Data cleaning pipeline.

Blocking

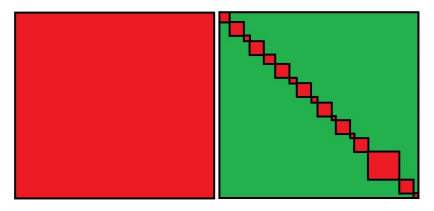


Figure 2: Left: All to all record comparison. Right: Example of resulting blocking partitions.

LSH

Locality sensitive hashing (LSH) is a fast method of blocking for record linkage that orginates from the computer science literature.

Citation data set

Recall that last time we worked with citation data set.

```
# get only the columns we want
n <- nrow(cora) # number of records
dat <- data.frame(id = seq_len(n)) # create id column</pre>
dat <- cbind(dat, cora[, c("title", "authors", "journal")]]</pre>
shingles <- apply(dat, 1, function(x) {
  # tokenize strings
  tokenize_character_shingles(paste(x[-1], collapse=" "), n
})
jaccard <- expand.grid(record1 = seq_len(n), # empty holder</pre>
                        record2 = seq_len(n))
# don't need to compare the same things twice
jaccard <- jaccard[jaccard$record1 < jaccard$record2,]</pre>
```

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 **low**.

Hashing shingles

Instead of storing the strings as shines, we can instead store *hashed* values

These are integers, and they take up less space.

Hashing shingles

```
# 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-sh
   hash_string(shingles) # return hashed shingles
})</pre>
```

Hashing shingles # Jaccard similarity on hashed shingles hashed_jaccard <-</pre>

expand.grid(record1 = seq_len(n), record2 = seq_len(n))
don't need to compare the same things twice

hashed_jaccard < hashed_jaccard[hashed_jaccard\$record1 < hashed_jaccard\$re
time <- Sys.time() # see how long this takes
hashed_jaccard\$similarity <-</pre>

This took up 6.53296×10^5 bytes, while storing the shingles took 8.411816×10^6 bytes; the whole pairwise comparison still took the

come amount of time (a. 1.6 minutes)

Similarity preserving summaries of sets

Sets of shingles are large (larger than the original data set)

If we have millions of records in our data set, 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 Jaccard similarity

Characteristic matrix

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

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

```
Characteristic matrix
   # return if an item is in a list
   item in list <- function(item, list) {</pre>
     as.integer(item %in% list)
   }
   # get the characteristic matrix
   # items are all the unique hash values
   # columns will be each record
   # we want to keep track of where each hash is included
   char_mat <- data.frame(item = unique(unlist(hashed_shingles))</pre>
   # for each hashed shingle, see if it is in each row
   contained <- lapply(hashed shingles, function(col) {</pre>
```

vapply(char_mat\$item, FUN = item_in_list,

})

FUN.VALUE = integer(1), list = col)

char_mat <- do.call(cbind, contained) # list to matrix

Minhashing

We want to create the signature matrix through minhashing

- 1. 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)

```
# set seed for reproducibility
set.seed(02082018)
# function to get signature for 1 permutation
get_sig <- function(char_mat) {</pre>
  # get permutation order
 permute order <- sample(seg len(nrow(char mat)))
  # get min location of "1" for each column (apply(2, ...))
 t(apply(char_mat[permute_order, ], 2,
          function(col) min(which(col == 1))))
# repeat many times
m <- 360
sig_mat <- matrix(NA, nrow=m,
                  ncol=ncol(char_mat)) #empty matrix
for(i in 1:m) {
  sig_mat[i, ] <- get sig(char_mat) #fill matrix</pre>
colnames(sig_mat) <- colnames(char_mat) #column names</pre>
```

Minhashing (cont'd)

```
# inspect results
kable(sig_mat[1:10, 1:5])
```

Record 1	Record 2	Record 3	Record 4	Record 5
3	3	3	3	3
38	38	38	38	38
46	46	46	46	46
36	36	36	36	36
31	31	31	31	31
124	124	124	124	124
21	21	21	21	21
9	9	9	9	9
85	85	85	85	85
44	44	44	44	44

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

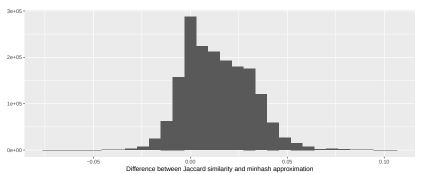
Jaccard similarity approximation

```
# add jaccard similarity approximated from the minhash to
# number of agreements over the total number of combination
hashed_jaccard$similarity_minhash <-
  apply(hashed_jaccard, 1, function(row) {
  sum(sig_mat[, row[["record1"]]]
      == sig_mat[, row[["record2"]]])/nrow(sig_mat)
})
# how far off is this approximation? plot differences
qplot(hashed_jaccard$similarity_minhash - hashed_jaccard$s
  xlab("Difference between Jaccard similarity and minhash a
## `stat bin()` using `bins = 30`. Pick better value with
30+05-
```



Jaccard similarity approximation

`stat_bin()` using `bins = 30`. Pick better value with



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)

Wait did I miss something?

We still haven not addressed the issue of pairwise comparisons

Locality Sensitive Hashing (LSH)

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

```
# view the signature matrix
print(xtable::xtable(sig_mat[1:10, 1:5]), hline.after = c(
```

	Record 1	Record 2	Record 3	Record 4	Record 5
1	3	3	3	3	3
2	38	38	38	38	38
3	46	46	46	46	46
4	36	36	36	36	36
5	31	31	31	31	31
6	124	124	124	124	124
7	21	21	21	21	21
8	9	9	9	9	9
9	85	85	85	85	85
_10	44	44	44	44	44

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

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

```
# library to get divisors of m
library(numbers)
# look at probability of binned together for various bin s
bin_probs \leftarrow expand.grid(s = c(.25, .75), h = m, b = divise
bin_probs$prob <- apply(bin_probs, 1, function(x) lsh_proba
# plot as curves
ggplot(bin probs) +
  geom_line(aes(x = prob, y = b, colour = factor(s), group
  geom_point(aes(x = prob, y = b, colour = factor(s)), size
  xlab("Probability") +
  scale color discrete("s")
```

300 -

"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)</pre>
```

"Easy" LSH in R (Continued)

"Easy" LSH in R (Continued)

```
# perform lsh to get buckets
buckets <- lsh(corpus, bands = b, progress = FALSE)

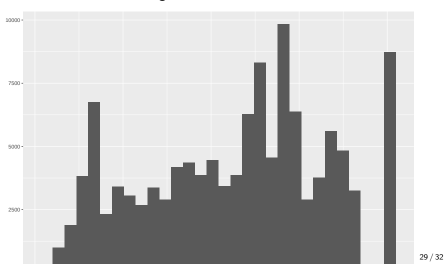
# grab candidate pairs
candidates <- lsh_candidates(buckets)

# get Jaccard similarities only for candidates
lsh_jaccard <- lsh_compare(candidates, corpus, jaccard_similarity)</pre>
```

"Easy" LSH in R (cont'd)

plot jaccard similarities that are candidates
qplot(lsh_jaccard\$score)

`stat_bin()` using `bins = 30`. Pick better value with



Putting it all together

##

library(igraph) #qraph package

Attaching package: 'igraph'

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

```
## The following objects are masked from 'package:stats':
##
##
       decompose, spectrum
## The following object is masked from 'package:base':
##
      union
##
# 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, is.vector((candidates[, 1:2]))) # candidates
g <- set_vertex_attr(g, "id", value = dat$id) # add id
                                                        30 / 32
```

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?
- 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