CS347

Lecture 4
April 18, 2001
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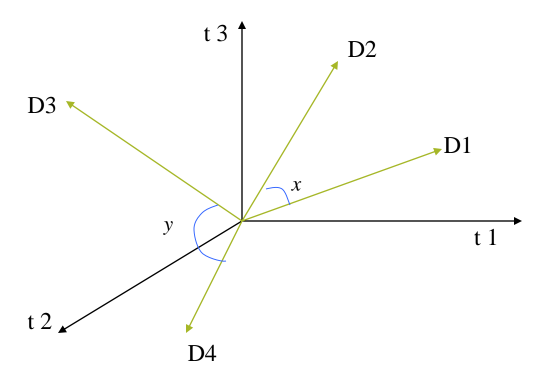
Today's topics

- Computing cosine-based ranking
- Speeding up cosine ranking
 - reducing the number of cosine computations
 - Union of term-wise candidates
 - Sampling and pre-grouping
 - reducing the number of dimensions
 - Random projection
 - Latent semantic indexing

Recall doc as vector

- Each doc j is a vector of $tf \times idf$ values, one component for each term.
- Can normalize to unit length.
- So we have a vector space
 - terms are axes
 - docs live in this space
 - even with stemming, may have 10000+ dimensions

Intuition



Postulate: Documents that are "close together" in vector space talk about the same things.

Cosine similarity

Cosine similarity of D_j, D_k :

$$sim(D_j, D_k) = \sum_{i=1}^{m} w_{ij} \times w_{ik}$$

Aka normalized inner product.

Can also compute cosine similarity from a query (vector of terms, e.g., truth forever) to each document.

Exercises

- How would you augment the inverted index built in lectures 1-3 to support cosine ranking computations?
- Walk through the steps of serving a query.

Why use vector spaces?

- **Key**: A user's query can be viewed as a (very) short document.
- Query becomes a vector in the same space as the docs.
- Can measure each doc's cosine proximity to query \rightarrow ranking.

Efficient cosine ranking

- Ranking consists of computing the k docs in the corpus "nearest" to the query $\Rightarrow k$ largest query-doc cosines.
- Efficient ranking:
 - Computing a single cosine efficiently.
 - Choosing the *k* largest cosine values efficiently.

Computing a single cosine

- For every term i, with each doc j, store term frequency tf_{ij} .
 - Tradeoffs on whether to store term count, freq, or weighted by idf_i . (Coding possibilities.)
- Accumulate component-wise sum

$$sim(D_j, D_k) = \sum_{i=1}^{m} w_{ij} \times w_{ik}$$

More on speeding up a single cosine, later in this lecture.

Computing the *k* largest cosines: selection vs. sorting

- Typically we want to retrieve the top *k* docs (in the cosine ranking for the query)
 - not totally order all docs in the corpus
 - just pick off docs with *k* highest cosines.

Use heap for selecting top k

- Binary tree in which each node's value > values of children
- Takes 2n operations to construct, then each of klog n "winners" read off in 2log n steps.
- For n=1M, k=100, this is about 10% of the cost of sorting.

Bottleneck

- Still need to first compute cosines from query to each of n docs \rightarrow several seconds for n=1M.
- Can select from only non-zero cosines; should be << 1M.

Can we avoid this?

- Yes, but may occasionally get an answer wrong
 - a doc *not* in the top *k* may creep into the answer.

Term-wise candidates

- <u>Preprocess</u>: Pre-compute, for each term, its *k* nearest docs.
 - (Treat each term as a 1-term query.)
 - lots of preprocessing.
 - Result: "preferred list" for each term.
- Search:
 - For a t-term query, take the union of their t preferred lists call this set S.
 - Compute cosines from the query to only the docs in *S*, and choose top *k*.

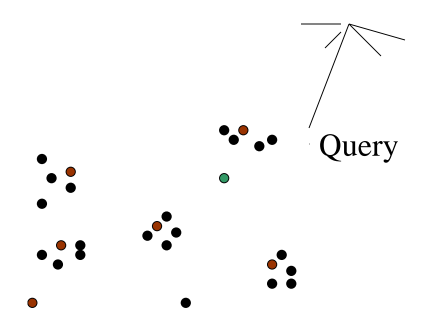
Exercises

- Fill in the details of the calculation:
 - Which docs go into the preferred list for a term?
- Devise a small example where this method gives an incorrect ranking.

Sampling and pre-grouping

- First run a pre-processing phase:
 - pick \sqrt{n} docs at random: call these leaders
 - For each other doc, pre-compute nearest leader
 - Docs attached to a leader: its followers;
 - <u>Likely</u>: each leader has $\sim \sqrt{n}$ followers.
- Process a query as follows:
 - Given query Q, find its nearest leader L.
 - Seek k nearest docs from among L's followers.

Visualization



Leader

Follower

Why use random sampling

- Fast
- Leaders reflect data distribution

General variants

- Have each follower attached to α =3 (say) nearest leaders.
- From query, find b=4 (say) nearest leaders and their followers.
- Can recur on leader/follower construction.

Exercises

- To find the nearest leader in step 1, how many cosine computations do we do?
- What is the effect of the constants a, b on the previous slide?
- Devise an example where this is *likely to* fail we miss one of the *k* nearest docs.
 - Likely under random sampling.

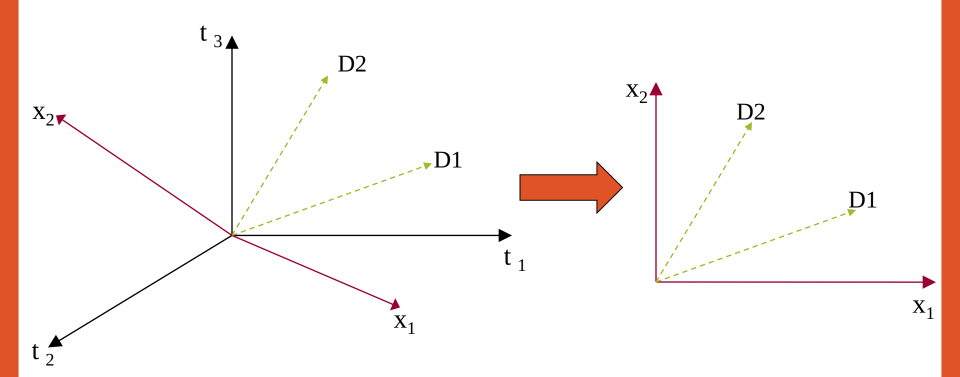
Dimensionality reduction

- What if we could take our vectors and "pack" them into fewer dimensions (say 10000→100) while preserving distances?
- (Well, almost.)
 - Speeds up cosine computations.
- Two methods:
 - Random projection.
 - "Latent semantic indexing".

Random projection onto *k*<<*m* axes.

- Choose a random direction x_i in the vector space.
- For i = 2 to k,
 - Choose a random direction x_i that is orthogonal to $x_1, x_2, \dots x_{i-1}$.
- Project each doc vector into the subspace $x_1, x_2, \dots x_k$.

E.g., from 3 to 2 dimensions



 x_1 is a random direction in (t_1, t_2, t_3) space. x_2 is chosen randomly but orthogonal to x_1 .

Guarantee

- With high probability, relative distances are (approximately) preserved by projection.
- Pointer to precise theorem in Resources.

Computing the random projection

- Projecting *n* vectors from *m* dimensions down to *k* dimensions:
 - Start with $m \times n$ matrix of terms \times docs, A.
 - Find random $k \times m$ orthogonal projection matrix R.
 - Compute matrix product $W = R \times A$.
- *j*th column of *W* is the vector corresponding to doc *j*, but now in *k* << *m* dimensions.

Cost of computation

This takes a total of kmn multiplications.



- Expensive see Resources for ways to do essentially the same thing, quicker.
- Exercise: by projecting from 10000 dimensions down to 100, are we really going to make each cosine computation faster?

Latent semantic indexing (LSI)

- Another technique for dimension reduction
- Random projection was data-independent
- LSI on the other hand is data-dependent
 - Eliminate redundant axes
 - Pull together "related" axes
 - car and automobile

Notions from linear algebra

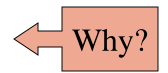
- Matrix, vector
- Matrix transpose and product
- Rank
- Eigenvalues and eigenvectors.

Overview of LSI

- Pre-process docs using a technique from linear algebra called <u>Singular Value Decomposition.</u>
- Have control over the granularity of this process:
 - create new vector space, details to follow.
- Queries handled in this new vector space.

Singular-Value Decomposition

- Recall $m \times n$ matrix of terms \times docs, A.
 - A has rank $r \le m, n$.
- Define term-term correlation matrix *T=AA*^t
 - A^t denotes the matrix transpose of A.
 - T is a square, symmetric $m \times m$ matrix.
- Doc-doc correlation matrix $D=A^tA$.
 - *D* is a square, symmetric $n \times n$ matrix.



Eigenvectors

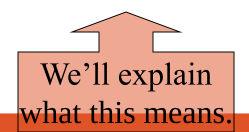
- Denote by P the $m \times r$ matrix of eigenvectors of T.
- Denote by R the $n \times r$ matrix of eigenvectors of D.
- It turns out A can be expressed (decomposed) as $A = PQR^t$
 - *Q* is a <u>diagonal</u> matrix with the eigenvalues of AA^t in sorted order.

Visualization

$$m \times n$$
 = $m \times r$ $r \times r$ $r \times n$
 A P Q R^t

Dimension reduction

- For some $s << r_r$ zero out all but the s biggest eigenvalues in Q.
 - Denote by Q_s this new version of Q.
 - Typically s in the hundreds while r could be in the (tens of) thousands.
- Let $A_s = P Q_s R^t$
- Turns out A_s is a pretty good approximation to A.



Visualization

The columns of A_s represent the docs, but in s << m dimensions.

Guarantee

- Relative distances are (approximately) preserved by projection:
 - Of all $m \times n$ rank s matrices, A_s is the best approximation to A.
- Pointer to precise theorem in Resources.

Doc-doc similarities

- •*A_sA_s^t* is a matrix of doc-doc similarities:
 - •the (j,k) entry is a measure of the similarity of doc j to doc k.

Semi-precise intuition

- We accomplish more than dimension reduction here:
 - Docs with lots of overlapping terms stay together
 - Terms from these docs also get pulled together.
- Thus *car* and *automobile* get pulled together because both cooccur in docs with *tires, radiator, cylinder*, etc.

Query processing

- View a query as a (short) doc:
 - call it row o of A_s .
- Now the entries in row o of $A_s A_s^t$ give the similarities of the query with each doc.
- Entry (o, j) is the score of doc j on the query.
- Exercise: fill in the details of scoring/ranking.

Resources

- Random projection theorem: http://citeseer.nj.nec.com/dasguptaggelementary.html
- Faster random projection: http://citeseer.nj.nec.com/frieze98fast.html
- Latent semantic indexing: http://citeseer.nj.nec.com/deerwestergoindexing.html
- Books: MG 4.6, MIR 2.7.2.