

NLP Galore

Various Sources

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Chapter 1

Introduction

This handbook is about natural language processing for internal use only.

Please **do not** cite.

BASICS OF NLP

Chapter 2

Dictionary-Based Methods

We first describe dictionary-based methods, which are by far the most common and least resource-intensive way of conducting textual analysis.

2.1 A Stylized Guide

Many papers employing dictionary-based methods follow this cookbook:

1. What do you want to measure, and why is it important?
2. How will you measure what you want to measure? Which corpus will you use? Which dictionary will you use?
3. Which economic variables are you interested in explaining with the measurement?

Many well-known papers follow this format:

- Cieslak and Vissing-Jorgensen (2021) study the Fed’s attention to the stock market, which is important given its prominent role in the conduct of monetary policy. To measure this object, they use the FOMC documents as corpus and utilize a self-created dictionary of stock market keywords. They then relate the measured attention to the Fed’s interest rate decisions and updates to its growth expectations.
- Sautner, Lent, Vilkov, and Zhang (2022) is interested in measuring firm-level exposure to climate change. Given rising climate risk and the accompanying regulator attention, measuring the exposures is of key importance. To measure this object, they use the earnings conference calls as corpus and utilize a dictionary created using a keyword-discovery algorithm. They relate the measured exposures to green technology hiring and patenting as well as prices of options and equity claims.
- Hassan et al. (2022) seeks to measure country risk as perceived by investors and firms, which is a critical determinant of capital movements

across countries. They use earnings conference calls as corpus and use synonyms of risk and uncertainty obtained from Oxford dictionary, library of tone words from Loughran and McDonald (2011), and a self-created dictionary of country-specific keywords. Ultimately, they relate the measured country risk to capital flows and sudden stops, firm-level investment and employment decisions, and fluctuations in exchange rates.

2.2 Often-Used Dictionaries

2.2.1 Loughran-McDonald (LM) Dictionary

Loughran and McDonald (2011) created six different word lists (negative, positive, uncertainty, litigious, strong modal, and weak modal) by examining word usage in a large sample of 10-Ks during the period 1994–2008. Their dictionary has become predominant in many studies, and papers have used the LM word lists (primarily negative words) to gauge the tone of the business communication. Data can be obtained at [this link](#).

Chapter 3

Regression-Based Methods

Using document characteristics, these methods try to predict word frequencies in a given document. A natural way to model these relationships is using multinomial logistic regression; however, due to the large dimensionality of vocabularies, standard multinomial logistic regressions are not computationally feasible to implement. In recent years, econometric breakthroughs have led to the development of computationally feasible approximations of the ideal multinomial logit regression (Taddy, 2015).

These models have several applications. Firstly, the method tells us how the text depends on observed covariates. This is relevant in settings where the dependent variable is the text, e.g., political speeches. Secondly, once estimated, the model can be inverted to get predicted values of covariates using text as an input. This is particularly powerful for (i) nowcasting/forecasting and (ii) backcasting when data is missing (Kelly et al., 2021). For example, measurements of real activity such as GDP are only available with a lag and a quarterly frequency. However, economic media coverage is available in real-time and with low latency. Hence, it can be used to nowcast GDP. Similarly, when data have short histories, text can be used to backcast the data.

3.1 Challenges of having text as the dependent variable

The typical unit of analysis for text data is at the frequency of an n -gram in a given document, e.g., the number of times the word “GDP” turned up in a document. These frequencies are unordered uncategorical data, i.e., it is not obvious how to combine information that a document has ten mentions of the word “GDP” with six mentions of the word “inflation.” Hence, frequency counts are represented by a document i specific vectors that sit in $\mathbf{c}_i \in \mathbb{R}^V$ where V is

the number of unique n-grams in our sample. Note that V is extremely large in practice which is why machine learning methods are needed.

The go-to model to explain the document i specific count vector \mathbf{c}_i using variables \mathbf{v}_i would be a multinomial logit regression,

$$p(\mathbf{c}_i | \mathbf{v}_i, m_i) = \text{MN}(\mathbf{c}_i; \mathbf{q}_i, m_i) \text{ for } i = 1 \dots n$$

$$q_{ij} = \frac{e^{\eta_{ij}}}{\sum_{k=1}^d e^{\eta_{ik}}} \text{ for } j = 1 \dots d$$

$$\eta_{ij} = \alpha_j + \mathbf{v}_i' \boldsymbol{\varphi}_j$$

where q_{ij} is the n-gram's probability, m_i is the total number of n-grams in the document, and (α_j, φ_j) are the $K + 1$ n-gram specific parameters.

Given vocabularies often exceed 10,000 n-grams, estimating the model above with even just two covariates would mean estimating 30,000+ Parameters! The issue with standard multinomial estimation is that it cannot be parallelized; hence, estimating them for a large number of parameters is computationally prohibitive.

Multinomial logit regressions cannot be parallelized because the denominator of q_{ij} which depends on all the parameters in the model. Hence, to estimate the model, we need a parallelizable approximation. Taddy (2015) provided precisely such an approximation and used it to develop the distributed multinomial regression (DMR)

3.2 Distributed multinomial regressions (DMR)

A canonical relationship is that Multinomial distributions can be exactly decomposed into independent Poissons,

$$\text{MN}(\mathbf{c}_i; \mathbf{q}_i, m_i) = \frac{\prod_j P_o(c_{ij}; e^{\eta_{ij}})}{P_o(m_i; \sum_{j=1}^d e^{\eta_{ij}})}$$

This decomposition does not solve the computation bottleneck since the denominator still depends on all parameters. However, Tadd (2015) used this decomposition to develop the following approximation,

$$p(\mathbf{c}_i | \mathbf{v}_i, m_i) = \text{MN}(\mathbf{c}_i; \mathbf{q}_i, m_i) \approx \prod_j \text{Po}(c_{ij}; m_i e^{\eta_{ij}})$$

Since this approximation doesn't depend on $\sum_{j=1}^d e^{\eta_{ij}}$ It can be parallelized! Essentially, each n-gram's parameters are estimated separately using (shifted) Poisson regressions.

Taddy (2015) also developed a method for inverting the DMR so that post-estimation n-grams can be projected into covariates.

3.3 Extension: Hurdle DMR (HDMR)

In practice, while DMR works well for explaining strictly positive frequency counts (i.e., counts > 0), it performs poorly in explaining whether a word is used at all (i.e., count 0 or 1). Kelly et al. (2021) developed the HDMR to simultaneously explain both the intensive margin (strictly positive frequency counts) and extensive margin (counts of zero and one) of word choice.

At a high level, the model combines a selection model with the DMR. The selection model approximates the extensive margin, and the DMR models the intensive margin.

Specifically, the selection model to include n-gram j in document i is standard,

$$h_{ij}^* = \gamma_i + \kappa_j + w_i' \delta_j + v_{ij}$$

$$h_{ij} = \mathbf{1}(h_{ij}^* > 0)$$

where w_i are observed variables and $(\gamma_i, \kappa_j, \delta_j)$ are parameters. And the count model has the following form,

$$c_{ij}^* = \lambda(\mu_i + \alpha_j + \mathbf{v}_i' \varphi_j) + \varepsilon_{ij}$$

$$c_{ij} = (1 + c_{ij}^*) h_{ij}$$

where v_i are observed variables and $(\mu_i, \alpha_j, \varphi_j)$ are parameters. $\lambda(\cdot) \geq 0$, hence the second equation restricts to positive counts. Like DMR, this method is also parallelizable and has an inversion method for projecting text to covariates.

3.4 Application 1: using text to measure traditionally difficult to quantify concepts

Gentzkow et al. (2019) used DMR to measure partisanship in US political—a traditionally allusive concept to quantify. They defined partisanship as the posterior probability, that an observer with a neutral prior on a politician, expects to identify a speaker’s party after hearing the speaker utter a single word. Given this definition, they use DMR to estimate the n-gram probabilities for Democrats and Republicans. They then map this empirical distribution to the posterior belief that an observer with a neutral prior assigns to a speaker being Republican if she utters phrase j in session t and has characteristics x . This definition of partisanship only considers the extensive margin of word choice e.g. whether the politician chooses to use the n-gram “pro-life” in their speeches or not.

Kelly et al. (2021) expand the definition of partisanship above to include both the extensive and intensive margin of word choice i.e. partisanship as the posterior probability, that an observer with a neutral prior on a politician, expects to identify a speaker’s party after hearing the speaker utter ~~a single word~~ the words in their speech. This definition not only considers whether the word “pro-choice” was used, but also how often was it used. Since, Kelly et al. (2021) definition of partisanship considers both the extensive and intensive margin of word choice they use HDMR.

The figure below shows the resulting partisanship measures for the two measures. The left hand side figure shows the results from the using the DMR with the extensive margin definition of partisanship. The DMR estimate suggests partisanship has significantly increased in the last 20 years. The right hand side figure whows that when the intensive margin is also considered, we see spikes in partisanship in the 20’s in addition to the current period too.

3.5 Application 2: Using text data to forecast, nowcast and backcast hard data

Once a linear regression model $y_i = \sum_k x_{i,k} \beta_k$ is estimated, we can use y_i , $\hat{\beta}$ and $x_{i,2}, \dots, x_{i,K}$ to get a fitted value for $x_{i,1}$. Similarly with DMR and HDMR, once the model is estimated, you can use text data to get fitted values for covariates in an *inverse regression*. Hence, text data can be used to forecast, nowcast and backcast hard data.

For example, text data can help with better nowcasting and forecasting of macro variables. See Kelly et al. (2021) for this application. Text data is typically available at a higher frequency and with less latency than most macro data, and hence, can potentially improve macro variable nowcasts and forecasts.

Similarly, often many data series have short-time series, whereas newspaper articles on the topic are available much further back. In this case news articles can be used to backcast data with short time series. For example, Kelly et al. (2021) demonstrate how text data (along with other financial data) can be used to extend back the intermediary capital ratio (ICR) back to the 1930s—ICR is only available for the post 70s period.

Chapter 4

Bag of Words: TF-IDF

4.1 Bag of Words

A bag of words is a representation of text that describes the occurrence of words within a document. We just keep track of word counts and disregard the grammatical details and the word order. It is called a “bag” of words because any information about the order or structure of words in the document is discarded. The model is only concerned with whether known words occur in the document, not where in the document.

4.1.1 Bag of N-Grams

4.1.1.1 The Idea

In general, bigrams make tokens more understandable. Suppose we have two sentences:

- Sentence 1: “This is a good job. I will not miss it for anything”
- Sentence 2: ”This is not good at all”

and let us take the vocabulary of 5 words only: “good”, “job”, “miss”, “not”, and “all.”

In this case, the respective vectors for these sentences are:

- Sentence 1: “This is a good job. I will not miss it for anything” = $[1,1,1,1,0]$
- Sentence 2: ”This is not good at all” = $[1,0,0,1,1]$

Sentence 2 is a negative sentence, yet this distinction is not reflected in the vectors.

Now, suppose instead we use bigrams, in which case Sentence 2 can be broken into: “This is”, “is not”, “not good”, “good at”, and “at all.” In this case, the

model can differentiate between sentence 1 and sentence 2.

4.1.1.2 Pre-Processing to Reduce Feature Space

Using N-grams increases the feature space, so a1. few techniques are used to reduce it:

- Removing stopwords such as a, the, and, it, is, etc.
- Stemming: This is the process of reducing a word to its word stem. For example, words like swimmer, swimming, swim, will be mapped to one-word swim.
- Chunking and Parts-of-speech tagging: One can use these techniques to find meaningful words in a sentence and use them as the feature vector

4.2 TF-IDF

TF-IDF is intended to reflect how relevant a term is in a given document. It makes rare words more prominent and effectively ignores common words. Therefore, unlike bag-of-words, it creates a normalized count where each word count is divided by the number of documents this word appears in.

It is computed by multiplying two different metrics:

- **Term Frequency (TF)** of a word t in document d is given as

$$tf(t, d) = \frac{f_{t,d}}{\sum_{t' \in d} f_{t',d}}$$

where $f_{t,d}$ is the raw count of a term in a document. The denominator is the total number of terms in document d .

- **Inverse Document Frequency (IDF)** of a word t in a set of documents D is given as

$$idf(t, D) = \log \frac{N}{|\{d \in D : t \in d\}|}$$

where $N = |D|$ and the denominator is the number of documents where the term t appears.

Then TF-IDF is then calculated as:

$$tfidf(t, d, D) = tf(t, d) \times idf(t, D)$$

A high weight in TF-IDF is reached by a high term frequency (in the given document) and a low document frequency of the term in the whole collection of documents

4.3 Limitations of Bag of Words

Although Bag-of-Words is quite efficient and easy to implement, there still are disadvantages.

1. **It suffers from the curse of dimensionality** as the total dimension is generally the vocabulary size. Therefore, Its vocabulary needs to be designed carefully to manage the size. At the same time, bag of words also often leads to sparse vectors, which require more memory and computational resources.
2. **The model ignores the location information of the word.** The location information is a piece of very important information in the text. For example “today is off” and “Is today off”, have the exact same vector representation in the BoW model.
3. **The model ignores the semantics of the word.** For example, words ‘soccer’ and ‘football’ are often used in the same context. However, the vectors corresponding to these words are quite different in the bag of words model.
4. **The range of vocabulary is a big issue faced by the Bag-of-Words model.** If the model comes across a new word, it ends up ignoring the word.

Chapter 5

Basic Word Embeddings: Word2Vec & Glove

5.1 Overview

Word embedding simply refers to representing words as word vectors. The idea is that we have a large corpus of text and every word in a fixed vocabulary is represented by a vector.

Need for Word Embedding

Various word encoding methods like Integer/ Label Encoding or One-Hot Encoding have many limitations. The main limitation is that these encoding doesn't have the semantic relationship between the words. Therefore, an alternative approach is to learn to encode similarity in the vectors themselves.

In addition, for one-hot encoding, memory requirement and the features space is increasing in the vocabulary size.

Generating Word Embeddings

There are generally two methods for generating word embeddings: (1) SVD based methods and (2) neural network (iteration) based methods.

- In SVD based embedding methods, first, we create the matrix of co-occurrence and then reduce the dimensionality of the matrix using SVD. After applying SVD, each word in the vocabulary has an embedding in reduced space.

While these methods effectively leverage global statistical information, they are primarily used to capture word similarities and do poorly on tasks such as word analogy, indicating a sub-optimal vector space structure.

- In neural net based methods, instead of computing and storing global information about some huge dataset, one can try to create a model that will be able to learn one iteration at a time.

These methods are shallow window-based, which learn word embeddings by making predictions in local context windows.

5.2 SVD Based Methods

One basic idea is to accumulate word co-occurrence counts in matrix X and then perform Singular Value Decomposition (SVD) on X to get USV^T decomposition. Then the first k columns of U can be used as k -dimensional word vectors.

5.2.1 Limitations

There are numerous limitations of SVD based methods:

- Generally, the matrix is of very high dimension, which results in high training cost $O(mn^2)$ where m is the size of a dictionary and n is the embedding size. So for a corpus with a large number of words, it becomes very difficult to train.
- The matrix quickly becomes imbalanced due to high frequency words.
- The dimension of the matrix changes as soon as a new word is introduced.

To get around this problem, one can use the following tricks:

- Ignore the common words like “is”, “the” etc.
- The weight of the two different words should be considered based on the distance between the two words, rather than just raw count.

5.3 Iteration Based Methods

The idea is to design a model whose parameters are the word vectors.

5.3.1 Word2Vec

Word2Vec is one of the most popular technique to learn word embeddings using shallow neural network, developed by Tomas Mikolov in 2013 at Google. It is a shallow, two-layer neural network that is trained to reconstruct linguistic contexts of words.

- It takes a large corpus of words as input and outputs a vector space with hundreds of dimensions, with each unique word in the corpus allocated to a corresponding vector in the space.

It can be obtained using two model architectures: Continuous Bag of Words (CBOW) and Skip-gram. CBOW aims to predict a center word from the sur-

rounding context in terms of word vectors. Skip-gram does the opposite, and predicts the distribution (probability) of context words from a center word.

5.3.1.1 Method #1: CBOW

The core idea is to predict a center word from the surrounding context. For each word w_i , we learn 2 vectors: v_i , the representation when the word is in the context, and u_i , the representation when the word is in the center.

Using the Model

The goal is to learn two matrices, $\mathcal{V} \in \mathbb{R}^{n \times |V|}$ and $\mathcal{U} \in \mathbb{R}^{|V| \times n}$. \mathcal{V} is the input word matrix where the i th column is v_i , and \mathcal{U} is the output word matrix where the j th row is u_j . Equipped with this matrix, the model then predicts the center word through the following steps:

1. For the input context of size m , one hot word vectors are generated: $x^{(c-m)}, \dots, x^{(c-1)}, x^{(c+1)}, \dots, x^{(c+m)} \in \mathbb{R}^{|V|}$
2. We obtain the embeddings for each vector via $v = \mathcal{V}x$.
3. We average the vectors to get \hat{v} and generate a score vector $z = \mathcal{U}\hat{v} \in \mathbb{R}^{|V|}$.
4. Turn the scores into probabilities via $\hat{y} = \text{softmax}(z) \in \mathbb{R}^{|V|}$, which we'd like to be close to the true probability y – which happens to be one hot vector of the actual word – as much as possible.

Training the Model

In training the model, we often use the cross entropy $H(\hat{y}, y)$ as the measure of distance:

$$H(\hat{y}, y) = - \sum_{j=1}^{|V|} y_j \log(\hat{y}_j)$$

Therefore, the optimization problem can be framed as minimizing the objective J where

$$J = -\log P(w_c | w_{c-m}, \dots, w_{c-1}, w_{c+1}, \dots, w_{c+m})$$

Using the notations for the embeddings,

$$= -\log P(u_c | \hat{v}) = -\log \frac{\exp(u_c^\top \hat{v})}{\sum_{j=1}^{|V|} \exp(u_j^\top \hat{v})}$$

Therefore we can use stochastic gradient descent to update all relevant word vectors u_c and v_j .

5.3.1.2 Method #2: Skip-Gram

The core idea is to predict surrounding context words given a center word. The setup is largely the same as CBOW with the role of center and context words reversed.

Using the Model

1. Generate one hot input vector $x \in \mathbb{R}^{|V|}$ of the center word.
2. Use the embeddings to get the embedded word vector for the center word:
 $v_c = \mathcal{V}x \in \mathbb{R}^n$.
3. Generate a score vector $z = \mathcal{U}v_c$.
4. Turn the score vector into probabilities $\hat{y} = \text{softmax}(z)$ which yields the probabilities of observing each context word: $\hat{y}_{c-m}, \dots, \hat{y}_{c-1}, \hat{y}_{c+1}, \dots, \hat{y}_{c+m}$.
 As before, we want these probability vectors to match the one hot vectors of the actual output.

Training the Model

Given this task is a bit more daunting, we need one additional assumption of strong conditional independence. In other words, given the center word, all output words are completely independent.

Therefore, the objective can be written as:

$$J = -\log P(w_{c-m}, \dots, w_{c+m} | w_c) = -\log \prod_{j=0, j \neq m}^{2m} P(w_{c-m+j} | w_c)$$

Once again, applying the embeddings yields:

$$= -\log \prod_{j=0, j \neq m}^{2m} P(u_{c-m+j} | v_c) = -\log \prod_{j=0, j \neq m}^{2m} \frac{\exp(u_{c-m+j}^\top v_c)}{\sum_{k=1}^{|V|} \exp(u_k^\top v_c)}$$

In essence, Skip-gram treats each context word equally: the model computes the probability for each word of appearing in the context independently of the distance to the center word.

5.3.1.3 Implementation Details

According to the original paper, it is found that Skip-Gram works well with small datasets, and can better represent less frequent words. However, CBOW is found to train faster than Skip-Gram, and can better represent more frequent words.

The original authors of the method also provide two implementation details that improve the training performance: (i) negative sampling and (ii) hierarchical softmax.

Negative Sampling

By defining a new objective function, negative sampling aims at maximizing the similarity of the words in the same context and minimizing it when they occur in different contexts. However, instead of doing the minimization for all the words in the dictionary except for the context words, it randomly selects a handful of words depending on the training size and uses them to optimize the objective.

Heirarchical Softmax

Heirarchical softmax is a replacement for softmax which is much faster to evaluate: softmax is $O(n)$ time, while hierarchical softmax is $O(\log n)$ time.

You can view the softmax function as a tree, where the root node is the hidden layer activations (or context vector C), and the leaves are the probabilities of each word. A hierarchical softmax instead computes the values of the leaves with a multi-layer tree. To evaluate the probability of a given word, take the product of the probabilities of each edge on the path to that node.

5.4 Best of Both Worlds: GloVe

The starting point for GloVe is to reconcile linear algebra algorithms with neural updating algorithms. The main advantage of GloVe is that unlike Word2vec, GloVe does not just rely on local statistics but also incorporates global statistics (word co-occurrence) to obtain word vectors.

The main insight is that the ratio of the co-occurrence probabilities of two words (rather than their co-occurrence probabilities themselves) is what contains information and so we look to encode this information as vector differences.

The Model

Let X denote the co-occurrence matrix where X_{ij} is the number of times word j occurs in the context of word i . And denote $P_{ij} = P(w_j|w_i) = X_{ij}/X_i$ as the probability of j appearing in the context of word i where $X_i = \sum_k X_{ik}$.

- As one can see, populating X and computing P requires a single pass through the entire corpus, which is a one-time up-front cost.

The objective J is the global cross-entropy loss:

$$J = - \sum_{i \in \text{corpus}} \sum_{j \in \text{context}(i)} \log Q_{ij}$$

where Q_{ij} is the probability of word j appearing in the context of word i as in the skip-gram model. We can then rewrite the objective as

$$J = - \sum_{i=1}^W \sum_{j=1}^W X_{ij} \log Q_{ij}$$

Since Q_{ij} requires normalization and thus incurs a summation over the entire vocabulary, a least squares objective is used instead:

$$\hat{J} = \sum_{i=1}^W \sum_{j=1}^W X_i (\hat{P}_{ij} - \hat{Q}_{ij})^2$$

where $\hat{P}_{ij} = X_{ij}$ and $\hat{Q}_{ij} = \exp(u_j^\top v_i)$ are the unnormalized distribution. But since X_{ij} can be large, we will instead minimize the squared error in the log

space and introduce a more general weighting scheme:

$$\hat{J} = \sum_{i=1}^W \sum_{j=1}^W f(X_{ij})(u_j^\top v_i - \log X_{ij})^2$$

Chapter 6

Topic Modelling: LDA

The goal of topic modeling is to extract latent topics from observed documents. For example, we may have a collection of news articles where any given article may discuss the economy, geopolitics, or both; the goal of topic modeling is to extract these topics in an unsupervised manner. Alternatively one can also think of topic modeling as a dimension reduction technique; documents are high-dimensional objects containing a large collection of words, and we would like to map them to a lower-dimensional topic space.

A popular topic model is the Latent Dirichlet Allocation (LDA) which was first proposed by Blei et al. (2003). The LDA is an unsupervised technique, which takes as its input a corpus of documents and outputs latent topics i.e. factors that load on words. In this chapter, we will cover LDAs.

6.1 Notation and Terminology

- **Words** indexed $1, \dots, V$ are represented by the basis vector—the collection of words that make up a corpus will be referred to as the vocabulary.
- **Topics** are denoted z_1, \dots, z_K . In all topic models discussed here, the researcher will need to specify the number of K latent topics we are trying to extract.
- **Document** m is represented by a sequence of sequence of N words, $d_m = \{w_1, w_2, \dots, w_N\}$ e.g. a newspaper article. *For simplicity, unless otherwise stated, throughout this chapter we are going to assume the length of documents is exactly N words.*
- **Corpus** is composed of a collection of M documents $\mathcal{D} = \{d_1, \dots, d_M\}$ e.g. the New York Times.
- **Document-term matrix** is a way to describe how often a words occur across documents in the corpus $X \in \mathbb{R}^{M \times V}$.

- A very basic version would involve simply doing a frequency count of all possible V words in each of the M documents.

6.2 Models that preceded the LDA:

Before diving into LDAs, it is useful to cover a couple of models that preceded the LDA. This serves three pedagogical purposes. Firstly, it helps see the deep parallels of topic modeling techniques with more commonly used dimension reduction methods such as principle component analysis (PCA). Secondly, it motivates the exact challenges faced by past methods LDAs were trying to address, and also shows the challenges that still remain.

6.2.1 Latent Semantic Models (LSA)

One way to think about topic modeling is that it is trying to get a low-dimensional representation of the documents, where these low-dimensional representations can be thought of as topics. Hence, one might attempt to simply apply a PCA to the document-term matrix, and interpret the principle components as topics. This is precisely the idea behind LSAs.

LSA applies a Singular Value Decomposition (SVD) to the document-term matrix and interprets the extracted factors as topics. While this method is excellent for dimension reduction, it does not allow for interpretability, since PCA only identifies the loadings up to orthogonal rotations. Additionally, SVDs require a large amount of data to get reasonable estimates. For both of these reasons, researchers decided to put a more parametric structure on the problem.

6.2.2 Probabilistic Latent Semantic Models (pLSA)

The next iteration on the LSA put more structure on the problem by using a two-level hierarchical Bayesian set-up to model the data generating process of a document. The DGP provides a joint distribution for documents, words, and topics. We then try to estimate the posterior of latent topics conditional on observed documents and words. We can then use this to map documents to topics.

This parameterization has direct parallels to the SVD (see great explanation here), and it allows for greater interpretability and more efficient estimation. It is worth discussing some of the specifics of the pLSA, as the LDA directly builds on it.

The pLSA's DGP attempts to capture two key properties of how we intuitively think of documents, topics and words are related: (i) a document can be composed of multiple topics (e.g. an article can be about both economics and geopolitics), and (ii) topics tend to be associated with a set of words (e.g. economics is associated with words like GDP, inflation, currency, etc.). The pLSA captures the former by modeling each document as a multinomial distribution from which

topics are drawn—the document-specific multinomial parameter captures the document-specific topic distribution. Similarly, it captures the latter by modeling words as being drawn from topic-specific multinomial distributions. More formally, the DGP for a given document m containing N words are as follows,

1. For each N words w_n in document m :
 1. Choose topic $z_n \sim \text{Multinomial}(\theta_m)$.
 2. Choose word w_n from the multinomial $p(w_n|z_n; \beta)$ where β is a $V \times K$ matrix such that each column is a topic-specific word distribution.

The pLSA algorithm can be represented using a flow chart (augmented from Blei et al. (2003)) can be graphically represented as follows,

pLSA's were a major improvement on the LSA. Namely, by assuming the latent topics were drawn from multinomial distributions, they were able to uniquely identify the topic loadings and allow for greater interpretability. However, a major limitation of the pLSA is that it requires estimating the multinomial parameter θ_m for each document separately. As a result, the pLSA requires each document to have a large number of words to estimate these parameters consistently. Hence, in practice, the pLSA can be quite unstable. LDA's built on top of pLSAs to solve this issue.

6.3 Latent Dirichlet Allocation (LDA)

To more efficiently use the data across documents, LDAs assume that the document-specific distribution is drawn from a Dirichlet distribution $\theta_m \sim \text{Dir}(\alpha)$. This Bayesian hierarchical structure helps parameter shrinkage; θ_m for documents with fewer words is shrunk towards the population mean. This makes the model more stable than pLSA.

The LDA algorithm is as follows,

1. For each document m choose $\theta_m \sim \text{Dir}(\alpha)$
2. For each N words w_n in document m :
 1. Choose topic $z_n \sim \text{Multinomial}(\theta)$.
 2. Choose word w_n from the multinomial $p(w_n|z_n; \beta)$ where β is a $V \times K$ matrix such that each column is a topic-specific word distribution.

Similarly, this can be graphically represented as follows (from Blei et al. (2003)).

Hopefully, it is clear from both the algorithm and the figures, that the main difference between LDAs and pLSAs is the additional assumption that θ_m is drawn from a Dirichlet distribution.

6.3.1 Challenges of working with LDA

Note that while the Dirichlet prior helps with shrinkage of the topic-document distribution, we are still left with the issue that the number of parameters grows as the vocabulary growth— is $V \times K$ and is $V \times 1$. Hence, to make LDA's work in practice considerable pre-processing of the data is needed to ensure V isn't too big and the model can be estimated easily.

More recently, methods combining word embeddings and LDAs have been proposed to solve this challenge.

6.3.2 Intractable likelihood and estimation

The probability of observing given words in terms of model parameters is,

$$p(\mathbf{w} \mid \alpha, \beta) = \frac{\Gamma(\sum_i \alpha_i)}{\prod_i \Gamma(\alpha_i)} \int \left(\prod_{i=1}^k \theta_i^{\alpha_i - 1} \right) \left(\prod_{n=1}^N \sum_{i=1}^k \prod_{j=1}^V (\theta_i \beta_{ij})^{w_n^j} \right) d\theta$$

As you can see this is quite messy, which makes it intractable to find an analytical expression for model parameters α and β . Hence, we need to rely on numerical methods such as MCMC or variational inference (VI). As is standard in many ML settings, due to the large dimensionality of the problem, the preferred method for estimating LDAs is VI.

6.3.3 Packages for implementing the standard LDA

Python has packages that can implement LDAs out of the box,

- `sklearn.decomposition.LatentDirichletAllocation`
- `gensim.models.ldamodel.LdaModel`

6.3.4 Extensions to the standard LDA

6.3.4.1 Dynamic LDA

The LDA assumes the parameters α and β are constant over time. However, this may not be an appropriate assumption over longer horizons. For example, the topics being discussed in scientific journals have considerably changed over the last century. A naive method for addressing this issue would be to repeatedly estimate the LDA for a year at a time. However, this approach throws out a lot of useful information, as topics in a given year should be somewhat informative about topics in the following year. Hence, the dynamic LDA was developed Blei and Lafferty (2006).

The dynamics LDA attempts to estimate a sequence of parameters (α_t, β_t) . It essentially connects a many LDA's together by assuming the α_t and β_t evolve as random walked with Gaussian shocks (so can be extracted using Kalman filters). The algorithm is,

1. Draw topics $\beta_t \mid \beta_{t-1} \sim \mathcal{N}(\beta_{t-1}, \sigma^2 I)$.
2. Draw $\alpha_t \mid \alpha_{t-1} \sim \mathcal{N}(\alpha_{t-1}, \delta^2 I)$.
3. For each document:
 - (a) Draw $\eta \sim \mathcal{N}(\alpha_t, a^2 I)$
 - (b) For each word:
 - i. Draw $Z \sim \text{Mult}(\pi(\eta))$.
 - ii. Draw $W_{t,d,n} \sim \text{Mult}(\pi(\beta_{t,z}))$.

Where π maps the multinomial natural parameters to the mean parameters,

$$\pi(\beta_{k,t})_w = \frac{\exp(\beta_{k,t,w})}{\sum_w \exp(\beta_{k,t,w})}.$$

Graphically this is can be represented as,

6.3.4.2 Supervised LDA

In some contexts, we both have text data and also a response variable. For example, with movie ratings, we can see people's reviews and the rating they give the movie. We could use this paired data to help better identify topics (but also to predict responses using text data). This is the idea behind supervised LDA (sLDA) (Blei and McAuliffe, 2007).

The sLDA has each word is a function of the latent topic (like the standard LDA), and it also has the document-specific response variable be a function of all the latent topics in a given document. The DGP is described by the algorithm,

1. Draw topic proportions $\theta \mid \alpha \sim \text{Dir}(\alpha)$.
2. For each word
 - (a) Draw topic assignment $z_n \mid \theta \sim \text{Mult}(\theta)$.
 - (b) Draw word $w_n \mid z_n, \beta_{1:K} \sim \text{Mult}(\beta_{z_n})$.
3. Draw response variable $y \mid z_{1:N}, \eta, \sigma^2 \sim \text{N}(\eta^\top \bar{z}, \sigma^2)$.

where \bar{z} is the mean of all topics in the document.

Graphically this is can be represented as,

Chapter 7

Sequence Models: RNNs and LSTMs

7.1 Sequence Models

Unlike models that work with inputs consisting of a single feature vector $\mathbf{x} \in \mathbb{R}^d$, sequence models work with inputs that consist of an ordered list of feature vectors $\mathbf{x}_1, \dots, \mathbf{x}_m$ where each feature vector is now indexed by time or sequence step t .

In the context of natural language processing, we often talk about language models. Specifically, the goal of language models is to compute the probability of sequence of m words, which is usually conditioned on a window of n previous words, as opposed to all previous words:

$$P(w_1, \dots, w_m) = \prod_{i=1}^{i=m} P(w_i | w_1, \dots, w_{i-1}) \approx \prod_{i=1}^{i=m} P(w_i | w_{i-n}, \dots, w_{i-1})$$

7.1.1 n-gram Language Models

One way to compute the above probabilities is to compare the frequency of each word against the frequency of each n -gram that contains the word.

- Bigram language model:

$$P(w_2 | w_1) = \frac{\text{count}(w_1, w_2)}{\text{count}(w_1)}$$

- Trigram language model:

$$P(w_3 | w_1, w_2) = \frac{\text{count}(w_1, w_2, w_3)}{\text{count}(w_1, w_2)}$$

This approach runs into some obvious issues. First, the numerator may be zero due to sparsity. One solution is to add a small δ to the count for each word in the vocabulary. Second, the denominator may be zero due to sparsity, in which case we may “back off” and condition on $\text{count}(w_2)$ rather than on $\text{count}(w_1, w_2)$. Third, since this step requires storing all n -grams, there may be issues with storage.

7.2 RNN: Neural Networks with Memory

Recurrent Neural Networks (RNNs) are a family of neural networks that capture the dynamics of sequences via recurrent connections. The idea is that instead of burdening our model with predicting an output in one go, we allow it the much easier task of predicting iterative sub-outputs, where each sub-output is an improvement or refinement on the previous step.

Hidden State = Memory

The core idea is that instead of modeling $P(w_m | w_{m-1}, \dots, w_{m-n+1})$ it is instead preferable to use a latent variable model:

$$P(w_t | w_{t-1}, \dots, w_1) = P(w_t | h_{t-1})$$

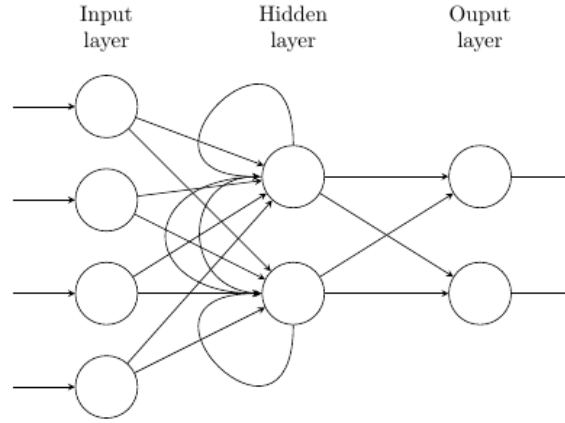
where h_{t-1} is a hidden state that stores the sequence information up to time step $t - 1$. In general, the hidden state can depend on both the current input and the previous hidden state: $h_t = f(x_t, h_{t-1})$.

Recurrence

We can also view RNNs as traditional neural networks enhanced with a loop, one that allows for information to persist across timesteps – hence the name, “recurrent.” [This also implies that the same weight matrix is applied.]

7.2.1 Basic RNN Architecture

An RNN can be represented by an internal hidden state h_t and output o_t :



$$\begin{cases} h_t = f(W_{xh}x_t + W_{hh}h_{t-1} + b_h) \\ o_t = g(W_{hy}h_t + b_o) \end{cases}$$

- $W_{xh}x_t$: Input x_t is multiplied by weight matrix W_{xh} to extract information from the input
- $W_{hh}h_{t-1}$: Previous hidden state h_{t-1} is multiplied by weight matrix W_{hh} to extract information from “memory”
- Activation functions: f is usually **tanh** or **ReLU** and g is often **softmax**
- b_h and b_o are biases.

Advantages

As we continue to emphasize, recurrent neural networks can model sequence of data – either time-series or words in a sentence – so that each sample is assumed to be dependent on the previous ones. From a modelling perspective, it is also very convenience since it can process input sequences of any length, and the model size does not increase for longer input sequence lengths. This is possible because computation for step t can use information from many steps back.

Disadvantages

One notable disadvantage is that training RNNs is slow since it is sequential and cannot be parallelized. In practice, it is difficult to access information from many steps back due to problems like vanishing and exploding gradients, which we discuss later in this section.

7.2.2 Training RNNs

Backpropagation Through Time (BPTT)

Backpropagation through time (BPTT) is simply backpropagation applied to sequence models with a hidden state and thus it is used to train recurrent neural networks. We relegate the details of the implementation to here.

7.2.3 Vanishing and Exploding Gradients

Vanishing Gradients

In backpropagation, the gradients frequently become smaller and the new model weights end up being virtually identical to the old weights without any updates. As a result, the gradient descent algorithm never converges to the optimal solution. This is known as the problem of vanishing gradients.

Why does this happen? Vanishing gradients issue typically occur when using sigmoid or tanh activation functions in the hidden layer, which effectively compresses a large input space into a small output space. When the inputs become fairly small or fairly large, the derivatives become extremely close to zero and there is no gradient value to propagate back.

One obvious solution is to replace the activation function of the network by using ReLU instead of sigmoid. It keeps linearity for regions where sigmoid and tanh are saturated, thus responding better to gradient vanishing. Another solution is to consider a different architecture such as the LSTM, which we discuss later below. This post discusses other potential changes.

Exploding Gradients

If gradients get larger as the backpropagation progresses, we may end up with outsized weight updates, thereby leading to the divergence of the gradient descent algorithm.

Why does this happen? This problem happens because of weights and not because of the activation function. When the weight values are large, the derivatives will also be higher, thereby changing the weights significantly and preventing the gradient from converging.

A common solution is “gradient clipping” in which one may simply clip the parameter gradient element-wise before a parameter update or clip the norm of the gradient before a parameter update.

7.2.4 Applications

Part-of-speech tagging

The process of classifying words into their parts of speech and labeling them accordingly is known as part-of-speech tagging, or simply POS-tagging. This post provides an implementation using RNN.

Text Generation

Text generation is one obvious application of the RNN architecture. If we are training the neural network to predict the next character, it is called Character Level Model. Similarly, we can train the model to predict the next word, given a sequence of words called Word Level Models.

7.3 LSTM

Long Short-Term Memory RNNs (LSTMs) is a type of RNN proposed in 1997 as a solution to the vanishing gradients problem. As foreshadowed, its major strength is in capturing long-term dependencies in the sequence. This post provides a very detailed explanation for understanding LSTM networks. Here we provide the most essential details.

7.3.1 Basic LSTM Architecture

Recall that all RNNs have the form of a chain of repeating modules of neural network. In standard RNNs, this repeating module will have a very simple structure, such as a single tanh layer. LSTMs also have this chain like structure, but the repeating module has a different structure. Instead of having a single neural network layer, there are four, interacting in a very special way.

LSTM addresses the issues of the RNN by maintaining a cell state (c_t), which is the state at any given time. This cell state is updated at each time step, and the output hidden state is derived from the input (x_t), the previous hidden state (h_{t-1}), and the updated cell state (c_t).

To read, erase, and write from the cell, there are also three corresponding gates. First is the **forget gate**, which is the first orange box on the left. It takes in the previous hidden state, the input and the learned weights to produce a number between 0 and 1. The second is the **input gate**, which consists of the next two orange boxes in the diagram. The first sigmoid layer decides which values to update, and the next tanh layer creates a vector of candidate values that can be added to the states.

Whether or not the update indeed happens is determined the by the last **output gate**, which consists of the last two orange boxes. The first sigmoid layer decides what parts of the cell state we're going to output, and then the cell state is put through the tanh layer and multiplied by the output of the sigmoid gate, so that we only output the parts we decided to. Note that the cell state also needs to be updated from c_{t-1} to c_t . This is done at the horizontal arrow at the very top of the diagram.

7.3.2 Variations

One popular LSTM variant, introduced by Gers & Schmidhuber (2000), is adding “peephole connections.” This means that we let the gate layers look at the cell state. Another variant is the Gated Recurrent Unit (GRU), which combines the forget and input gates into a single “update gate.” It also merges the cell state and hidden state, and makes some other changes. The resulting model is simpler than standard LSTM models, and has been growing increasingly popular. As GRU exposes the complete memory unlike the LSTM and is simpler, it is easier to modify and faster to train.

7.4 Other Extensions

7.4.1 Bidirectional RNNs

It is possible to make predictions based on future words by having the RNN model read through the corpus backwards. Such bi-directional RNN therefore maintains two hidden layers, one for the left-to-right propagation and another for the right-to-left propagation.

Since bidirectional RNNs require access to the entire input sequence, they are not applicable to language modeling in which only the left context is available. BERT is one such system built on bidirectionality.

7.4.2 Multi-layer RNNs

One can stack RNNs to construct a multi-layer RNNs. In such system, the hidden states from RNN layer i are the inputs to RNN layer $i + 1$. This allows the network to compute more complex representations.

High-performing RNNs are often multi-layer, ranging from 2 to 4 layers. This is actually not as deep as convolutional or feed-forward networks. Transformer-based networks such as BERT are usually deeper like 12 or 24 layers.

Chapter 8

Attention Models and Transformers

Transformers are one particular form of the encoder-decoder architecture that uses only attention and does not use RNNs. Main applications are BERT (Bidirectional Encoder Representations from Transformers) and GPT (Generative Pre-Training Transformer). In this section, we first describe the attention mechanism and then the transformer architecture.

8.1 Attention

8.1.1 Basic Idea

In the context of language translation, the basic concept of attention is that each time the model predicts an output word, it only pays attention to some input words. Specifically, for each word in the output sentence, it will map the important and relevant words from the input sentence and assign higher weights to these words as shown in the figure below:

In the traditional Seq2Seq model, we discard all the intermediate states of the encoder and use only its final states (vector) to initialize the decoder. The central idea behind attention is not to throw away these intermediate encoder states but to utilize all the states.

8.1.2 Steps for Computing Attention

The general attention mechanism makes use of three main components: queries (Q), keys (K), and values (V). It performs the following computations.

1. For each query vector q , it is matched against a database of keys to compute a score value for each key. In other words, for each key k_i , the dot

product is computed: $e_{q,k_i} = q \cdot k_i$.

2. The scores, which are computed for all keys, are then passed through a softmax operation to generate the weights: $\alpha_{q,k_i} = \text{softmax}(e_{q,k_i})$.
3. The value vector v_{k_i} , which is paired with a corresponding key, is weighted by the previously computed weights: $\sum_i \alpha_{q,k_i} v_{k_i}$.

In the context of machine translation, each word in an input sentence would be attributed its own query, key and value vectors. There are three different weight matrices that generate these three vectors. There are also different types of attention mechanisms, which are detailed at this blog.

8.1.3 Self-Attention

Self-attention is equivalent to a generalized attention mechanism where the query, key, and values take the same input. This is the attention mechanism used in the transformer architecture. It is an attention mechanism relating different positions of a single sequence in order to compute a representation of the same sequence.

For clarity, suppose the input sequence is “Today is Monday.”

1. For each token in the input sequence, compute the **query**, **key**, and **value** vectors using the three weight matrices. If the embedding vector size is 5 and the vector size of Q, K, and V is 3, then the weight matrices have the dimension 5×3 .
2. Compute the attention score using the **query** and **key** vector by computing the dot product. For example, if the **query** word is “Today”, then one computes the dot product with key vectors for “Today”, “Is”, and “Monday.” This yields a vector of three numbers, since there are three tokens in the input sequence.
3. Using the softmax() function, compute the weights. This still has a vector of three numbers that now sum up to 1.
4. Multiply the weights to the **value** vector. Since the **value** vector has dimension 1×3 , the resulting output also has the same dimension.

In sum, when the sequence “Today is Monday” is inputted into the self-attention layer as three 5×1 vectors, the output is three 3×1 vectors. In short notation, this is often expressed as:

$$\text{Attention}(Q, K, V) = \text{Softmax}(QK^T)V$$

which summarizes the above steps in a simple form.

Scaled Dot-Product Attention

The Transformer implements a scaled dot-product attention. As the name suggests, it first computes a dot product for each query with all of the keys. It subsequently divides each result by $\sqrt{d_k}$ where d_k is the dimension of the key vector.

Multi-Head Attention

Multi-head attention simply means that the self-attention is applied repeatedly. Specifically, we not only have one but multiple sets of Query/Key/Value weight matrices, and each of these sets is randomly initialized. Then, after training, each set is used to project the input embeddings (or vectors from lower encoders/decoders) into a different representation subspace.

8.2 Transformers

8.2.1 Overview

In its simplest form, a transformer takes in a sentence to translate and outputs a translated sentence:

$$input \rightarrow transformer \rightarrow output$$

The transformer consists of an encoder and a decoder:

$$input \rightarrow encoder \rightarrow decoder \rightarrow output$$

The encoder and the decoder consist of layers:

- Notice that the sublayers are stacked linearly for both the encoder and the decoder, but the decoder takes an additional input: the output from the last encoder sub-layer.

Finally, each layer of the encoder and decoder take the following structure:

- The encoder layer computes the self-attention and then feeds it into the feed-forward network.
- The decoder layer is identical to the encoder layer with additional middle step: the encoder-decoder attention, which uses the information carried over from the last step of the encoder.

8.2.2 Step 1. Embedding

Before the input sentence can be fed into the architecture, it needs to be embedded via (1) word embedding and (2) positional encoding. Obviously, this embedding only happens in the bottom-most encoder. Typically, the word embedding is done through a pre-trained neural network instead of a simple one-hot encoding.

Positional encoding is also necessary because the transformer architecture does not include a default method for analyzing the order of words in the input sentence. In transformers, each position (index) is mapped to a vector, and hence the output of the positional encoding layer is a matrix where the i th row is a vector representing the i th token in the input sequence. This vector is then added to the original word embedding and fed into the very first encoder layer. (See this post for details on the construction of this matrix.)

8.2.3 Step 2. Self-Attention in Encoders

Self-attention is a layer that helps the encoder look at other words in the input sentence as it encodes a specific word. For example, suppose we are given the following input sentence: “Simon called Manav because he felt overwhelmed.” Self-attention allows the model to associate “he” with “Simon” and not “Manav.”

For each word embedding vector, there is a vector of the attention layer. But we require the entire input sequence (i.e. the list of words) for computing self-attention because (1) we need it to compute the weights (= normalized scores) and (2) we need to combine the weights with the value vectors, each of which is associated with each word input. This post describes the step-by-step process for computing self-attention very nicely, which is summarized below:

Step 1. Create Query, Key, and Value vectors

The first step is to create, for each word, a Query vector, a Key vector, and a Value vector. These vectors are created by multiplying the embedding by three matrices that trained during the training process. Note that typically these new vectors are smaller in dimension than the embedding vector.

Step 2. Compute the Score

For each word, the score is calculated by taking the dot product of the query vector with the key vector of the respective word. So for the word in position i , the first score would be the dot product of q_i and k_1 and the second score would be the dot product of q_i and k_2 .

Step 3. Convert the Scores to Weights

The scores are divided by \sqrt{d} where d is the dimension of the key vectors, which is intended to lead to more stable gradients. Then they are passed through a softmax operation so that the scores are converted to weights. The resulting softmax score determines how much each word will be expressed at this position.

Step 4. Construct the self attention layer

Each value vector is multiplied by the softmax score and then the weighted value vectors are summed to produce the output of the self-attention layer.

8.2.4 Step 3. Self-Attention in Decoders

Similarly as before, the decoder passes its input into a multi-head self-attention layer. Unlike the one in the encoder, it is only allowed to attend to earlier positions in the sequence. This is done by masking future positions.

8.2.5 Step 4. Encoder-Decoder Attention in Decoders

This additional layer works like self-attention except that it combines two sources of inputs: the self-attention layer below it as well as the output of the encoder stack. **Importantly, the output from the encoder stack**

is passed to the value and key parameters, while the output of the self-attention module is passed to the query parameter.

8.3 References

1. This post describes the mathematics behind the attention mechanism.
2. This post describes the Transformer architecture in a simplified form, and this post provides a nice illustration of how transformers work.

Chapter 9

Pretrained Models and Fine-Tuning

In this section, we provide an overview

9.1 BERT

9.2 GPT

Chapter 10

Other Useful Methods for Textual Analysis

10.1 Convolutional Neural Networks (CNNs)

10.2 Hidden Markov Models (HMMs)

APPLICATIONS IN ECON/FINANCE

Chapter 11

Sentiment Analysis

CODE SNIPPETS

Chapter 12

Data Scraping

Chapter 13

Data Cleaning

13.1 Word Tokenization

Tokenizers can easily become complex.

13.1.1 Implementing Tokenization

Several Python libraries implement tokenizers, each with its own advantages and disadvantages:

1. spaCy—Accurate , flexible, fast, Python
2. Stanford CoreNLP—More accurate, less flexible, fast, depends on Java 8
3. NLTK—Standard used by many NLP contests and comparisons, popular, Python

NLTK and Stanford CoreNLP have been around the longest and are the most widely used for comparison of NLP algorithms in academic papers.

13.1.1.1 N-Grams

An n-gram is a sequence containing up to n elements that have been extracted from a sequence of those elements, usually a string.

When a sequence of tokens is vectorized into a bag-of-words vector, it loses a lot of the meaning inherent in the order of those words. By extending your concept of a token to include multiword tokens, n-grams, your NLP pipeline can retain much of the meaning inherent in the order of words in your statements.

13.1.1.2 Techniques for Normalizing the Vocabulary

One can normalize the vocabulary so that tokens that mean similar things are combined into a single, normalized form. Doing so reduces the number of tokens you need to retain in your vocabulary and also improves the association of meaning across those different “spellings” of a token or n-gram in your corpus

Case Folding

Case folding is when you consolidate multiple “spellings” of a word that differ only in their capitalization. We can normalize the capitalization using list comprehension: `[x.lower() for x in tokens]`

Stemming

Another common vocabulary normalization technique is to eliminate the small meaning differences of pluralization or possessive endings of words, or even various verb forms. Stemming removes suffixes from words in an attempt to combine words with similar meanings together under their common stem. A stem isn’t required to be a properly spelled word, but merely a token, or label, representing several possible spellings of a word.

It’s important to note that stemming could greatly reduce the “precision” score for your search engine, because it might return many more irrelevant documents along with the relevant ones.

Two of the most popular stemming algorithms are the Porter and Snowball stemmers. The Porter stemmer is named for the computer scientist Martin Porter. Porter is also responsible for enhancing the Porter stemmer to create the Snowball stemmer.

```
from nltk.stem.porter
import PorterStemmer
stemmer = PorterStemmer()
''.join([stemmer.stem(w).strip("'") for w in "dish washer's washed dishes".split()]])
```

Lemmatization

If you have access to information about connections between the meanings of various words, you might be able to associate several words together even if their spelling is quite different. This more extensive normalization down to the semantic root of a word—its lemma—is called lemmatization.

Lemmatization is a potentially more accurate way to normalize a word than stemming or case normalization because it takes into account a word’s meaning. A lemmatizer uses a knowledge base of word synonyms and word endings to ensure that only words that mean similar things are consolidated into a single token.

So lemmatizers are better than stemmers for most applications. Stemmers are only really used in large-scale information retrieval applications (keyword

search).

```
nltk.download('wordnet')
from nltk.stem import WordNetLemmatizer
lemmatizer = WordNetLemmatizer()
lemmatizer.lemmatize("better", pos="a") # if POS is not specified, it assumes noun
```