

Computer Science Tripos - Part II Project

# Language Modelling for Text Prediction

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supervised by Dr Marek Rei & Dr Ekaterina Shutova

## Proforma

Name: Devan Kuleindiren College: Robinson College

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Project Originator: Devan Kuleindiren & Dr Marek Rei Supervisors: Dr Marek Rei & Dr Ekaterina Shutova

## Original Aims of the Project

The primary aim of the project was to implement and benchmark a variety of language models, comparing the quality of their predictions as well as the time and space that they consume. More specifically, I aimed to build an n-gram language model along with several smoothing techniques, and a variety of recurrent neural network-based language models. An additional aim was to investigate ways to improve the performance of existing language models on error-prone text.

## Work Completed

All of the project aims set out in the proposal have been met, resulting in a series of language model implementations and a generic benchmarking framework for comparing their performance. I have also proposed and evaluated a novel extension to an existing language model which improves its performance on error-prone text. Additionally, as an extension, I implemented a mobile keyboard on iOS that uses my language model implementations as a library.

## Special Difficulties

None.

## Declaration

I, Devan Kuleindiren of Robinson College, being	a candidate for Part II of the Computer
Science Tripos, hereby declare that this dissertat	on and the work described in it are my
own work, unaided except as may be specified be	elow, and that the dissertation does not
contain material that has already been used to $\epsilon$	ny substantial extent for a comparable
purpose.	
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# Chapter 1

## Introduction

My project investigates the performance of various language models in the context of text prediction. I started by implementing a series of well-established models and comparing their performance, before assessing the tradeoffs that occur when you attempt to apply them in a practical context, such as in a mobile keyboard. Finally, I proposed a novel extension to an existing model which aims to improve its performance on error-prone text.

## 1.1 Language Models

Language models (LMs) produce a probability distribution over a sequence of words, which can be used to estimate the relative likelihood of words or phrases occurring in various contexts. This predictive power is useful in a variety of applications. For example, in speech recognition, if the speech recogniser has estimated two candidate word sequences from an acoustic signal; 'it's not easy to wreck a nice beach' and 'it's not easy to recognise speech', then a language model can be used to determine that the second candidate is more probable than the first. Language models are also used in machine translation, handwriting recognition, part-of-speech tagging and information retrieval.

My project focuses on language modelling in the context of text prediction. That is, given a sequence of words  $w_1w_2...w_k = w_1^k$ , I want to estimate  $\mathbb{P}(w_{k+1}|w_1^k)$ . For instance, if a user has typed 'do you want to grab a', then a language model could be used to suggest probable next words such as 'coffee', 'drink' or 'bite', and these predictions could further be narrowed down as the user continues typing.

#### 1.2 Motivation

#### Benchmarking

Language models are central to a wide range of applications, but there are so many different ways of implementing them. Before using a language model in a particular context, it is important to understand how it performs in comparison to other methods available. In this project I focused in depth on the two most prominent types of language model: n-gram models and recurrent neural network-based models. n-gram models are typically coupled with smoothing techniques, which are explained in section 2.1. I investigated 5 different smoothing techniques and 3 different recurrent neural network architectures on a variety of datasets.

#### **Error-prone Text**

One problem with existing language models is that their next-word predictions tend to be less accurate when they are presented with error-prone text. This is not surprising, because they are only ever trained on sentences that do not contain any errors. Unfortunately, the assumption that humans will not make any mistakes when typing text is almost never valid. For this reason, I also investigated ways to narrow the gap in performance between language model predictions on error-prone text and language model predictions on error-free text.

#### 1.3 Related Work

Chelba et al. [1] from Google explore the performance of a variety of language models on a huge, one billion word dataset. Their work presents the limits of language modelling, when vast quantities of data and computational resources are available. Chen and Goodman [2] compare the performance of a series of smoothing techniques for n-gram models, and later use their results to propose an extension to Kneser-Ney smoothing [3] which is implemented in this project.

In recent years, there have been joint efforts from Ng et al. at CoNLL to improve and compare the performance of grammatical error correction [4] [5]. Language modelling on error-prone text, however, has an important distinction: A test time, a language model cannot use words ahead of its current position to make predictions. In other words, if a language model is predicting the probability of the next word in the middle of an error-prone sentence, then it can only use the first half of the sentence for making corrections and predictions.

# Chapter 2

# Preparation

My preparation consisted of thoroughly understanding n-gram and RNN-based language models, as well as planning how to tie them all together in an efficient implementation.

## 2.1 *n*-gram Models

This section describes n-gram language models and the various smoothing techniques implemented in this project.

#### 2.1.1 An Overview of *n*-gram Models

Language models are concerned with the task of computing  $\mathbb{P}(w_1^N)$ , the probability of a sequence of words  $w_1w_2...w_N = w_1^N$ , where  $w_i \in V$  and V is some predefined vocabulary<sup>1</sup>. By repeated application of the product rule, it follows that:

$$\mathbb{P}(w_1^N) = \prod_{i=1}^{N} \mathbb{P}(w_i | w_1^{i-1})$$

n-gram language models make the Markov assumption that  $w_i$  only depends on the previous (n-1) words. That is,  $\mathbb{P}(w_i|w_1^{i-1}) \approx \mathbb{P}(w_i|w_{i-n+1}^{i-1})$ :

$$\mathbb{P}(w_1^N) \approx \prod_{i=1}^N \mathbb{P}(w_i|w_{i-n+1}^{i-1})$$

Using the maximum likelihood estimation,  $\mathbb{P}(w_i|w_{i-n+1}^{i-1})$  can be estimated as follows:

$$\mathbb{P}(w_i|w_{i-n+1}^{i-1})_{MLE} = \frac{c(w_{i-n+1}^i)}{\sum_{w} c(w_{i-n+1}^{i-1}w)}$$

where c(W) denotes the number of times that the word sequence W was seen in the training set.

To a first approximation, the aforementioned n-gram language model provides reasonable results and is simple to compute. However, it does have one major issue: if, as an example, a 3-gram (trigram) model does not encounter the trigram 'the cat sat' in the data

<sup>&</sup>lt;sup>1</sup>The vocabulary is typically taken as all of the words that occur at least k times in the training set. k is typically around 2 or 3.

it is trained upon, then it will assign a probability of 0 to that word sequence. This is problematic, because 'the cat sat' and many other plausible sequences of words might not occur in the training data. In fact, there are  $|V|^n$  possible n-grams for a language model with vocabulary V, which is exponential in the value of n. This means that as the value of n is increased, the chances of encountering a given n-gram in the training data becomes exponentially less likely.

One way to get around this problem is to exponentially increase the size of the training set. This does, however, require significantly more memory and computation, and assumes that additional training data is available in the first place. An alternative solution is to adopt a technique called *smoothing*. The idea behind smoothing is to 'smooth' the probability distribution over the words in the vocabulary such that rare or unseen *n*-grams are given a non-zero probability. There are a variety of methods that achieve this. The ones which I have implemented are described in the next section.

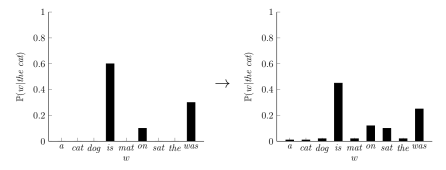


Figure 2.1: A toy example of smoothing. The probabilities of words that frequently follow 'the cat' are distributed to other less frequently occurring words in the vocabulary. The vocabulary is  $V = \{a, cat, dog, is, mat, on, sat, the, was\}$ .

## 2.1.2 Smoothing Techniques

#### Add-One Smoothing

Add-one smoothing [6] simply involves adding 1 to each of the n-gram counts, and dividing by |V| to ensure the probabilities sum to 1:

$$\mathbb{P}(w_i|w_{i-n+1}^{i-1})_{\text{ADD-ONE}} = \frac{c(w_{i-n+1}^i) + 1}{\sum_{w} c(w_{i-n+1}^{i-1}w) + |V|}$$

One issue with add-one smoothing is that it gives an equal amount of probability to all n-grams, regardless of how likely they actually are. As an example, if both 'the cat' and 'pizza cat' are unseen in the training data of a bigram model, then  $\mathbb{P}(cat \mid the)_{\text{ADD-ONE}} = \mathbb{P}(cat \mid pizza)_{\text{ADD-ONE}}$ , despite the fact that 'the' is much more likely to precede 'cat' than 'pizza'. This problem can be reduced by employing backoff, a technique whereby you recurse on the probability calculated by the (n-1)-gram model. In this case, it is likely that  $\mathbb{P}(the)_{\text{ADD-ONE}} > \mathbb{P}(pizza)_{\text{ADD-ONE}}$ , which could be used to deduce that 'the cat' is more likely than 'pizza cat'.

#### Absolute Discounting

Absolute discounting employs backoff by interpolating higher and lower order n-gram models. It does this by subtracting a fixed discount  $0 \le D \le 1$  from each non-zero count:

$$\mathbb{P}(w_i|w_{i-n+1}^{i-1})_{\text{ABS}} = \frac{max\{c(w_{i-n+1}^i) - D, 0\}}{\sum_{w} c(w_{i-n+1}^{i-1}w)} + \frac{D}{\sum_{w} c(w_{i-n+1}^{i-1}w)} N_{1+}(w_{i-n+1}^{i-1}\bullet) \mathbb{P}(w_i|w_{i-n+2}^{i-1})_{\text{ABS}}$$

where

$$N_{1+}(w_{i-n+1}^{i-1} \bullet) = |\{w \mid c(w_{i-n+1}^{i-1} w) \ge 1\}|$$

and the base case of recursion  $\mathbb{P}(w)_{\text{ABS}}$  is given by the maximum likelihood unigram model.  $N_{1+}(w_{i-n+1}^{i-1}\bullet)$  is the number of unique words that follow the sequence  $w_{i-n+1}^{i-1}$ , which is the number of n-grams that D is subtracted from. It is not difficult to show that the coefficient attached to the  $\mathbb{P}(w_i|w_{i-n+2}^{i-1})_{\text{ABS}}$  term ensures that the probabilities sum to 1.

Ney, Essen and Kneser [7] suggested setting D to the value:

$$D = \frac{n_1}{n_1 + 2n_2} \tag{2.1}$$

where  $n_1$  and  $n_2$  are the total number of n-grams with 1 and 2 counts respectively.

#### **Kneser-Ney Smoothing**

Kneser and Ney proposed an extension to absolute discounting which takes into account the number of unique words that precede a given n-gram [8]. As a motivating example, consider the bigram 'bottle cap'. If 'bottle cap' has never been seen in the training data, then the absolute discounting model would backoff onto the unigram distribution for 'cap'. Using the unigram distribution, 'Francisco' might be given a higher probability than 'cap' (assuming 'Francisco' occurs more frequently than 'cap'). This would result in the bigram 'bottle Francisco' being given a higher probability than 'bottle cap'. Clearly, this is undesirable, because 'Francisco' only ever follows 'San'.

From this example, it seems intuitive to assign more probability to those n-grams that follow a larger number of unique words. Kneser and Ney encapsulate this intuition by replacing some of the absolute counts  $c(w_i^j)$  with the number of unique words that precede the word sequence  $w_i^j$ ,  $N_{1+}(\bullet w_i^j)$ :

$$N_{1+}(\bullet w_i^j) = |\{w \mid c(w_i^j w) \ge 1\}|$$

Kneser-Ney smoothing  $^2$  is defined as follows:

$$\begin{split} \mathbb{P}(w_i|w_{i-n+1}^{i-1})_{\text{KN}} &= \frac{max\{\gamma(w_{i-n+1}^i) - D, 0\}}{\sum_{w} \gamma(w_{i-n+1}^{i-1}w)} \\ &+ \frac{D}{\sum_{w} \gamma(w_{i-n+1}^{i-1}w)} N_{1+}(w_{i-n+1}^{i-1}\bullet) \mathbb{P}(w_i|w_{i-n+2}^{i-1})_{\text{KN}} \end{split}$$

<sup>&</sup>lt;sup>2</sup>This is actually the interpolated version of Kneser-Ney smoothing, which differs slightly in form to the equation presented in the original paper.

where

$$\gamma(w_{i-k+1}^i) = \begin{cases} c(w_{i-k+1}^i) & \text{for the outermost level of recursion (i.e. } k = n) \\ N_{1+}(\bullet w_{i-k+1}^i) & \text{otherwise} \end{cases}$$
(2.2)

and the unigram probability is given as:

$$\mathbb{P}(w_i)_{\mathrm{KN}} = \frac{N_{1+}(\bullet w_i)}{\sum_{w} N_{1+}(\bullet w)}$$

#### Modified Kneser-Ney Smoothing

Chen and Goodman experimented with different discount values D in Kneser-Ney smoothing and noticed that the ideal average discount value for n-grams with one or two counts is substantially different from the ideal average discount for n-grams with higher counts. Upon this discovery, they introduced a modified version of Kneser-Ney smoothing [3]:

$$\mathbb{P}(w_i|w_{i-k+1}^{i-1})_{\text{MKN}} = \frac{\max\{\gamma(w_{i-k+1}^i) - D(c(w_{i-k+1}^i), 0\}}{\sum_{w} \gamma(w_{i-k+1}^{i-1}w)} + \lambda(w_{i-k+1}^{i-1})\mathbb{P}(w_i|w_{i-k+2}^{i-1})_{\text{MKN}}$$

where  $\gamma$  is defined in equation 2.2, and  $\lambda$  is defined as:

$$\lambda(w_{i-k+1}^{i-1}) = \frac{D_1 N_1(w_{i-k+1}^{i-1} \bullet) + D_2 N_2(w_{i-k+1}^{i-1} \bullet) + D_{3+} N_{3+}(w_{i-k+1}^{i-1} \bullet)}{\sum_{w} \gamma(w_{i-k+1}^{i-1} w)}$$

and

$$D(c) = \begin{cases} 0 & \text{if } c = 0\\ D_1 & \text{if } c = 1\\ D_2 & \text{if } c = 2\\ D_{3+} & \text{if } c \ge 3 \end{cases}$$

Chen and Goodman suggest setting the discounts to be:

$$D_1 = 1 - 2D\frac{n_2}{n_1}$$
  $D_2 = 2 - 3D\frac{n_3}{n_2}$   $D_{3+} = 3 - 4D\frac{n_4}{n_3}$ 

where D is as defined in equation 2.1.

#### Katz Smoothing

Katz smoothing is a popular smoothing technique based on the Good-Turing estimate [9]. The Good-Turing estimate states an n-gram that occurs r times should be treated as occurring  $r^*$  times, where:

$$r^* = (r+1)\frac{n_{r+1}}{n_r}$$

where  $n_r$  is the number of n-grams that occur r times. Converting this count into a probability simply involves normalising as follows:

$$\mathbb{P}(w_i|w_{i-n+1}^{i-1})_{\text{GT}} = \frac{c^*(w_{i-n+1}^i)}{\sum_{r=0}^{\infty} n_r r^*}$$
 (2.3)

Katz smoothing [10] is then defined as:

$$\mathbb{P}(w_i|w_{i-n+1}^{i-1})_{\text{KATZ}} = \begin{cases} \mathbb{P}(w_i|w_{i-n+1}^{i-1})_{\text{GT}} & \text{if } c(w_{i-n+1}^i) > 0\\ \alpha(w_{i-n+1}^{i-1})\mathbb{P}(w_i|w_{i-n+2}^{i-1})_{\text{KATZ}} & \text{otherwise} \end{cases}$$
(2.4)

where

$$\alpha(w_{i-n+1}^{i-1}) = \frac{1 - \sum_{\{w_i \mid c(w_{i-n+1}^i) > 0\}} \mathbb{P}(w_i | w_{i-n+1}^{i-1})_{\text{KATZ}}}{1 - \sum_{\{w_i \mid c(w_{i-n+1}^i) > 0\}} \mathbb{P}(w_i | w_{i-n+2}^{i-1})_{\text{KATZ}}}$$

In practice, the infinite sum in equation 2.3 cannot be computed. To get around this issue, Katz takes n-gram counts above some threshold k as reliable and only applies the Good-Turing estimate to those with a count less than or equal to k. Katz suggests k=5. This modification requires a slightly different equation to 2.4 and is presented in Katz's original paper.

#### 2.2 Recurrent Neural Network Models

In this section I give a brief introduction to neural networks, recurrent neural networks (RNNs) and how RNNs can be used in the context of language modelling. A thorough description of the more complex RNN architectures, gated recurrent units (GRUs) and long short-term memory (LSTM), is given in chapter 3.

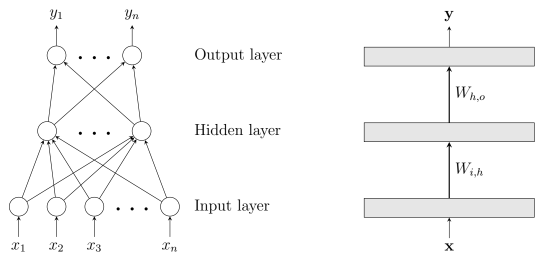
#### 2.2.1 An Overview of Neural Networks

The human brain is a furiously complicated organ, packed with a network of approximately 86 billion neurons<sup>3</sup> that propagate electrochemical signals across connections called synapses. Artificial neural networks, or neural networks, were originally developed as a mathematical model of the brain [12], which despite being substantially oversimplified, now provides an effective tool for classification and regression in modern-day machine learning.

Neural networks consist of a series of nodes which are joined by directed and weighted connections. Inputs are supplied to some subset of the nodes and then propagated along the weighted connections until they reach the designated output nodes. In the context of the brain, the nodes represent neurons, the weighted connections represent synapses and the flow of information represents electrochemical signals.

An important distinction to be made is whether the neural network is cyclic or not. Acyclic neural networks are called feed-forward neural networks (FNNs), whereas cyclic neural networks are denoted recurrent neural networks (RNNs) which are covered in section 2.2.2. There are a variety of FNNs, but the most prominent is the multilayer perceptron (MLP) [13], which I will outline below.

<sup>&</sup>lt;sup>3</sup>According to a study by Azevedo et al. [11].



- (a) An example multilayer perceptron.
- (b) A simplified notation for (a).

Figure 2.2: Two alternative notations for the same network: a multilayer perceptron with one hidden layer. In the simplified notation, the grey rectangle represents a vector of neurons, the thin arrow represents a vector and the thick arrow represents a matrix of weights.

#### The Multilayer Perceptron

The multilayer perceptron consists of layers of neurons, where each layer is fully connected to the next one. The first layer is the input layer, the last is the output layer and any layers in between are called *hidden layers*. If there is more than one hidden layer then it is called a *deep* neural network. The input neurons simply pass on the input values that they are given. The neurons in the subsequent layers of the network compute the weighted sum of their inputs, before passing that value through an *activation function* and outputting it:

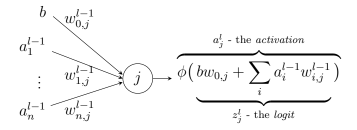


Figure 2.3: The computation carried out by each neuron. The outputs or *activations* of the neurons from the previous layer, (l-1), are denoted  $a_i^{l-1}$ , the weights on the input connections are  $w_{i,j}^{l-1}$ , the activation function is  $\phi$  and b is the bias input.

The unusual looking input b is a bias input. This is a fixed-value input, typically set to 1, which allows the neuron to shift the output of the activation function left and right by adjusting the weight  $w_0$ . Bias inputs are usually attached to every non-input neuron in the network.

It is also worth noting that activation functions should be chosen to be non-linear. Any

combination of linear operators is linear, which means that any linear MLP with multiple hidden layers is equivalent to an MLP with a single hidden layer. Non-linear neural networks, on the other hand, are more powerful, and can gain considerable performance by adding successive hidden layers to re-represent the input data at higher levels of abstraction [14] [15]. In fact, is has been shown that a non-linear MLP with a single hidden layer containing a sufficient number of neurons can approximate any continuous function on a compact input domain to arbitrary precision [16].

Frequently used activation functions include the sigmoid and the hyperbolic tangent functions. These are both non-linear functions squashed within the ranges (0,1) and (-1,1) respectively. Their steep slope at the origin is supposed to mimic an axon, which fires its output after the input reaches a certain potential. More importantly, these functions are differentiable, which allows for the network to be trained using gradient descent. This is discussed below.

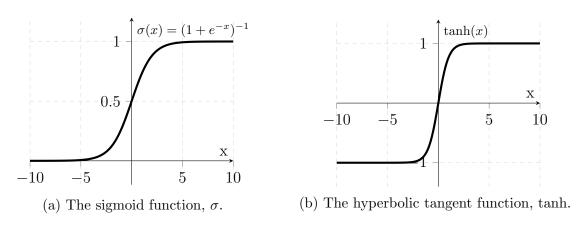


Figure 2.4: Two popular activation functions.

#### **Backpropagation and Gradient Descent**

FNNs compute a function, f, parameterised by the weights  $\mathbf{w}$  of the network, mapping an input vector  $\mathbf{x} = (x_1, ..., x_n)^T$  to an output vector  $\mathbf{y} = (y_1, ..., y_m)^T$ .

$$\mathbf{y} = f_{\mathbf{w}}(\mathbf{x})$$

The entire point of training a neural network is to get it to learn a particular mapping from input vectors to output vectors. What the inputs and outputs represent depends on the problem at hand. For example, in the context of classifying pictures of animals, the input vector might represent the pixel values of an image and the output vector might represent a probability distribution over the set of animals in the classification task.

In order to train a neural network, you supply it with a training set, which is just a list of input-target pairs:

$$\mathbf{s} = ((\mathbf{x}_1, \mathbf{t}_1), ..., (\mathbf{x}_N, \mathbf{t}_N))$$

Where  $\mathbf{t}_i$  is the vector that the network should output given the example input  $\mathbf{x}_k$ . A differentiable loss function,  $\mathcal{L}(\mathbf{y}, \mathbf{t})$ , is also defined, which essentially says how badly the

network output  $\mathbf{y}$  matches the target output  $\mathbf{t}$ . Then, a measure of how much error the network produces over the whole training set can be defined as follows:

$$\mathcal{L}_{ ext{total}} = \sum_{k=1}^{N} \mathcal{L}(f_{\mathbf{w}}(\mathbf{x}_k), \mathbf{t}_k)$$

The end product of the backpropagation algorithm is the partial derivative:

$$\frac{\partial}{\partial w_{i,j}^l} (\mathcal{L}_{ ext{total}})$$

for every weight  $w_{i,j}^l$  in the neural network. It is a two stage algorithm that works as follows:

- 1. Calculate,  $f_{\mathbf{w}}(\mathbf{x}_k)$  for each input example  $\mathbf{x}_k$ , and use those values to derive  $\mathcal{L}_{\text{TOTAL}}$ . The process of calculating the output of the neural network given the input is known as *forward propagation*, because the input is propagated through each layer in the network.
- 2. Given  $\mathcal{L}_{\text{TOTAL}}$ , calculate  $\frac{\partial}{\partial w_{i,j}^l} (\mathcal{L}_{\text{TOTAL}})$  for each weight in each layer of the network by applying the chain rule backwards from the output layer to the input layer. This step is described in more detail below.

 $\frac{\partial}{\partial w_{i\ j}^{l}}(\mathcal{L}_{ ext{TOTAL}})$  is calculated as follows:

$$\frac{\partial}{\partial w_{i,j}^{l}} \left( \mathcal{L}_{\text{TOTAL}} \right) = \sum_{k=1}^{N} \underbrace{\frac{\partial \mathcal{L}}{\partial f_{\mathbf{w}}(\mathbf{x}_{k})} \frac{\partial f_{\mathbf{w}}(\mathbf{x}_{k})}{\partial z_{j}^{l+1}}}_{\delta_{j}^{l+1}} \underbrace{\frac{\partial z_{j}^{l+1}}{\partial w_{i,j}^{l}}} = \delta_{j}^{l+1} a_{i}^{l}$$
(2.5)

The term  $\delta_j^{l+1}$  varies in form for each layer. For the output layer, it has the following form:

$$\delta_j^L = \frac{\partial \mathcal{L}}{\partial f_{\mathbf{w}}(\mathbf{x}_k)} \phi'(z_j^L) \tag{2.6}$$

For the preceding layers,  $\delta_j^l$  can be defined recursively in terms of the delta values for the neurons in subsequent layers in the network. This is defined below, and a detailed justification of it is given in appendix A:

$$\delta_i^l = \phi'(z_i^l) \sum_j \delta_j^{l+1} w_{i,j}^l \tag{2.7}$$

Gradient descent is an optimisation technique that uses the derivatives produced by the backpropagation algorithm to adjust the weights such that the loss is minimised over the training set. The simplest form of gradient descent does this by changing each weight value in the direction of the negative gradient of the loss. This is equivalent to moving the network output downwards on the error surface defined by plotting the loss against each of the weights in the network:

$$w_{i,j}^l \leftarrow w_{i,j}^l - \eta \frac{\partial}{\partial w_{i,j}^l} (\mathcal{L}_{\text{total}})$$

 $\eta$  is known as the *learning rate*, which determines how large the steps are in the negative direction of the gradient. Ways for setting  $\eta$ , and other useful techniques when training neural networks in practice are described in chapter 3.

#### 2.2.2 Recurrent Neural Networks

In the context of language modelling, the input is a sequence of words and the output should be the same sequence of words shifted ahead in time by one word. For example, given ('the', 'cat', 'sat', ...) as input, the network should produce something similar to ('cat', 'sat', 'on', ...) as output. The problem with FNNs is that these sequences may have a varying number of words in them, yet FNNs have fixed input and output vector sizes. One way to get around this problem is to look at a finite window of words at any given time, and encode that window into a vector, but even for modest window sizes this can lead to a huge number of weights in the network, which becomes more difficult to train. A better solution is to use a recurrent neural network.

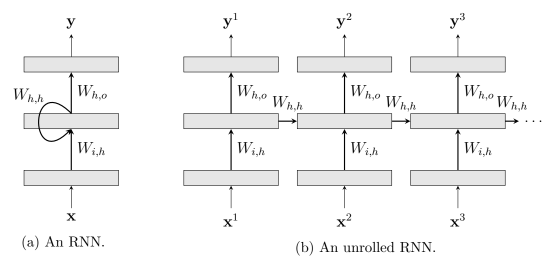


Figure 2.5: Two different representations of the same RNN.

A recurrent neural network can be constructed from an FNN by adding connections from each neuron in a hidden layer back to every neuron in that layer, including itself. This gives something like the network shown in figure 2.5a. An easier way to visualise an RNN is by *unrolling* it as shown in figure 2.5b. A nice property about RNNs is that they can be unrolled an arbitrary number of steps without the need for any additional parameters, because the same weight matrices are reused at each step. For this reason, RNNs are well suited for operating over sequences of input data. In the case of language modelling, this is ideal, because the inputs are sequences of words. The problem of representing textual words as numerical input vectors for an RNN is addressed in section 2.2.3.

As shown in figure 2.5b, RNNs can take a sequence of vectors as input. In most applications, the vectors in such sequences are typically ordered by time, so I use the notation  $\mathbf{x}^t$  and  $\mathbf{y}^t$  to denote the input and output vectors at time step t.

#### Backpropagation Through Time (BPTT)

Training an RNN is not so much different to training an FNN: the only difference is that gradients must also be calculated across time steps. As shown figure 2.5b, each weight is reused at each time step. For this reason, the weight update given in equation 2.5 is

summed across the time steps:

$$\frac{\partial}{\partial w_{i,j}^l} (\mathcal{L}_{\text{total}}) = \sum_t \delta_j^{l+1,t} a_i^{l,t}$$

The delta term for the output layer is defined analogously to equation 2.8:

$$\delta_j^{L,t} = \frac{\partial \mathcal{L}}{\partial f_{\mathbf{w}}(\mathbf{x}_k^t)} \phi'(z_j^{L,t})$$
(2.8)

The delta term for the preceding layers can be shown to have the form below. This is justified in more detail in appendix A.

$$\delta_i^{l,t} = \phi'(z_i^{l,t}) \left( \sum_j \delta_j^{l+1,t} w_{i,j}^l + \sum_j \delta_j^{l,t+1} v_{i,j}^l \right)$$
 (2.9)

where  $v_{i,j}^l$  denotes the weight on the recurrent connection from neuron i in layer l back to neuron j in the same layer.

#### Vanilla Recurrent Neural Networks

So far, the equations presented have assumed a vanilla RNN. That is, each neuron, or cell, simply computes the weighted sum of its inputs and then applys an activation function. A significant problem associated with this architecture is known as the vanishing gradient problem. Namely, the influence of a particular input on the hidden layers and subsequently the output layer either decays or blows up exponentially as it cycles through the recurrent connections, which makes it difficult for this architecture to retain long-term information across several steps. Various improvements have been made to improve upon this problem, two of which are outlined below.

#### Long Short-Term Memory

The most effective solution so far for tackling the vanishing gradient problem is the Long Short-Term Memory architecture proposed by Hochreiter and Schmidhuber [17]. In their architecture, each cell maintains a state, **c**, which is manipulated according to the inputs it receives to produce a new state and the cell output. At each cell, the state passes through three gates: the *forget gate*, *input gate* and the *output gate*. The effect of such gates and a more detailed discussion of this architecture is given in chapter 3.

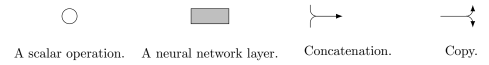


Figure 2.6: A notation for drawing RNN architectures.

#### Gated Recurrent Unit

Whilst the LSTM architecture provides a significant improvement for retaining long-term information, it is computationally more expensive to run. Cho et al. propose the Gated Recurrent Unit [18], which is a simplified version of LSTM that is cheaper to compute. The most significant changes are that it combines the forget and input gates into a single 'update gate', and it merges the cell state and cell output. This architecture is also described in more detail in chapter 3.

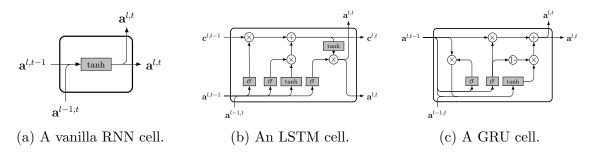


Figure 2.7: Various architectures of RNN neurons, using the notation from figure 2.6

#### 2.2.3 Word Embeddings

RNNs take a sequence of vectors as input, and produce a sequence of vectors as output. In the context of language modelling, these inputs and outputs should represent sequences of words. In order to do this, each word is represented by a single vector, which is known as a word *embedding*.

The aim behind word embeddings is that similar words should be represented by vectors that are closer together in the vector space than those for dissimilar words. One effective way to obtain such embeddings is to learn them through backpropagation, which is the method that was used in this dissertation. The idea is that each word is mapped to a large, randomly initialised vector. Then, when the RNN is trained on sequences of words, the embedding values are updated in a way that is analogous to how the weights are updated in the network.

## 2.3 Software Engineering

In this section details the requirements of the project and the early design decisions that were made to target those goals.

## 2.3.1 Requirements

The success requirements set out in the project proposal were:

	Language models (LMs) using the following techniques are implemented:
	• n-gram LMs with various smoothing techniques.
R1	• A vanilla RNN-based LM.
	• An LSTM-based LM.
	Comprehensible and reliable comparisons between the various LM
R2	implementations and their combinations are made regarding their accuracy,
	speed and resource consumption during both training and inference.
R3	A simple console application is developed to demonstrate the capability of
пэ	the aforementioned language models in the context of next-word prediction.

The project proposal also suggested two possible extensions:

<b>E</b> 1	Explore possible extensions to existing language models to improve their performance on error-prone text.
<b>E2</b>	Build a mobile keyboard that exhibits the functionality of the language models.

All requirements and extensions depend on **R1**, so I implemented that first. Afterwards, I built **R2** and **R3** in parallel before moving onto **E1** and **E2**.

#### 2.3.2 Tools and Technologies Used

Below I describe and justify where necessary the tools and technologies that I selected to build my project.

#### Version Control and Build Tools

I hosted my project in a repository on GitHub, used Git for version control, and used Bazel for running builds and tests on my project.

#### Machine Learning

I chose to use TensorFlow, an open source machine learning library, to assist in the implementation of my recurrent neural networks. There are two key reasons for this:

- 1. TensorFlow supports automatic differentiation. That is, each tensor operation has a partial derivative associated with it, so when you want to apply the second stage of backpropagation, TensorFlow can automatically apply the chain rule through any sequence of tensor operations. Without this, I would have to hard-code new gradient calculations every time I change the structure of the network, which is tedious and error-prone.
- 2. TensorFlow supports GPU-accelerated matrix operations. Without this, training my recurrent neural networks would take significantly longer.

Of course, there are many other machine learning libraries that offer comparable features. The reason I picked TensorFlow in particular was because I was already familiar with its codebase from using and contributing to it during an internship at Google in 2016.

In order to train large scale models on NVIDIA GPUs, I used the University's High Performance Computing service.

#### Languages

At the time of writing my code, TensorFlow had both a C++ and a Python API. However, the C++ API only had support for loading and running inference on models, not for training them.

One of the possible extensions I set out in my project proposal was to implement a mobile keyboard that makes use of my language model implementations. Android and iOS both provide little or no support for applications written in Python, but they do support code written in C++ via the JNI and Objective-C++ respectively.

I used C++ to write the benchmarking framework and all n-gram language models, so that they could easily be used within a mobile application. As for the RNN-based language models, I wrote code to train and export them in Python, and then wrote some C++ classes for loading and running inference on them. This way, I could leverage the

power of TensorFlow's Python API whilst also supporting the use of such models in a mobile keyboard. In order to export and import trained language models across different languages, I used Google's protocol buffers, a language neutral mechanism for serialising structured data.

#### **Testing**

I used Google Test for C++ unit tests and the unittest package for testing in Python.

#### 2.3.3 Starting Point

My project codebase was written from scratch, with the assistance of the tools and libraries mentioned in section 2.3.2. Before the project, I had no understanding of language modelling. Apart from a basic knowledge of recurrent neural networks, I had to learn about language modelling, n-gram smoothing techniques, long short-term memory and gated recurrent units through personal reading.

Experience	Tools and Technologies			
Significant	C++, Python, Git, GitHub, TensorFlow			
Some	Bazel, iOS, Protocol Buffers			
None	Google Test, Objective-C++, SLURM			

Figure 2.8: A summary of my prior experience with the tools and technologies used in this project.

# Chapter 3

# Implementation

3.1	Develo	opment	Strategy
		~	,

- 3.1.1 Version Control and Build Tools
- 3.1.2 Testing Strategy
- 3.2 System Overview
- 3.2.1 Interface to Language Models
- 3.3 *n*-gram Models
- 3.3.1 Counting *n*-grams Efficiently
- 3.3.2 Precomputing Smoothing Coefficients
- 3.4 Recurrent Neural Network Models
- 3.4.1 TensorFlow
- 3.4.2 Long Short-Term Memory
- 3.4.3 Gated Recurrent Units
- 3.4.4 Word Embeddings

Show some cool embedding plots with PCA.

## 3.4.5 Parameter Tuning

Dropout, embedding, learning rate decay, momentum?, gradient clipping

- 3.5 Extending Models to Tackle Error-Prone Text
- 3.5.1 Preprocessing the CLC Dataset
- 3.5.2 Error Correction on Word Context
- 3.6 Mobile Keyboard
- 3.6.1 Updating Language Model Predictions On the Fly

# Chapter 4

## **Evaluation**

In this chapter, I will first describe the benchmarking framework that I built to evaluate the language models, before proceeding onto the results. The results section is threefold: firstly I present the performance of the existing language models that I implemented, secondly I focus on the tradeoffs faced when employing those models on a mobile device, and finally I display my findings in language modelling on error-prone text.

**TODO:** Include these too:

- Examples of generated sentences.
- Embedding visualisations or perhaps put this into the implementation?
- Visualisations of the neuron activations for each word in a sentence.
- The number of parameters included in the LM in the PTB table (like Google did).
- The training time of each LM.

## 4.1 Evaluation Methodology

#### 4.1.1 Metrics

In the context of text prediction, there are essentially two questions one might want to answer when evaluating a language model:

- 1. How accurately does the language model predict text?
- 2. How much resource, such as CPU or memory, does the language model consume?

In order to answer these questions, I implemented a generic benchmarking framework that can return a series of metrics that fall into one of the two aforementioned categories when given a language model. These metrics include perplexity, average-keys-saved, memory usage and average inference time. The first two are concerned with the accuracy of language models and the latter two relate to the resource usage. I also recorded how long it took to train each model.

#### Perplexity

Perplexity is the most widely-used metric for language models, and is therefore an essential one to include so that my results can be compared with those of other authors. Given a sequence of words  $w_1^N = w_1 w_2 ... w_N$  as test data, the perplexity PP of a language model L is defined as:

$$PP_L(w_1^N) = \sqrt[N]{\frac{1}{\mathbb{P}_L(w_1^N)}} = \sqrt[N]{\prod_{i=1}^N \frac{1}{\mathbb{P}_L(w_i|w_1^{i-1})}}$$
(4.1)

where  $\mathbb{P}_L(w_i|w_1^{i-1})$  is the probability computed by the language model L of the word  $w_i$  following the words  $w_1^{i-1}$ . The key point is that **lower values of perplexity indicate** better prediction accuracy for language models trained on a particular training set.

This somewhat arbitrary-looking formulation can be better understood from a touch of information theory. In information theory, the cross-entropy H(p,q) between a true probability distribution p and an estimate of that distribution q is defined as:<sup>1</sup>

$$H(p,q) = -\sum_{x} p(x) \log_2 q(x)$$

It can be shown that  $H(p,q) = H(p) + D_{KL}(p||q)$  where  $D_{KL}(p||q) \ge 0$  is the Kullback-Leibler distance between p and q. Generally speaking, the better an estimate q is of p, the lower H(p,q) will be, with a lower bound of H(p), the entropy of p.

The perplexity PP of a model q, with respect to the true distribution p it is attempting to estimate, is defined as:

$$PP = 2^{H(p,q)} \tag{4.2}$$

Language models assign probability distributions over sequences of words, and so it seems reasonable to use perplexity as a motivation for a measure of their performance. In the context of language modelling, however, we do not know what the underlying distribution of p is, so it is approximated with Monte Carlo estimation by taking samples of p (i.e. sequences of words from the test data) as follows:

$$PP_L(w_1^N) = 2^{-\frac{1}{N} \sum_i \log_2 \mathbb{P}_L(w_i|w_1^{i-1})}$$

With a little algebra, this can be rearranged to give equation (4.1).

One issue with perplexity is that it is undefined if  $\mathbb{P}_L(w_i|w_1^{i-1})$  is 0 at any point. To get around this in my implementation, I replaced probability values of 0 with the small constant 1e-9. Results that use this approximation are marked.

#### Average-Keys-Saved

It is typical for the top three next-word predictions to be displayed and updated as the user types in a mobile keyboard, as described in section **TODO**: XXX. Clearly, it is in the interest of the mobile keyboard developer to minimise the amount of typing a user

<sup>&</sup>lt;sup>1</sup>Note that H(p,q) is often also used to denote the joint entropy of p and q, which is a different concept.

has to do before the correct prediction is displayed. Average-keys-saved is based on this incentive, and is defined as the number of keys that the user would be saved from typing as a result of the correct next word appearing in the top three predictions, averaged over the number of characters in the test data.

As an example, if the user is typing science and the word science appears in the top three predictions after they have typed sc, then that would count as 5 characters being saved, averaging at  $\frac{5}{7}$  keys saved per character. Averaging over the number of characters in the test data ensures that the results are not biased by the data containing particularly long words, which are easier to save characters on.

#### Memory Usage

This is measured as the amount of physical memory in megabytes occupied by the process in which the language model under test is instantiated.

#### **Training Time**

The amount of time it took to train the language model.

#### Average Inference Time

This is measured as the amount of time in milliseconds that the language model takes to assign a probability to all of the words in its vocabulary given a sequence of words, averaged over a large number of sequences.

#### 4.1.2 Datasets

I used three datasets throughout the evaluation of my project:

#### Penn Tree Bank (PTB) Dataset

The Penn Tree Bank is a popular dataset for measuring the quality of language models, created from text from the Wall Street Journal. It has already been preprocessed such that numbers are replaced with N, rare words are replaced with <unk> and the text has been split up into one sentence per line. It has been split up into a training, validation and test set. The training set has 10,000 unique words and 887,521 words overall.

Given that the Penn Tree Bank is so widely adopted, I used it for all tests in which the size of the training data is fixed, and will refer to it as PTB.

#### One Billion Word (1BW) Benchmark

This is a much larger dataset produced by Google of approximately 1 billion words [1]. I used this dataset for tests in which the size of the training data is a variable under investigation, and will refer to it as 1BW.

#### Cambridge Learner Corpus (CLC)

The Cambridge Learner Corpus is a dataset of 1,244 exam scripts written by candidates sitting the Cambridge ESOL First Certificate in English (FCE) examination in 2000 and 2001 [19]. The original dataset contains the scripts annotated with corrections to all of the mistakes by the candidates. In this project I make use of a preprocessed version of the dataset, in which there is one file containing the error-free version of the exam scripts and there is another file containing the original exam scripts with their errors. These two files are aligned line by line. I used this dataset when exploring the performance of language models on error-prone text, and will refer to it as CLC.

#### 4.2 Results

#### 4.2.1 Existing Models

#### Smoothing techniques and the value of n in n-gram models

The first set of language models that I built were n-gram models, along with a series of smoothing techniques for improving their predictions on less frequent n-grams.

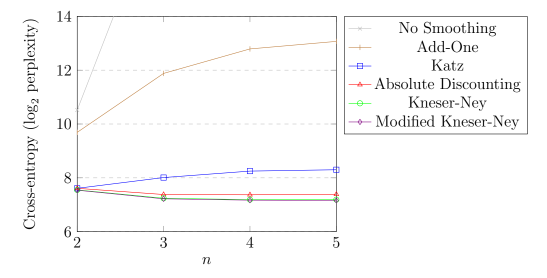


Figure 4.1: Cross-entropy of *n*-gram models trained on the PTB dataset.

Recall from equation (4.2) that cross-entropy is just the binary logarithm of perplexity, and that lower perplexity scores indicate better prediction accuracy. With this in mind, it is clear that modified Kneser-Ney smoothing offers the best prediction performance amongst the n-gram models.

The change in performance with the value of n is interesting. Intuitively, one might expect that increasing n will always yield better results, because this corresponds to increasing the number of words you use to make a prediction. However, for n-gram models with no smoothing, add-one smoothing or Katz smoothing, this is not the case. For n-gram models with add-one or no smoothing, this is because they do not employ backoff. At higher values of n, n-grams are much more sparse, so without any backoff higher n-gram models can only rely on sparse counts, resulting is lower probabilities being assigned to

plausible sequences of words. Katz smoothing does employ backoff, and achieves much better performance, but it still distributes too much probability to rare n-grams.

#### A comparison of RNN-based models with *n*-gram models

As described in the implementation chapter, I also implemented three RNN-based language models which differ in the RNN cell architecture: vanilla RNN, Gated Recurrent Unit and Long Short-Term Memory.

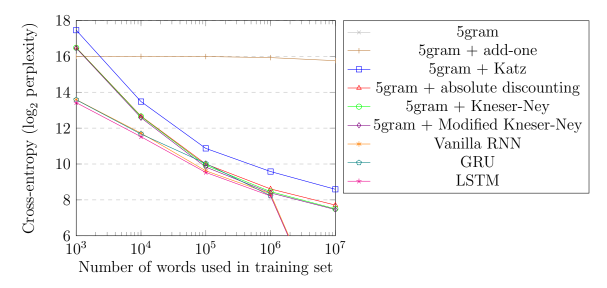


Figure 4.2: Cross-entropy of various language models with respect to the training set size, using the 1BW dataset.

As can be seen:

- RNNs better than *n*-gram models.
- etc, etc.

Some datasets are more predictable than others, which means that metrics like perplexity and average-keys-saved depend on which dataset is used. In order to produce metrics that are comparable with the work of other authors, I have tested my language models on the Penn Tree Bank dataset, as shown in figure 4.3. The perplexity was calculated over the whole of the test set, whereas average-keys-saved, a more expensive metric to compute, was taken over the first 1000 words of the test set. Indeed, my results for 3-gram modified Kneser-Ney smoothing, 5-gram modified Kneser-Ney smoothing and vanilla RNN are comparable to those from Mikolov [20].

There is a lot of information that can be drawn from figure 4.3, so I will highlight only the most important and interesting points:

• In general, the RNN-based language models outperform the *n*-gram based models in terms of prediction accuracy. In my implementation, they run slightly faster at inference time and also consume less memory, although this is partly due to the fact that they are entirely unpruned. The RNN-based language models take significantly longer to train and obtain the correct hyperparameters for.

Language Model	Perplexity	Average- Keys- Saved	Memory Usage (MB)	$egin{array}{c}  ext{Training} \  ext{Time} \  ext{(min,secs)}^\dagger \end{array}$	Average Inference Time (ms)
3-gram	$4.54 \times 10^{5*}$	0.35014	266.91	11s	62
3-gram $+$ add-one	3764.96	0.53063	266.94	11s	41
3-gram + Katz	256.95	0.68482	266.71	14s	88
3-gram $+$ absolute disc.	166.03	0.72178	266.78	13s	63
3-gram + KN	150.73	0.72466	266.88	14s	54
3-gram + modified KN	149.54	0.72355	266.97	14s	54
5-gram	$1.96 \times 10^{8*}$	0.07167	737.36	26s	130
5-gram $+$ add-one	8610.45	0.33886	737.30	26s	63
5-gram + Katz	314.49	0.67154	737.43	41s	156
5-gram + absolute disc.	167.38	0.72333	737.43	40s	126
5-gram + KN	146.35	0.72598	737.37	44s	114
5-gram + modified KN	142.68	0.72554	737.53	50s	116
Vanilla RNN	131.03	0.72776	253.67	15m 10s	39
Gated Recurrent Units	114.52	0.73993	271.39	$28m\ 35s$	37
Long Short-Term Memory	112.47	0.73617	287.13	18m 59s	38
LSTM, 5-gram + MKN (av)	96.07	0.75719	929.39	19m 49s	189
LSTM, 5-gram + MKN (int)	94.70	0.75830	927.20	19m 49s	190

<sup>\*</sup>These perplexity scores use the approximation mentioned in section 4.1.1 where 0-valued probabilities are replaced with a small constant to avoid division by 0. †The n-gram models were trained on my laptop, whereas the neural models were trained on a GPU cluster on the High Performance Computing Service. Nevertheless, the n-gram models were still much faster to train.

Figure 4.3: A benchmark of various language models on the PTB dataset.

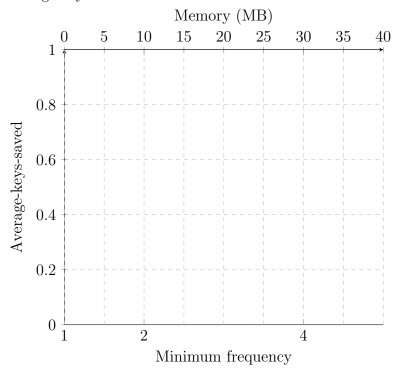
- Ignoring insignificant differences in memory and time, the discounting-based smoothing techniques (absolute discounting, Kneser-Ney smoothing and modified Kneser-Ney smoothing) equal or outperform add-one and Katz smoothing in every metric.
- LSTM yields marginally better prediction accuracy than the GRU architecture, but requires more memory. Apart from the memory overhead, GRUs significantly outperform the vanilla RNN architecture. The differences in training time for the RNN-based models should be taken lightly, because each model was trained for a different number of iterations, depending on how long it took to converge to a steady perplexity score on the validation set.
- Interestingly, a substantial improvement in perplexity and average-keys-saved can be obtained by simply averaging the probabilities produced by an *n*-gram and an RNN-based language model. This seems to imply that the two classes of language model complement one another, in the sense that RNN-based models make strong predictions when some of the *n*-gram predictions are weak, and vice versa.
- The combined language models can be improved even more by interpolating their probabilities rather than just averaging them. The result shown for the interpolation

of 5-gram modified Kneser-Ney with LSTM used  $\lambda = 0.38$ , where the probability was calculated as  $\lambda(n\text{-gram probability}) + (1 - \lambda)(\text{LSTM probability})$ .

#### 4.2.2 On a Mobile Device

Incorporating an RNN-based language model implementation into a mobile keyboard, as described in section **TODO**: XXX, presented a different series of challenges. The strict memory and CPU limitations of App Extensions in iOS forced me to explore the tradeoffs between resource consumption and prediction performance.

As shown in figure 4.3, the vanilla RNN architecture presents the smallest memory overhead, and so this is what I use in the following experiments. I investigated the effect of both vocabulary size and the number of hidden neurons on the memory usage and the average-keys-saved:



Here, I want to focus on the tradeoff between accuracy and resource consumption. Specifically, I could look at the following:

- The effect of increasing the minimum frequency for a word to be considered in the vocabulary. (I.e. the effect of changing the vocabulary size).
- The effect of changing the number of hidden layer neurons.
- The effect of using RNN vs GRU vs LSTM.
- (Perhaps also the effect of pruning on *n*-gram models).
- ALSO: Perhaps compare using float 32 vs float 16 in terms of memory vs accuracy.

The trade-off between the vocabulary size and resource usage

The trade-off between the number of hidden neurons and resource usage

The trade-off between the number of hidden neurons and resource usage

Other optimisations (not graphed):

- Remove softmax layer.
- Store weights as float 16 rather than float 32.

#### 4.2.3 On Error-Prone Text

I split the CLC dataset up into six files:

```
train.correct.txt train.incorrect.txt
valid.correct.txt valid.incorrect.txt
test.correct.txt test.incorrect.txt
```

The training, validation and test sets all consisted of one file containing uncorrected text and another file containing the corresponding corrected text. I trained an 2-layer LSTM-based language model with 256 hidden neurons on train.correct.txt, and used valid.correct.txt to guide the learning rate decay. I filtered the test set files such that they only contained pairs of lines that were not identical and that did not contain any insertion or deletion corrections.

Before exploring ways to improve the performance on error-prone text, I first established an approximate upper and lower bound. To obtain the upper bound (i.e. the best possible perplexity), I evaluated the LSTM using test.correct.txt for the input words and test.correct.txt for the target words. and 'In order to establish a rough upper and lower bound on the performance of the LSTM on error-prone text, I first evaluated Things I aim to evaluate here are:

- The hypothetical upper and lower bounds on accuracy (i.e. the LM results on correct input and on incorrect input respectively).
- The effect of using the vocabulary vs different sized dictionaries for determining if a word should be replaced or not.
- The effect of edit distance on performance.
- An intuitive explanation behind the gap remaining between the current performance and the upper bound. Perhaps some suggestions for future work.

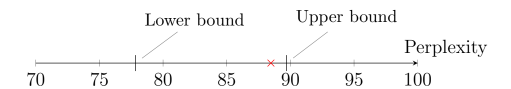


Figure 4.4: Perplexity on Cambridge Learner Corpus

# Chapter 5

# Conclusions

Conclusions:

•

Future work/possible extensions:

• Using Neural Machine Translation to correct the context, making use of attention.

•

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# Appendix A

# Backpropagation Recurrence Relation

$$\begin{split} \delta_i^l &= \sum_j \delta_j^{l+1} \frac{\partial z_j^{l+1}}{\partial z_i^l} \\ &= \sum_j \delta_j^{l+1} \frac{\partial z_j^{l+1}}{\partial y_i^l} \frac{\partial y_i^l}{\partial z_i^l} \\ &= \sum_j \delta_j^{l+1} w_{i,j}^l \phi'(z_i^l) \\ &= \phi'(z_i^l) \sum_j \delta_j^{l+1} w_{i,j}^l \end{split}$$

# Appendix B<br/> Project Proposal