

## IMPROVING THE COMPUTATIONAL EFFICIENCY OF TRAINING AND APPLICATION OF NEURAL LANGUAGE MODELS FOR AUTOMATIC SPEECH RECOGNITION

by

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A dissertation submitted to The Johns Hopkins University

in conformity with the requirements for the degree of

**Doctor of Philosophy** 

Baltimore, Maryland

October 2020

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### Abstract

A language model is a vital component of automatic speech recognition systems. In recent years, advancements in neural network technologies have brought vast improvement in various machine learning tasks, including language modeling. However, compared to the conventional backoff *n*-gram models, neural networks require much greater computation power and cannot completely replace the conventional methods.

In this work, we examine the pipeline of a typical *hybrid speech recognition system*. In a hybrid speech recognition system, the acoustic and language models are trained separately and used in conjunction. We propose ways to speed up the computation induced by the language model in various components. In the context of neural-network language modeling, we propose a new loss function that modifies the standard cross-entropy loss that trains the neural network to self-normalize, which we call a *linear loss*. The linear loss significantly reduces inference-time computation and allows us to use an importance-sampling based method in computing an unbiased estimator of the loss function during neural network training.

We conduct extensive experiments comparing the linear loss and several

commonly used self-normalizing loss functions and show linear loss's superiority. We also show that we can initialize with a well-trained language model trained with the cross-entropy loss and convert it into a self-normalizing linear loss system with minimal training. The trained system preserves the performance and also achieves the self-normalizing capability.

We refine the sampling procedure for commonly used sampling-based approaches. We propose using a sampling-without-replacement scheme, which improves the model performance and allows a more efficient algorithm to be used to minimize the sampling overhead. We propose a speed-up of the algorithm that significantly reduces the sampling run-time while not affecting performance. We demonstrate that using the sampling-without-replacement scheme consistently outperforms traditional sampling-with-replacement methods across multiple training loss functions for language models.

We also experiment with changing the sampling distribution for importancesampling by utilizing longer histories. For batched training, we propose a method to generate the sampling distribution by averaging the *n*-gram distributions of the whole batch. Experiments show that including longer histories for sampling can help improve the rate of convergence and enhance the trained model's performance. To reduce the computational overhead with sampling from higher-order *n*-grams, we propose a 2-stage sampling algorithm that only adds a small overhead compared to the commonly used unigram-based sampling schemes.

When applying a trained neural-network for lattice-rescoring for ASR, we

propose a pruning algorithm that runs much faster than the standard algorithm and improves ASR performances.

The methods proposed in this dissertation will make the application of neural language models in speech recognition significantly more computationally efficient. This allows researchers to apply larger and more sophisticated networks in their research and enable companies to provide better speech-based service to customers. Some of the methods proposed in this dissertation are not limited to neural language modeling and may facilitate neural network research in other fields.

**Primary reader**: Daniel Povey

Secondary reader: Sanjeev Khudanpur

Third reader: Philipp Koehn

### Acknowledgments

First of all, I would want to thank Dr. Daniel Povey, my primary advisor, for his close guidance in the past six years of my Ph.D. program. Also, for his relentless efforts in protecting the servers of Center for Language and Speech Processing at Johns Hopkins University. What happened to Dan that caused his firing from the university was unfair. While I hope things could be different, I am glad that he found employment opportunities with Xiaomi China. I am glad that now Dan can work with some of the best speech researchers globally and have complete freedom to improve speech recognition and help build products that reach hundreds of millions of users worldwide.

I also want to thank Prof. Sanjeev Khudanpur, my advisor, for his guidance in my research projects and for sharing his life wisdom in our frequent conversations. Sanjeev showed me how to be a better researcher and taught me how to be a better human being. I also owe a lot of gratitude to Sanjeev for doing an impeccable job maintaining my funding over the past years.

I am also grateful to Prof. Kai Yu for first taking me to the beautiful world of speech research; to Prof. Philipp Koehn, Prof. Shinji Watanabe, Dr. Guoguo Chen, for their guidance on research projects at JHU. I also owe much gratitude to Dr. Bhuvana Ramabhadran, my current manager at Google, for her guidance in my work and also being supportive while I prepared for my defense and wrote my thesis while working with her.

I also want to express my utmost gratitude to Dr. Leslie Leathers, Dr. Scott Spier, Dr. Michael Lint, and Dr. Alex Szablya, who at different times were my therapist. Working on a Ph.D. takes many tolls on a person's mental health, and I would not have been able to make it if it were not for the counseling sessions I have had with you.

I also want to thank Dr. Jordan Peterson. I know Dr. Peterson is a controversial figure, and I cannot entirely agree with everything he said. But I benefit immensely from his advice to "compare yourself to who you were yesterday, not to who someone else is today", as well as the most straightforward advice to just "clean your room before you criticize the world". Personal development is an endless pursuit, and I am glad that I have taken Dr. Peterson's advice to do my best to work on myself, and I will try to keep doing that going forward.

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## Chapter 1

## Introduction

#### 1.1 The Speech Recognition Problem

Speech is one of the, if not the most, natural way[s] for people to communicate with each other and to convey information. In this information age, there is a lot of demand for computers to "understand" human speech, and researchers have been working for decades trying to solve this task. While the word "understand" is hard to define, few would argue that to extract the text information from the speech is a vital first step before any understanding is possible. This defines the task of *automatic speech recognition* (ASR) [1], i.e., of designing a system that can accurately transcribe a representation of audio (for example, waveform or MP3 audio that we could play on a computer, or extracted acoustic features including Mel-frequency cepstral coefficients [2] (MFCC) or Perceptual linear predictive [3] (PLP), etc.) into text. While the ability to recognize speech

comes naturally to most humans, it is no easy task for machines. Although researchers have come up with different speech processing models and made significant progress in ASR [4, 5, 6] for the last five decades, speech recognition remains a challenging problem, especially in noisy environments and/or with spontaneous/accented speech.

The basic principles of solving the speech recognition problem have experienced many shifts in the past decades. In the earliest speech recognition systems, researchers built a "template" for each word in the vocabulary and used *dynamic time warping* [7] or similar methods for computing a "distance" between the template word and audio input to be recognized. This method works well for *isolated word-recognition*, e.g., recognizing single word commands. However, suppose an input sequence has multiple words. In that case, a separate stage of processing is needed to find boundaries between words before the speech recognition model recognizes the word from each segment. Overall, while there was progress, this class of methods did not make the speech recognition technology applicable to real-life uses.

With the introduction of the Dragon System [8], *Hidden Markov Models* (HMM) began their long success in the speech recognition community, up to this day. At the same time, the problem of speech recognition also became modular, where a typical system would comprise of several sub-components, including an acoustic model, a lexicon, a language model, and a decoder. In each of those components, researchers have worked on improving model per-

formance. For acoustic modeling, we have seen a shift from adopting Gaussian*mixture Models* (GMM), to sub-space GMM [9], to *Deep neural network* (DNN) based models [10]; we have also seen different training schemes proposed, including Maximum-likelihood Estimation (MLE) [11], Maximum A-Posteriori *Estimation* (MAP) [12] and discriminative training objectives [13], including Maximum Mutual Information (MMI) training [14] and Minimum Phone Error*rate* (MPE) training [15]. Some linguistic knowledge was also shown to help improve ASR performance, especially in earlier times; for example, expert knowledge of a phone set of a language was used in constructing questions for building phonetic decision trees [16] in order to cluster acoustic model units. Although gradually, the reliance on linguistic knowledge started to disappear. Rumor has it that the great speech scientist Fred Jelinek has said, Every time I fire a linguist, the performance of our speech recognition system goes up. Although there is a certain level of exaggeration in this quote, it does reflect the real trend, namely, that we are moving towards building a speech recognition system from data alone without external knowledge about languages. Even the lexicon (a pronunciation dictionary), which gives vital information on how words are pronounced, is sometimes unnecessary for many languages. Researchers have discovered that graphemic (letter-based) systems [17] can work surprisingly well, not only for languages like Spanish, which has a simple spelling to pronunciation mapping, but even for languages like English, whose spelling to pronunciation mapping is quite complicated and lacks structure. The recent

success seen in *end-to-end speech recognition* further proves this trend.

Several techniques, including model adaptation [18] and model combination [19], are shown to be useful in improving ASR performance, either in their generic forms or within the context of phonetic decision trees [20, 21]. With the incorporation of phonetic decision trees to classify context-dependent phones for generating modeling units, and the complexity of representing a phonetic dictionary and a language model, there was a high bar for researchers to implement a correct decoder for *large vocabulary continuous speech recognition* (LVCSR). Fortunately, those sub-components of ASR systems, in the last decade, have been elegantly unified in the *weight finite-state transducer* (WFST) framework [22, 23], which not only provides an elegant mathematical foundation of the methods but also makes the system more efficient and the algorithms easier to implement.

In the last ten years, as *neural machine translation* (NMT) models [24] have gradually surpassed the performance of traditional *statistical machine translation* (SMT) models [25], *end-to-end speech recognition* [26] has emerged. Since then, much of the work in end-to-end speech recognition has been inspired by similar work in machine translation, such as data cleaning [27], alternative representations of words [28] etc. However, although the end-to-end methods have caught up with or even surpassed the traditional hybrid systems on very large datasets, overall, it still cannot replace the traditional hybrid methods in terms of performance, the ease of performing domain adaptation [29], and interpretability [30].

#### **1.2** Mathematical Analysis of Speech Recognition

As of this moment, the most successful approach to tackle the speech recognition problem is through probabilistic and statistical modeling of speech, and that requires us to look at the speech recognition problem through a mathematical lens, which happens later in this section.

If we denote the speech observation as O and a word sequence as W, then the problem of speech recognition is to find the most "probable" word sequence  $W^*$  given the observation O, i.e.,

$$W^* = \underset{W}{\arg\max} P(W|O). \tag{1.1}$$

Now the problem becomes how to compute the term P(W|O). While it is possible to model the distribution P(W|O) directly, which is the foundation of end-to-end speech recognition techniques, the most successful approach, at least at the moment of writing, is to first break up the conditional probability using Bayes' Rule,

$$W^{*} = \arg \max_{W} P(W|O)$$
  
=  $\arg \max_{W} \frac{P(O|W)P(W)}{P(O)}$   
=  $\arg \max_{W} P(O|W)P(W).$  (1.2)

With this decomposition, an ASR system is now broken into two components,

- 1. an acoustic model which computes P(O|W).
- 2. a language mode that computes P(W).

While there has been much exciting work on acoustic modeling, this dissertation focuses on the language modeling part. However, a major part of the work proposed in this dissertation is generic enough to apply to any neural network, so some of the techniques proposed in this paper apply to other tasks, including acoustic modeling as well.

#### 1.3 Language Models

The task of language modeling is to design a mathematical system that can take the input W, which is a sequence of words, i.e.  $W = w_1, w_2, ..., w_n$ , and compute a score for it, for example, its probability P(W). Usually the joint probability itself is hard to estimate, and a common approach is to use the chain rule to break it into a product of conditional probabilities,

$$P(W) = P(w_1, ..., w_n)$$

$$= P(w_1)P(w_2|w_1)...P(w_n|w_1, ..., w_{n-1}).$$
(1.3)

This is the basic idea, and in practice people find that it helps to imagine there are always an implicit "begin-of-sentence" (<s>) and an "end-of-sentence" (</s>) word in each sentence, and then compute sentence probabilities as,

$$P(W) = P(\langle s \rangle, w_1, ..., w_n, \langle s \rangle)$$
  
=  $P(w_1 | \langle s \rangle) P(w_2 | \langle s \rangle, w_1) ... P(w_n | \langle s \rangle, w_1, ..., w_{n-1})$  (1.4)  
 $P(\langle s \rangle | \langle s \rangle, w_1, ..., w_n).$ 

Note we omit the  $P(\langle s \rangle)$  term because it's always  $1^1$ .

With this decomposition, it is now possible to use a counting-based method to estimate those conditional probability distributions from corpora. However, the longer the history is, the harder it is to get a reliable estimate for its distribution because of data sparsity issues. Researchers have come up with different techniques to alleviate this issue. While the actual methods differ, the fundamental idea is to map all possible (of which there are infinite) histories into a finite space to make estimations feasible. Mathematically, given a word

<sup>&</sup>lt;sup>1</sup>Since "<s>" is always the 1st word in any sentence in this representation.

w and a history h, it assumes

$$P(w|h) = P(w|f(h)).$$
 (1.5)

for some mapping function f(.), whose structure and output domain depend on the model assumption. We now briefly introduce the standard *n*-gram language models and then recurrent neural models, which are the focus of this dissertation.

#### **1.3.1** *n*-gram Language Models

An *n*-gram language model [31] is the most standard method in language modeling, which achieved significant success in the early stages of ASR research. An *n*-gram model only identifies the last n - 1 words in any history, for a pre-determined *n*. If we view an *n*-gram model in the context of Equation (1.5), the *f*(.) function for an *n*-gram model is defined as,

$$f(h) = f(w_1, ..., w_{t-1}) = w_{t-n+1}, ..., w_{t-1}.$$

Plugging this in the original equation we have,

$$P(w_t|w_1,...,w_{t-1}) = P(w_t|w_{t-n+1},...,w_{t-1}).$$
(1.6)

The *n*-gram assumption, along with some backoff/smoothing methods,

makes it possible to build a language model that gives a good performance in tasks like speech recognition and machine translation. To this day, although other forms of language models have far surpassed the performance of *n*-grams, it is still a vital component in most ASR systems. One of its major benefits is that since the target of the mapping function f(.) for any *n*-gram is finite-sized (there can be at most  $|V|^{n-1}$  unique histories for an *n*-gram, where *V* is the vocabulary of words), they can always be compiled into a *finite-state automaton*, which enables efficient processing that is vital for any ASR systems in production. Even without the advantage in terms of efficiency, researchers have observed that for a language model that outperforms an *n*-gram model, further performance gains could be seen if this model is combined<sup>2</sup> with a well-trained *n*-gram model [32].

The limitation of *n*-gram language models is evident in that they are not capable of learning long-term dependencies between words with distances larger than n - 1. For example, if one English speaker sees the following partial sentence: "I heard a joke earlier today and it was so…", she or he would usually have no problem in guessing correctly that the next word is probably "funny" or "hilarious", based on the word "joke" in history; however, any n-gram model with n < 7 would not be able to capture this dependency. This is due to the limitation of model capacity, and no amount of data could make the model learn this association.

<sup>&</sup>lt;sup>2</sup>For example, linearly or log-linearly interpolated.

#### **1.3.2** Neural-network Language Models

In the past decade, with the advent of greater computation power, neuralnetworks have become popular and successful in solving various tasks, including language modeling [33]. In particular, because a language model needs to model variable length texts, a recurrent neural network (RNN) becomes a natural choice for this task. We have seen great success in RNN language models (RNNLM) [34], especially with the application of more sophisticated recurrent structures, e.g., Long-short Term Memory (LSTM) networks [35]. Note that while we acknowledge that in some literature, the term RNNLM only means "vanilla RNNLMs" where simple linear layers coupled with non-linear activation function are used in building the network, here we use the term in its broader sense, to mean any network that has a recurrence in the t dimension, including more sophisticated networks like LSTM, GRU, etc. Formally, an RNN is a neural-network where at time t, its hidden state  $s_t$  depends not only on its input at t (which we denote as  $w_t$  in the context of language modeling) but also its hidden state at the previous time step  $s_{t-1}$ . In order to compute P(w|h), an RNNLM maps the history h into a real vector of a fixed dimension *d* for some *d*, i.e.

$$f(h) \in \mathbb{R}^d$$

where *d* is the vector dimension. Again, f(.) here refers to the definition in Equation (1.5).

Let's assume that the RNN takes input w from an alphabet V, and the hidden state s is represented as a real vector with dimension d, i.e.  $s \in \mathbb{R}^d$ . Then an RNN is defined by two functions:

- a state-transition function  $\delta$ :  $(s, w) \rightarrow s$
- a function *f*: *s* → (0,1)<sup>|V|</sup> that converts a hidden representation to an output distribution.

When an RNN models a sequence, the two functions are computed alternately. At any time *t*, first, a new hidden representation  $s_t$  is computed based on the old hidden representation  $s_{t-1}$  and the new word  $w_t$ ,

$$s_t = \delta(s_{t-1}, w_t). \tag{1.7}$$

Then a distribution over all words is computed based on the current hidden state,

$$p(w|s_t) = f(s_t).$$
 (1.8)

To make the recursion well-defined, an RNN would also need to identify an *initial state*  $s_0$  for the initial computation. Researchers usually set  $s_0$  as an all-zero vector/tensor; an alternative is to make  $s_0$  part of the model parameter learned during training.

In the context of recurrent neural-network language models (RNNLM), the

*w* above represents words, and *h* represents the encoding of a "history". The model assumes,

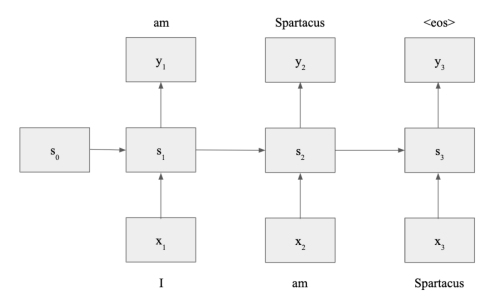
$$P(w_t|w_1, ..., w_{t-1}) = p(w_t|s_{t-1})$$

where

$$s_{t-1} = \delta(s_{t-2}, w_{t-1})$$
  
=  $\delta(\delta(s_{t-3}, w_{t-2}), w_{t-1})$   
= ...  
=  $\delta(\delta(...(\delta(\delta(s_0, w_1), w_2), ...), w_{t-2}), w_{t-1}).$  (1.9)

We see that the computation for any hidden state  $s_t$  would depend on **all** histories from  $w_1$  to  $w_t$ , and thus theoretically RNNLMs can encode infinite history information. Although in practice, RNNLMs rarely acquire the power of infinite memory, they have outperformed *n*-gram models significantly in various language modeling tasks, including language generation, speech recognition and machine translation.

In recent years, a special type of RNN, namely *long short-term memory* (LSTM) [36] based language models [35] have brought furthers gains in various language-related tasks. Compared to vanilla RNNLMs where both  $\delta$  and f functions are implemented as simple affine transforms, or stacks of affine transforms and non-linearity layers, an LSTM uses a gated mechanism that enables the system to learn long-term dependencies. A *Gated Recurrent Unit* (GRU) [37]



**Figure 1.1:** A Demonstration of a Simple RNNLM Working on Text Sequence "I am Spartacus".

is similar to LSTMs in using gated mechanisms but is less complicated and uses fewer parameters than LSTM.

Figure 1.1 shows how a simple 1-hidden-layer RNNLM works on the sentence "I am Spartacus". In this example, the hidden states of the RNNLM are represented in boxes with text s, and the arrows represent functional dependency. We see that state  $s_1$  depends on the initial state  $s_0$  and the input at t = 1, which is "I";  $s_1$  outputs a symbol "am", which is passed as input at t = 2 which alongside  $s_1$  generate  $s_2$ . This procedure repeats for the words in the sentence, until the last "end of sentence" symbol <eos> is generated.

#### 1.4 Application of Language Models in ASR

Suppose we already have a well-trained language model that computes P(W), and a well-trained acoustic model that computes P(W|O), we can use Equation (1.2) to compute for the most-likely word sequence W given any speech input O. However, it is impossible to apply Equation (1.2) directly in an ASR system, for it has to enumerate all possible sequences, of which there are infinitely many. Even when constraining the maximum lengths of hypotheses, using Equation (1.2) results in an algorithm that is not feasible in practice – we would need to iterate over all possible word sequences under the length constraint, which grows exponentially with the length. To solve this problem, an efficient *decoding* procedure is required.

The essence of the *decoding* procedure is to find the best hypotheses in a hypothesis set, represented as a graph, which could theoretically be infinite, and it is a challenging problem. If there are structures in the graph to exploit (for example, if the Markov assumption holds for the graph and/or the graph is finite), it is possible to make decoding more efficient. As an *n*-gram model can be compiled into a finite graph that satisfies the Markov assumption, it is a convenient choice for performing ASR decoding. In this case, an exact decoding process with an *n*-gram language model is equivalent to performing a *Viterbi Shortest Path* algorithm [38].

Because RNNLMs can theoretically encode infinite history, it is impossible

to compile a static decoding graph from an RNNLM. How to apply an RNNLM in speech recognition has been an ongoing topic in the speech community [39, 40] and is one of the primary focus of this dissertation.

#### 1.4.1 2-pass Method

Previously we mentioned that an *n*-gram may be compiled into a finite-state graph, and an efficient decoding procedure is therefore possible. The same thing, however, cannot be said about RNNLMs. While researchers have attempted to use RNNLMs for decoding directly, a more common approach is to adopt a 2-pass method or a *coarse-to-fine* method. In this method, we use an *n*-gram in the first pass decoding to generate a set of hypotheses, and in the second pass, refine the scores of hypotheses with an RNNLM.

The commonly used 2-pass methods for speech recognition utilizing RNNLMs differ in representing the hypothesis set and generally fall into two categories, (1) *n*-best rescoring and (2) lattice-rescoring.

#### **1.4.1.1** *n*-best Rescoring

In *n*-best rescoring, the hypothesis set is chosen to contain the *n* highest-scoring sentences for the input audio, where *n* is commonly chosen between 10 and 1000 in practice. The benefit of such methods is that it is relatively easier to implement, and also, for any sentence in the *n*-best list, the RNNLM can compute the exact score for it. However, since the hypothesis class represented

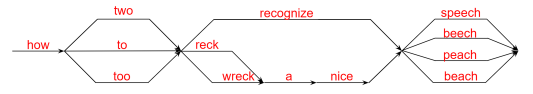


Figure 1.2: An Example of a Simple Word Lattice.

with n-best list has a simple structure with no sharing mechanism, it might not cover enough hypotheses unless a large enough n is chosen. For a small n, it might not contain enough hypotheses, which could hurt performance. However, if n is too large, it could be computationally very costly.

#### 1.4.1.2 Lattice Rescoring

In lattice-rescoring, the hypothesis set is represented in the form of a word lattice [41], where hypotheses with common prefixes can share their common structure, which enables more efficient computation and also more spaceefficiency in representing hypotheses.

An example of a word-lattice is shown in Figure 1.2. This lattice contains 7 states and 13 arcs, yet it contains 36 sentence hypotheses. However, in n-best rescoring, to rescore all hypotheses in this lattice, we need to run forward computation for 36 sentences separately, with minimal sentence-length being 4, and this inevitably adds a large overhead in computation.

In this dissertation, when we evaluate language models, both methods are used. The *n*-best rescoring method is mostly used for comparing different types of language models since it is easy to implement; we also focus on lattice-

rescoring later and propose ways to improve both the performance and the computational complexity of the algorithm.

#### **1.5** Evaluation of Language Models in ASR

Previously, we have mentioned that different language models achieve different "performance" in speech recognition tasks, but how do we evaluate a language model's performance?

A common measure for the quality of a language model is *perplexity* [42] (PPL). Given a language model m, and a corpus text T consisting of sentences  $[t_1, t_2, ..., t_n]$ , the perplexity of the text under the model is computed as,

$$perplexity = \exp(-\frac{\log(P_m(T))}{N}) = \exp(-\frac{\sum_i \log(P_m(t_i))}{N}), \quad (1.10)$$

where  $P_m(t)$  is the probability that the model *m* assigns to the sentence *t*, and *N* is the total length of all sentences in *T*, with the convention that all occurrences of the end-of-sentence symbol </s> are included in both  $P_m(t)$  and *N*.

By examining the perplexity computation, we note that it directly measures the text's likelihood under the model. A smaller perplexity indicates that the model assigns a larger probability to the text and is thus "better". While perplexity is a good measurement for language models, when we use a language model for speech recognition, the perplexity computation does not use any ASR system information. Thus it might not give us the complete picture of the language model's impact in speech recognition tasks.

When a language model is used in speech recognition, the direct method to measure its performance is through the ASR system's accuracy that incorporates the model. The most commonly used measure for speech recognition accuracy is *word error rates* (WER).

Say we run an ASR system on a dataset and generate a list of hypotheses  $[h_1, h_2, ..., h_n]$  where *n* is the number of utterances to recognize, and  $h_i$  is the hypothesis output of the ASR system for the *i*-th utterance. Let us represent the reference text as  $[r_1, ..., r_n]$ , then the WER of the ASR output is computed as,

WER = 
$$\frac{S+I+D}{N}$$
,

where *S*, *I*, *D* represent the number of *substitution errors*, *insertion errors* and *deletion errors*, respectively; *N* represent the total length of the reference sentences. The values of *S*, *I* and *D* are computed by performing a *Levenshtein Distance* [43] computation between the hypothesis and the reference text, at the word level, summed over all  $(h_i, r_i)$  pairs. Thus the numerator represents the minimal number of edits on words (possible edits include adding a word, deleting a word, and substituting one word with another) to change the hypothesis text to the reference text.

In the scenario above, the WER is computed between the one-best hypothesis for every utterance and the reference; we could generalize WER for use with lattices. If we have a set of hypotheses instead of just one, we could compute *oracle WER*, as the lowest possible WER we could get by comparing all hypotheses with the reference text. Depending on how we represent the hypothesis set, we could have *lattice oracle WER* and *n-best oracle WER*.

## **1.6 Limitations of RNNLMs in ASR**

In this section, we identify several issues of RNNLMs and their application in ASR.

#### 1.6.1 Training

Compared to traditional *n*-grams where the model parameters are computed by simple counting-based methods on the training data, a neural network model requires estimating its parameters from data, with many training iterations. Training a feed-forward neural language model even with fully paralleled computation is much slower than estimating an *n*-gram, let alone a recurrent neural network, whose training is harder to parallelize because of the sequential dependencies of the computation. Besides, to achieve better performance, we use more sophisticated networks (e.g., LSTM or GRU) instead of simple RNNs, which are more complex and take longer to run.

Compared to other tasks, the computational cost for neural networks is particularly more extensive for language modeling related tasks due to its vocabulary size of hundreds of thousands (or even millions) of words. In the input layer, having a large vocabulary size does not necessarily require more computation time since the computation could be implemented as selecting a row from the embedding matrix. However, in the output layer, full-sized matrix multiplication is inevitable, which is a bottleneck during the computation.

#### 1.6.2 Inference

After an RNNLM is well-trained, an ASR system uses it to score sentences, and the scores are combined with the acoustic model scores to recognize the input audio. We call this *model inference*. Usually, running inference with an RNNLM requires a full forward-propagation on the network, which is orders of magnitudes slower than an *n*-gram, where the inference "computation" is essentially a table-lookup.

Like the training, a large vocabulary is also a problem for RNNLM inference. The output-embedding is usually large, making the matrix-to-vector multiplication very costly.

It is worth mentioning that this costly computation with the output-embedding layer is rather unnecessary. In the 2-pass method, we only care about the scores for words that occur in the hypotheses for any given history in the hypothesis set. For example, let us take a look at the lattice shown in Figure 1.2. For the history "wreck a nice", scores are only needed for the words "speech" and "beach"; however, for a typical RNNLM, the score for any output word would depend on scores for all words, and this increases the complexities of the computation.

#### **1.6.3** Rescoring algorithms

As mentioned before, part of this work focuses on the 2-pass rescoring framework, where the hypothesis set is represented as a word-lattice. Because RNNLMs theoretically encode infinite history, a trivial rescoring algorithm on the lattice would expand it to an exponentially-sized tree, where each path from the root to a leaf is a sentence in the hypothesis set, whose size could be huge, making the computation infeasible.

An *n*-gram approximation method is usually adopted in practice to limit the search space, where the *n*-gram order is chosen to be 4 or 5. Under such a setup, two histories are merged into one if they share the last (n - 1) words.

However, even with the *n*-gram approximation method, the computational complexity still grows exponentially w.r.t. *n*, and that is why, in practice, an *n* larger than 5 is usually not quite computationally tractable. On top of that, using an *n*-gram approximation, we would be merging states representing different histories; thus, some of the following states' computation would be based on wrong histories, thus computing inaccurate scores for some of the hypotheses. Thus, this *n*-gram approximation method usually hurts performance [44] relative to the full exponential-complexity computation.

#### 1.6.4 Contribution of this Dissertation

In the previous section, we identified several computational issues regarding neural language model training, inference, and applications in speech recognition. Those are the issues that we work on in this dissertation. The dissertation makes the following contributions.

- 1. We provide an alternative loss function to cross-entropy loss, which we call *linear loss*.
  - (a) In terms of modeling performance, we show that linear loss either outperforms or is on-par with cross-entropy.
  - (b) Linear loss trains the model to self-normalize. It also outperforms the commonly used self-normalizing *noise contrastive estimation* (NCE) loss.

The self-normalization property brings significant speed up for *model inference*.

- 2. We propose using importance-sampling for linear loss training, which significantly speeds up *model training*.
- 3. We show that it is easy to "convert" a well-trained cross-entropy model to a self-normalizing model, with just one epoch of training with linear loss.

- 4. We propose an efficient method to perform lattice rescoring with neural language models for speech recognition.
  - (a) The method significantly speeds up the rescoring procedure, *and* it outperform the standard method in terms of speech recognition performance.

The methods proposed in this paper could be easily applied to ASR systems working on smartphones, tablets, and other smart devices with a voice interface and could improve the quality of the service they provide and drive down the computation costs. This page was left intentionally blank.

# PART I:

# IMPROVING THE COMPUTATIONAL

# EFFICIENCY OF RNNLMS

## Part I Outline

In Part I of this thesis, we focus on improving the computational efficiency of RNNLMs. We focus on both training and inference of RNNLMs and propose methods to make the computation more efficient for both of those scenarios. Part I spans from Chapter 2 to Chapter 6, and is structured as follows: we first introduce a new loss function for RNNLM training in Chapter 2, which we call linear loss; in Chapter 3, we propose an importance-sampling based training scheme that works in combination with the linear loss; we compare the performance of linear loss with common methods in Chapter 4; in Chapter 5, we conduct a comprehensive study on the choice of sampling algorithms used, investigating their impact on training speed as well as model performance. Furthermore, we study the effect of including longer histories in the sampling distribution in Chapter 6.

# Chapter 2

# Linear Loss: an Alternative to Cross-entropy Loss

This chapter first briefly introduces the standard cross-entropy loss and then introduces a new training loss for RNNLMs as an alternative to cross-entropy. We name the loss *linear loss* as it may be viewed as a linear-approximation of cross-entropy by applying Taylor expansion. We show how it is derived and its computational benefits compared to cross-entropy.

## 2.1 Cross-Entropy Loss Function

Cross-entropy is a standard loss function used in neural network training. When training a network by minimizing the cross-entropy between the output distribution predicted by the model, and the empirical distribution of the training data, the log-likelihood of the data under the model is maximized, or equivalently, the KL-divergence between the empirical data distribution and the model output distribution is minimized.

#### 2.1.1 Log-softmax Function

To understand cross-entropy, let us first take a look at some background in neural network modeling. Usually, a neural network model's output represents a probability distribution over *k* classes, where *k* is the dimension of the model output, and in language modeling, it would equal the size of the vocabulary. In order for the output to represent a proper probability distribution, normalization of the output is required. This is usually achieved by performing a *softmax* operation on the output. Let  $\sigma : \mathbb{R}^k \to \mathbb{R}^k$  represent the softmax function, as defined by

$$\sigma(z)_i = \frac{\exp(z_i)}{\sum_j \exp(z_j)}, \forall i \in \{1, 2, \dots, k\},$$
(2.1)

where  $z_i$ 's represent the "pre-softmax" output of the network, and  $\sigma()_i$  represents the *i*-th component of the softmax output.

In the actual implementation of the cross-entropy loss, *log-softmax* values are usually *stored* instead of the values of the softmax. The difference is that the log-softmax function would perform a per-element log function after the the softmax function, i.e.

$$\log\text{-softmax}(z) = \log \sigma(z), \tag{2.2}$$

where log-softmax(z)  $\in \mathbb{R}^k$ , and log(z)<sub>*i*</sub> is given by

$$\log(z)_i = \log(z_i), \forall i \in \{1, 2, ..., k\}.$$
(2.3)

By plugging in the definitions, we can see that

$$\log\operatorname{-softmax}(z)_i = z_i - \log\left(\sum_j \exp(z_j)\right), \forall i \in \{1, 2, ..., k\}.$$
(2.4)

#### 2.1.2 Cross-entropy Implementation

In training, the correct word after a history is sometimes referred to as the *gold label*, and could be represented as a one-hot vector  $v \in \mathbb{R}^k$ , where *k* is the vocabulary size, and

$$v_{i} = \begin{cases} 1, & \text{if } i \text{ is the index of the gold label,} \\ 0, & \text{otherwise.} \end{cases}$$
(2.5)

Given those definitions, the cross-entropy loss could be implemented as a negated dot-product between the one-hot vector representing the gold label and the network output, representing the different classes' log-probabilities. To ensure the output represents a valid probability distribution, we perform a log-softmax function as the last step of the network.

Mathematically, let *V* be the (output) vocabulary of an *n*-layer neural net-

work, *c* represent an input data point,  $\theta$  the current parameters,  $h_{n-1}(c;\theta) \in \mathbb{R}^d$ the hidden layer activations before the last affine layer of the network,  $A_n$  the last affine layer of dimension  $d \times |V|$ , and *w* the one-hot vector representation of the gold label corresponding to *c*. Then the cross-entropy loss for this one data point is the negated dot product,

$$-w \cdot \log \operatorname{softmax} \left( A_n(h_{n-1}(c;\theta)) \right).$$
(2.6)

In the context of language modeling, the linear component of  $A_n$ , which is of size  $d \times |V|$ , is usually referred to as a *word embedding* matrix, where each row of this matrix is a vector embedding of the correspond word. As the log-softmax function may be thought of as subtracting a scalar (acting as the normalization term), this setup may be understood as, firstly, computing *w* as the "predicted word embedding", then computing the dot product between *w* and all word embeddings, the log-probability for a certain word is the dot-product minus the normalization term.

Let us define  $\mathbf{y}(c; \theta) = A_n(h_{n-1}(c; \theta))$ , then the normalization term to be subtract is computed as

$$\log \sum_{i} \exp(y_i(c;\theta)), \tag{2.7}$$

and the computed log-probability of a word *w* is,

$$y_w(c;\theta) - \log \sum_i \exp(y_i(c;\theta)).$$
(2.8)

In cross-entropy training, the negated sum in Equation (2.8) over all (c, w) pairs in the data is used as the loss function that we want to minimize. If we view the data  $\mathcal{D}$  as a collection of (history context, correct word) pairs, then the cross-entropy loss on the whole dataset is computed as,

$$\mathcal{L}(\mathcal{D};\theta) = -\frac{1}{|\mathcal{D}|} \sum_{(c,w)\in\mathcal{D}} \left[ y_w(c;\theta) - \log\sum_i \exp(y_i(c;\theta)) \right].$$
(2.9)

## 2.2 Linear Loss

In this section, we propose a new loss function that takes a linear approximation of the standard cross-entropy loss, and analyze how the new function might impact training for neural networks.

Recall that for a continuously differentiable function f(x), we can use *Taylor expansion* at a point  $x_0$  to approximate it with a linear function, i.e.

$$f(x) \approx f(x_0) + g(x_0)(x - x_0),$$

where  $g(x) = \frac{df}{dx}$ 

We derive our *linear loss* by applying this linear approximation to the log function in cross-entropy. Note that,

$$\log x \approx \log(x_0) + \frac{1}{x_0}(x - x_0)$$
  
=  $\log(x_0) + \frac{x}{x_0} - 1.$  (2.10)

We also note that log is a convex function, and the linear approximation would always be smaller or equal to the function value, i.e.

$$\log x \le \log(x_0) + \frac{x}{x_0} - 1.$$
 (2.11)

In particular, when  $x_0 = 1$ , we have

$$-\log x \ge 1 - x, \tag{2.12}$$

with equality iff x = 1, as illustrated in Figure (2.1).

Now, we define

$$\mathcal{L}'(\mathcal{D};\theta) = -\frac{1}{|\mathcal{D}|} \sum_{(c,w)\in\mathcal{D}} \left[ y_w(c;\theta) + 1 - \sum_i \exp(y_i(c;\theta)) \right].$$
(2.13)

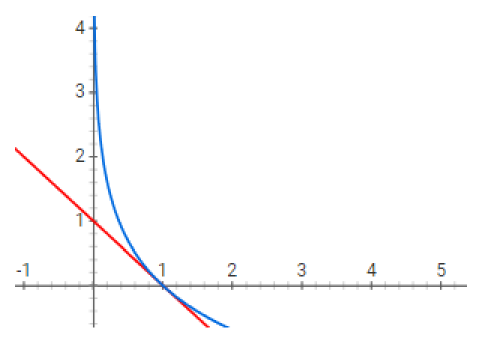
By this definition, we have

$$\mathcal{L}'(\mathcal{D};\theta) \ge \mathcal{L}(\mathcal{D};\theta), \forall \mathcal{D} \forall \theta,$$
(2.14)

with equality iff

$$\sum_{i} \exp(y_i(c;\theta) = 1, \forall c.$$
(2.15)

This inequality means that if we minimize the linear loss, the best it can do is to achieve the same value that we could get from cross-entropy loss, and at that point, Equation (2.15) would apply for the neural network output for all



**Figure 2.1:** Comparing -log(x) (blue) and 1 - x (red).

context in the training data<sup>1</sup>. In practice, though, due to a neural network's limited modeling capacity with finite parameters and details of optimization methods, we are not likely to reach that point. However, it is still reasonable to assume that a neural network's output should be reasonably close to being normalized when well trained under the linear loss.

## 2.3 Experiments

In this section, we empirically evaluate the proposed loss function and compare it with the cross-entropy loss. We implement the loss function with PyTorch [46], following the RNNLM example from [47]. We report both *perplexity* on development data as well as *word-error-rates* (WER) in ASR tasks, where

<sup>&</sup>lt;sup>1</sup>This could be easily proven if we assume that our neural network is indeed a universal approximator [45], and we have a perfect optimizer to do the training.

we include experiments in both hybrid speech recognition as well as end-toend speech recognition. As mentioned in the last section, we expect this loss function to train a network to self-normalize; therefore, we also report the normalization terms' mean and variance when running inference with the trained neural networks.

#### 2.3.1 Datasets

In the following sections, we report our numbers on two speech datasets, namely AMI [48, 49] and *Wall Street Journal* [50] (WSJ).

The AMI corpus consists of around 100 hours of meeting recordings. The language spoken in those meetings is English, where most speakers are not native speakers. The meetings were recorded in three rooms, each with different acoustic properties. The corpus has around 100,000 sentences and around 800,000 words (excluding the *end-of-sentence* symbol) in total, and the vocabulary size is 11,842 unique words. In this corpus, the most frequent five words are *the*, *yeah*, *uh*, *I* and *you*, with unigram probabilities of 0.0434, 0.0290, 0.0264, 0.0242 and 0.0219 respectively. Of all the 11,842 words in the vocabulary, 4387 appear only once in the corpus, and 1589 appear twice.

The Wall Street Journal (WSJ) corpus consists of around 284 hours of recording. The content comes from English news articles from the Wall Street Journal. The text consists of around 1,631,000 sentences and 37,000,000 words, and the vocabulary consists of 162,430 unique words. The most frequent five words are *the, of, to, a* and *and,* with unigram probabilities 0.0553, 0.0261, 0.0253, 0.0229 and 0.0222. Of all words in the vocabulary, 55,292 words appear only once in the corpus, and 20,837 appear twice.

#### 2.3.2 Language Modeling Performance

We train RNNLMs on the text corpus of the AMI dataset. We take its official training and development (dev) datasets for training and parameter tuning. We also randomly select a "training diagnostics" subset of 10000 sentences from the training set to report training perplexities. For the linear loss systems, we normalize the output to show valid perplexity numbers to make it comparable with cross-entropy systems. Three systems trained with the linear loss function are evaluated, where we choose different  $x_0$  values <sup>2</sup> to be 0.5, 1.0 and 2.0<sup>3</sup> in linearly approximating the log function. We also report the mean and variance of  $\sum \exp(y_i)$  for the unnormalized neural network outputs evaluated on datasets, as well as the ratio between standard deviation and mean<sup>4</sup>. We choose embedding-size to be 200 and use *Long-short term memory* (LSTM) for the network, and the number of layers is chosen to be two, with a dropout rate of 0.4. We report the results in Table 2.1. For each configuration, we select

<sup>&</sup>lt;sup>2</sup>Refer to Equation (2.11) on page 32 for its definition.

<sup>&</sup>lt;sup>3</sup>Those numbers are rather arbitrarily chosen and not tuned.

<sup>&</sup>lt;sup>4</sup>This ratio is a meaningful quantity to look at because, if the average mean of a network on a dataset is c, then we could trivially add a constant  $-\log(c)$  to every dimension of the bias parameter of the softmax layer and this would guarantee that the average mean is exactly one, to make this network "self-normalizing". However, the ratio between the standard deviation and the mean remains invariant during this operation, more accurately reflecting how well the network can normalize the output.

the model trained that gives the best (lowest) perplexities on the development set. For all systems, the batching is done by concatenating all sentences (the order of which is shuffled first to minimize between-sentence dependencies) before splitting into fixed-sized chunks, which we choose to be 35, and we use a batch-size of 64. For optimization, we use an Adam optimizer [51] with an initial learning rate of 0.001.

loss	Dataset	perplexity	mean	variance	stddev/mean
CE	train	50.78	0.2131	0.0196	0.6566
	dev	91.77	0.2209	0.0203	0.6451
Linear, $x_0 = 1.0$	train	49.56	1.0572	0.0326	0.1707
	dev	90.20	1.0623	0.0331	0.1713
Linear, $x_0 = 0.5$	train	48.26	0.5358	0.0080	0.1673
	dev	89.57	0.5419	0.0086	0.1711
Linear, $x_0 = 2.0$	train	49.95	2.1340	0.1274	0.1673
	dev	<b>89.21</b>	2.1441	0.1241	0.1643

**Table 2.1:** Model Stats on AMI Corpus Showing Perplexity, Mean, Variance and Ratio between Standard Deviation and Mean for the Normalization Term for the Output of Models Trained with Different Loss Functions.

Table 2.1 shows our experimental results, where we report perplexities of training and development set under different training schemes. By comparing the dev perplexities in the third column, we see that the linear loss function improves perplexity on the dev dataset compared to cross-entropy systems in all cases. We also see from the last column (stddev/mean) that using the linear loss would help reduce the variance of the sum of the exponential terms of the output. From the fourth column (mean), we see that it gives a very arbitrary mean in the cross-entropy system, while for the linear loss, the mean is quite

close to the value specified by  $x_0$ .

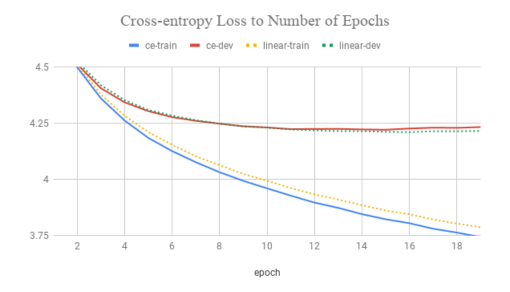
We also report perplexity results for the Wall Street Journal (WSJ) corpus. We use the standard script from the open-source toolkit Kaldi [52] repository to pre-process the corpus. We then take a random subset of 1000 sentences from this corpus as the development and the rest as the training set. We use a two-layered LSTM with dimension 800 and a dropout rate of 0.2 for training. All other parameters are kept the same as the AMI system described before.

system	train perplexity	dev perplexity
cross-entropy	62.96	84.12
linear, $x_0 = 1$	64.60	85.62
linear, $x_0 = 0.5$	65.55	85.98

Table 2.2: Perplexity of Models Trained with Different Loss Functions on WSJ Corpus

From Table 2.2, we see that using the linear loss achieves very similar perplexities compared with the baseline cross-entropy system. The numbers shown are slightly worse than cross-entropy for both training and development data. We show the impact of this on word-error-rates in ASR systems in later sections.

To investigate the convergence speed of training, we show the curve of train/dev loss during training at different epochs in Figure 2.2 with the AMI corpus, where we use the normalization constant  $x_0 = 1.0$  with linear loss. The curves for the linear loss system use dotted lines in order to highlight the difference. From the curves' overall trend, we see that the linear loss has



**Figure 2.2:** Comparison of Cross-entropy Loss to Number of Epochs for Models Trained with Different Loss Functions.

roughly the same convergence rate compared to the cross-entropy system. We also see that the linear loss achieves a better dev loss and a worse train loss than the cross-entropy system, which shows the linear loss's superiority compared to the cross-entropy loss.

#### 2.3.3 Initializing with Cross-entropy Systems

In the experiments reported in the previous section, RNNLMs are trained from scratch. As researchers widely use cross-entropy systems, it would be nice to easily "convert" a model trained with cross-entropy to one trained by linear loss. Here, we experiment in the framework of initializing our RNNLM model with a well-trained cross-entropy model and report results with continuing training with the linear loss for only one epoch. We take a well-trained crossentropy system for WSJ (the one we used to report in Table 2.2) to initialize the model and run only one epoch of training using the linear loss function. All parameters in the original network are updated during the training, and we use an Adam optimizer with an initial learning rate of 0.0001. We report the perplexities in Table 2.3.

system	ppl	mean	variance	stddev/mean
cross-entropy				108.6245
linear, $x_0 = 1$	85.62	1.1138	0.1888	0.5274

**Table 2.3:** Model Stats on WSJ Corpus Showing Perplexity, Mean, Variance and Ratio between Standard Deviation and Mean for the Normalization Term for the Output of Models Trained with Different Loss Functions. The Linear Loss System is Trained for One Epoch after Initializing with a Well-trained Cross-entropy System.

It may be seen that with the cross-entropy system, while it is giving good language modeling performance as measured by perplexity, the average mean is very large, and the variance is even larger, making normalization vital for the output to be interpreted as probabilities. With the linear loss training, even after one epoch, the system learns to normalize the output and significantly decrease the variance/mean ratio of the outputs.

#### 2.3.4 Hybrid Speech Recognition Performance

A practical benefit of a self-normalizing RNNLM is that it brings potential speed-up in the inference of an RNNLM. Note that while perplexity on development set is one crucial measurement for language modeling performance, it requires that the network output be normalized in order for the perplexity number to be valid, and thus the output of a self-normalizing network cannot be used to compute perplexity numbers. To investigate the effects of normalizing the outputs of RNNLMs, we compare their performance in speech recognition tasks.

#### 2.3.4.1 Speech Recognition Performance

We evaluate the WER on the AMI-SDM dataset when rescoring with different RNNLMs. We utilize the PyTorch RNNLMs by adopting the *n*-best rescoring method, where *n* is chosen to be 50. During rescoring, the RNNLM output is linearly interpolated with the original LM score, and the weight for the RNNLM scores is set as 0.8, while the original score has a weight of 0.2. For the linear loss systems, we choose  $x_0$  to be 1.0. We report both the "unnormalized" results where the neural network output is used without normalization, and "normalized" results where we force-normalize the output of RNNLMs. We evaluate on the AMI-SDM dataset and follow the standard script in Kaldi [52] to build a *Lattice-free MMI* [53] acoustic model for our evaluation. The results are shown in Table 2.4.

dev	eval
35.9	39.8
34.8	38.2
34.8 34.7	38.2 38.3
	35.9 34.8 34.8

Table 2.4: WER of AMI-SDM Corpus When Rescored by Different RNNLMs.

From Table 2.4, by comparing rows 1 and 2, we see that rescoring with a cross-entropy trained RNNLM significantly reduces WER; when comparing rows 3, 4, and 5, we see that the linear loss gives almost identical performance compared to the standard cross-entropy system, regardless whether normalization is performed on the neural-network outputs.

We also report the WER on the Wall Street Journal corpus in Table 2.5. From the table, we can see very similar trends in WER to those of the AMI corpus.

rescoring LM	dev93	eval92
no rescoring	4.8	3.4
cross-entropy	3.4	1.7
linear, unnormalized linear, normalized	3.2 3.3	1.3 1.3

Table 2.5: WER of WSJ Corpus When Rescored by Different RNNLMs.

Now we study if the specified normalization term ( $x_0$  in Equation (2.10)) impacts the language model performance. In Table 2.6, we report the WER performances on the AMI-SDM set of RNNLMs trained with different values of the specified normalization term, with or without normalization during inference. We adopt two types of normalization schemes, (i). exact normalization (denoted as exact), where we use a log-softmax function for the neural network output so that the neural network output could be interpreted as a valid probability distribution; (ii). approximate normalization (denoted as approx), where we add  $-\log(x_0)$  to the neural network output so that the sum of the "probabilities" is close to 1 (when  $x_0 = 1$ , this is equivalent to without normalization). The results are reported in Table 2.6. Again, we see no major difference when choosing a different normalization term, when either an exact or approximate normalization is performed; For the systems without any normalization, we notice that, interestingly, the system with specified  $x_0 = 0.5$  gives the best performance, and  $x_0 = 2.0$  gives the worst performance. Unfortunately, we do not see this trend in other datasets in later sections.

<i>x</i> <sub>0</sub>	normalization	dev	eval
no rescoring	n/a	35.9	39.8
cross-entropy	exact	34.8	38.2
1.0	no/approx	34.8	38.2
	exact	34.7	38.3
0.5	no	34.6	37.9
	approx	34.8	38.2
	exact	34.8	38.2
2.0	no	35.0	38.6
	approx	34.8	38.2
	exact	34.8	38.3

**Table 2.6:** WER in AMI-SDM When Rescored by Different RNNLMs with DifferentNormalization Schemes.

When we run rescoring experiments on the WSJ setup, we see some interesting results shown in Table 2.7. Remember previously in Table 2.2, we have seen slightly worse perplexities achieved using the linear loss than the cross-entropy one. However, from Table 2.7, we see that although their perplexity might be higher, language models trained using the linear loss consistently outperform the baseline cross-entropy systems, as similarly reported for the AMI dataset.

<i>x</i> <sub>0</sub>	normalization	dev93	eval92
no rescoring	n/a	4.8	2.7
cross-entropy	n/a	3.4	1.7
1.0	no/approx exact	3.2 3.3	1.3 1.3
0.5	no approx exact	3.1 3.1 3.0	1.6 1.5 1.5

**Table 2.7:** WER in WSJ When Rescored by Different RNNLMs with Different Normalization Schemes.

system	normalization	dev93	eval92
from scratch	no/approx	3.2	1.3
	exact	3.3	1.3
1 epoch from cross-entropy	no/approx	3.2	1.5
	exact	3.1	1.5

**Table 2.8:** WER in WSJ of Cross-entropy and Linear-loss Systems. The Linear loss System is Trained for One Epoch after Initializing with a Well-trained Cross-entropy Model.

#### 2.3.4.2 Speech Recognition Performance - One Epoch Training

Previously, we proposed converting a cross-entropy RNNLM to a self-normalizing one by training with the linear loss for one epoch and have shown that this achieves similar perplexities. Here we evaluate this training method in speech recognition tasks and show results in Table 2.8. Again, we use  $x_0 = 1$  and report the unnormalized results versus the normalized ones. We could see that, overall, the RNNLM trained from scratch performs similarly to the one trained with the linear loss for just one epoch when initialized from a cross-entropy

system. This is very encouraging and means that if researchers have already spent weeks or even months training a good cross-entropy language model, they could acquire a self-normalizing language model by quickly converting their old model. This new model retains the old model's performance but could run much faster in inference, as we show in the next section.

#### 2.3.4.3 Speed of RNNLM Computation

In this section, we report the inference speed for different methods. In Table 2.9, we compare the time used to rescore a subset of 10000 randomly selected sentences from the n-best list for all utterances with (i) the cross-entropy system and (ii) the linear loss system. We see that using the linear loss gives more than twice speed-up compared to the cross-entropy because there is no need to normalize the output.

	Time (seconds)	relative speed-up
cross-entropy	72.3	-
linear (unnormalized)	34.3	211%

**Table 2.9:** Time Spent Rescoring 10000 Randomly Selected Sentences from *n*-best Lists with Different RNNLMs on AMI Corpus. For Linear loss RNNLM, Normalization is Not Performed on the Output.

The AMI corpus has a small vocabulary (less than 12,000 words). We report the inference time of rescoring a random subset of 3000 sentences from all *n*-best sentences for the eval92 dataset of WSJ in Table 2.10. Note that the WSJ corpus's vocabulary size is around 162,000, which means a larger portion of the computation would be on the last softmax layer. From the results, we see that for such a model with an extensive vocabulary size, using our proposed linear loss and run unnormalized inference would save almost 97% of the time in inference computation. This is more similar to the real applications of RNNLMs used in companies like Google, Microsoft, and Apple. The vocabulary size could be hundreds of thousands or even millions, and this amount of run-time reduction would be very significant in reducing the actual costs of speech recognition systems.

	Time (seconds)	relative speed-up
cross-entropy	912.4	-
linear (unnormalized)	28.1	3247%

**Table 2.10:** Time Spent Rescoring 10000 Randomly Selected Sentences from *n*-best Lists with Different RNNLMs on WSJ Corpus. For Linear loss RNNLM, Normalization is Not Performed on the Output.

Combining the results here and previously shown tables, we conclude that the proposed linear loss improves the computational efficiency of RNNLMs in ASR tasks, which does not come at the cost of model performance.

#### 2.3.5 End-to-end Speech Recognition Performance

To further evaluate the proposed linear loss's effectiveness, we now turn to end-to-end speech recognition (E2E ASR). We use *Espresso* [54, 55] to carry out our experiments. We use the proposed linear loss in external language models used in E2E ASR in the context of *shallow-fusion* [56] for E2E ASR. We report on the Wall Street Journal corpus and compare it with shallow fusion results with the standard cross-entropy systems; for reference, we also report the results without language model fusion. We train a 2-layer LSTM language model with a hidden dimension 800. We use Adam optimizer and an initial learning rate of 0.001 and a dropout rate of 0.2. For decoding, we follow all the default parameters provided in the official Espresso release. The results are shown in Table 2.11. We can see that using the linear loss gives comparable WER in end-to-end ASR compared to the cross-entropy loss.

dev93	eval92
14.3	11.5
5.9 6.3	4.5 4.5
	5.9

**Table 2.11:** WER of Shallow Fusion with Different LMs for E2E ASR on WSJ Corpus. Normalization is not Performed on the Output of the Linear Model.

## 2.4 Chapter Summary

This chapter introduces a new loss function, which we call linear loss, for neural network training, and in particular neural network language modeling. From the experimental results, we see that linear loss trains the neural language model to self-normalize. We have presented a comprehensive comparison between the proposed loss and the widely used cross-entropy loss. Our experiments show that the linear loss gives an on-par performance as measured in the perplexity of development data and either outperforms or is on-par with the cross-entropy loss in both hybrid speech recognition systems and end-to-end speech recognition systems while being significantly faster. We have also shown that it is easy to "convert" a cross-entropy trained model to a self-normalizing one by training with the linear loss for just one epoch, saving much time. In particular, when rescoring the WSJ corpus, it brings a speed-up of more than 32X compared to the standard cross-entropy system while also giving better WER performance. This page was left intentionally blank.

# Chapter 3

# **RNNLM Training with Sampling**

In the previous chapter, we introduced a linear loss function that could replace the standard cross-entropy training for neural language models. We have shown that the linear loss function helps improve the inference speed of RNNLM computation and achieves comparable perplexities on held-out data and lower word error rates in speech recognition tasks.

This chapter shifts our focus to RNNLM training and proposes using an importance-sampling-based method to speed up training for RNNLMs. We use theoretical results as well as run evaluations to show the superiority of our method.

Note that in the current literature, the term "importance-sampling" is already used in the context of language modeling training to mean a particular method that adopts the importance-sampling technique to approximate the cross-entropy loss<sup>1</sup>. Here, we use the term in its broad sense in statistics and later apply it to compute the previously proposed linear loss. Readers are reminded not to be confused with those two different methods.

## 3.1 Importance-sampling

In statistics, importance-sampling is a general method to estimate specific properties of a distribution and a common such scenario is for computing summation of a discrete distribution or integration of a continuous distribution. In this work, we mainly focus on using importance-sampling for computing a summation term of discrete symbols. Say we want to compute the term,

$$S = \sum_{i} f(x_i). \tag{3.1}$$

The method to compute S exactly requires looping over all possible i's; however, we can define a random variable

.

$$y_{i} = \begin{cases} \frac{f(x_{i})}{p_{i}}, \text{ with probability } p_{i}, \\ 0, \text{ with probability } 1 - p_{i}. \end{cases}$$
(3.2)

If we compute the sum of all *y*'s, i.e.

$$S' = \sum_{i} y_i. \tag{3.3}$$

<sup>&</sup>lt;sup>1</sup>See Section 3.2.1 for details.

Note, now S' is also a random variable just like  $y_i$ 's. We see that,

$$\mathbb{E}[y'] = p_i \cdot \frac{f(x_i)}{p_i} + (1 - p_i) \cdot 0$$
  
=  $f(x_i).$  (3.4)

We can see that

$$\mathbb{E}[S'] = \mathbb{E}[\sum_{i} y_i] = \sum_{i} \mathbb{E}[y_i] = \sum_{i} x_i = S.$$
(3.5)

In this case, because  $\mathbb{E}[S'] = S$ , we say S' is an *unbiased estimator* for S.

One benefit of utilizing importance-sampling to compute the sum is that it could save much computation. In this case, depending on the choice of  $p_i$ 's, a large number of the *y*'s could be made 0, a novel scheme could be adopted that loops over only the non-zero terms to compute the summation term. We talk about that in the subsequent chapters.

# 3.2 RNNLM Training with Sampling-based Methods

#### 3.2.1 Importance-sampling for Cross-entropy Training

Importance-sampling has been used in the past for standard cross-entropy training of neural networks [57, 58]. Actually, the term "importance-sampling"

in the context of language model training usually refers to this use case. It is also referred to as "sampled-softmax" in parts of the literature [59, 60]. As mentioned in Chapter 2, in standard RNNLM training, we usually use the cross-entropy loss

$$\mathcal{L}(\mathcal{D};\theta) = -\sum_{(c,w)\in\mathcal{D}} \left[ y_w(c;\theta) - \log\sum_i \exp(y_i(c;\theta)) \right].$$
 (3.6)

Here it is possible to use importance-sampling methods to compute the term  $\sum_{i} \exp(y_i(c; \theta))$ . If we let

$$e_{i}'(c;\theta) = \begin{cases} \frac{\exp(y_{i}(c;\theta))}{p_{i}}, \text{ with probability } p_{i}, \\ 0, \text{ with probability } 1 - p_{i}, \end{cases}$$
(3.7)

then we can compute  $\mathcal{L}_s$  as an estimator for  $\mathcal{L}$ , i.e.

$$\mathcal{L}_{s}(\mathcal{D};\theta) = -\sum_{(c,w)\in\mathcal{D}} \left[ y_{w}(c;\theta) - \log\sum_{i} e_{i}'(c;\theta) \right].$$
(3.8)

However, even though we have the guarantee that y' is an unbiased estimator for y, i.e. that

$$\mathbb{E}\left[\sum_{i} e'_{i}(c;\theta)\right] = \sum_{i} \exp(y_{i}(c;\theta)), \tag{3.9}$$

note that the logarithm is a non-linear function, and

$$\mathbb{E}[\log\sum_{i} e_{i}'(c;\theta)] \neq \log \mathbb{E}[\sum_{i} e_{i}'(c;\theta)],$$
(3.10)

and therefore  $\mathcal{L}_s$  is not an unbiased estimator of  $\mathcal{L}$ :

$$\mathbb{E}[\mathcal{L}_{s}(\mathcal{D};\theta)] \neq \mathcal{L}(\mathcal{D};\theta).$$
(3.11)

The use of importance-sampling for computing the summation term in the standard cross-entropy training results in a biased estimator of the actual loss and this could potentially be a problem in practice during RNNLM training. Actually, since log is a convex function, by applying Jensen's Inequality we have

$$\mathbb{E}[\log\sum_{i} e_{i}'(c;\theta)] \le \log\sum_{i} \exp(y_{i}(c;\theta)),$$
(3.12)

and therefore,

$$\mathbb{E}[\mathcal{L}_s(\mathcal{D};\theta)] \le \mathcal{L}(\mathcal{D};\theta). \tag{3.13}$$

In other words, using importance-sampling inside the log function underestimates the true loss in practice.

#### 3.2.2 Importance-sampling for Linear Loss Training

In Chapter 2, we proposed a linear loss function,

$$\mathcal{L}'(\mathcal{D};\theta) = -\frac{1}{|C|} \sum_{(c,w)\in\mathcal{D}} \left[ y_w(c;\theta) + 1 - \sum_i \exp(y_i(c;\theta)) \right], \quad (3.14)$$

This new loss function, compared to the standard cross-entropy, does not have a log term, and is linear w.r.t the neural-network terms in the summation, and therefore, if we use importance-sampling techniques for computing the summation term  $\sum_{i} \exp(y_i(c; \theta))$ , and define

$$\mathcal{L}'_{s}(\mathcal{D};\theta) = -\frac{1}{|C|} \sum_{(c,w)\in\mathcal{D}} \left[ y_{w}(c;\theta) + 1 - \sum_{i} e'_{i}(c;\theta) \right], \quad (3.15)$$

then we have

$$\mathbb{E}[\mathcal{L}'_{s}(\mathcal{D};\theta)] = \mathcal{L}'(\mathcal{D};\theta).$$
(3.16)

This means using importance-sampling methods for computing the new loss yields an *unbiased estimator* for the actual loss. Note that this conclusion is reached without specifying the sampling-distribution p, which means we have the freedom to pick any p in practice, and the math would ensure the sample estimator's unbiasedness.

#### 3.2.3 Sampling Distributions

In the previous section, we have shown that when using importance-sampling for training neural networks with the linear loss, an unbiased estimator is always guaranteed without specifying the sampling distribution. A natural question readers might want to ask is, "Does this mean we could use any distribution for sampling? Does the sampling distribution have any impact on the estimator at all?" We answer that question in this section.

When we talk about an estimator being biased or unbiased, this limits us in only analyzing its *expectation*  $\mathbb{E}[\sum_{i} e'_{i}(c; \theta)]$ . Let's now consider another important aspect – the variance of the estimator,  $Var[\sum_{i} e'_{i}(c;\theta)]$ , which we want to minimize during training.

For simplicity of analysis, let's assume in  $\operatorname{Var}[\sum_{i} e'_{i}(c;\theta)]$ , all the terms in the summation are independent. This can be easily achieved if, during the sampling procedure, an independent decision is made whether a word *i* is picked in the sampled set, with probability  $p_{i}$ . Also, let's assume that the sum of all  $p_{i}$ 's is bounded, i.e.  $\sum_{i} p_{i} \leq c$  for c > 0, since without this constraint, we have a trivial solution to minimize the variance, by making  $p_{i} = 1, \forall i$ . In this case, the variance is 0 since we will be "sampling"/looping over all the words.

For a particular  $y'_i(c; \theta)$ , if we use the formula for Bernoulli distribution,

$$\operatorname{Var}[e_i'(c;\theta)] = p_i(1-p_i)\frac{\exp^2(y_i(c;\theta))}{p_i^2} = \frac{1-p_i}{p_i}\exp^2(y_i(c;\theta)), \quad (3.17)$$

then we have

$$\operatorname{Var}\left[\sum_{i} e_{i}'(c;\theta)\right] = \sum_{i} \operatorname{Var}\left[e_{i}'(c;\theta)\right]$$
$$= \sum_{i} \frac{1-p_{i}}{p_{i}} \exp^{2}(y_{i}(c;\theta))$$
$$= \sum_{i} \frac{\exp^{2}(y_{i}(c;\theta))}{p_{i}} - \sum_{i} \exp^{2}(y_{i}(c;\theta)).$$
(3.18)

A simple application of the Lagrange Multiplier method shows that the choice of  $p_i$  that minimizes the variance satisfies

$$p_i \propto \exp(y_i(c;\theta)),$$
 (3.19)

i.e., for any word *i*, we should set  $p_i$  to be proportional to the "true" distribution of words based on the current history. In practice, it usually suffices to set  $p_i$ proportional to the unigram probability distribution; for simplicity, we refer to this as *sampling from unigram*. Naturally, it will further minimize the variance if we utilize the history information in the context, for example, using a bigram or trigram distribution. However, so far, it is not apparent how to utilize any history when training models with batches of data since there could be multiple histories within the same batch. Hence, we first focus on experiments that sample from a unigram distribution in this chapter. We propose a method to leverage longer histories in Chapter 6.

#### 3.2.4 Noise-contrastive Estimation

*Noise-contrastive estimation* (NCE), although technically does not belong to the class of importance-sampling methods, is another commonly used method that utilizes sampling to speed up training, as well as trains RNNLMs to self-normalize. NCE works in different ways from the standard CE systems, which trains to maximize the likelihood of training text. Instead, NCE trains the system to differentiate between real data and generated noise (hence *noise-contrastive*).

NCE assumes that for any word history *h*, data could be generated from one of two distributions,

1. the true distribution  $P_{\text{TRUE}}(.|h)$  which the model would try to learn;

2. a noise distribution  $P_{\text{NOISE}}(.|h)$ ,

NCE trains the system to tell them apart. In training, for every word from the given training data (which represents the true distribution), the system samples k noisy words from the noise distribution  $P_{\text{NOISE}}(.|h)$ , so there are (1 + k) total words, only one of which is the "true" one. From any word w from all the (1 + k) choices, we could compute the posterior probability of "w is generated from the TRUE distribution", denoted as P(TRUE|w,h),

$$P(\text{TRUE}|w,h) = \frac{P_{\text{TRUE}}(w|h)}{P_{\text{TRUE}}(w|h) + kP_{\text{NOISE}}(w|h)},$$

and conversely, the probability of "w is generated from the noise distribution", denoted as P(NOISE|w,h),

$$P(\text{NOISE}|w,h) = \frac{kP_{\text{NOISE}}(w|h)}{P_{\text{TRUE}}(w|h) + kP_{\text{NOISE}}(w|h)}.$$

In training, we sample *k* words from the noise distribution for every correct word and maximize the joint-posterior probabilities (or equivalently, the sum of the log of those probabilities) of each word being in their correct class (TRUE vs. NOISE). The noise distribution is usually chosen to be the unigram distribution that is pre-computed from the training data. A sampling-with-replacement scheme is usually used in NCE.

Since there is no way to compute exactly  $P_{\text{TRUE}}(.|h)$ , we replace that with the probability output from RNNLM, i.e. using  $P_{\text{RNNLM}}(w|h)$  instead. However,

to compute  $P_{\text{RNNLM}}(w|h)$  is quite costly because it requires a normalization term to be computed; [61] pointed out that simply replacing the RNNLM output probabilities with RNNLM output "unnormalized scores" is enough to make training work. Furthermore, this would also train the RNNLM to self-normalize.

### 3.3 Importance-sampling for Variance-Regularization

The use of importance-sampling is not limited to any particular loss function, as we have shown with sampled softmax and the sampled version of linear loss. In this section, we turn to another commonly used self-normalizing loss function, namely the *Variance Regularization* (VR) loss [62].

Variance regularization (VR) [63, 62] is a common method in order to train an RNNLM to self-normalize. In some literature, it is also referred to as *selfnormalizing* methods [64]. This method's core idea is to add a quadratic penalty term to the standard cross-entropy loss to regulate the normalization term's growth. The mathematical definition of the loss function is shown in Equation (3.20).

$$\mathcal{L}_{vr}(\mathcal{D};\theta) = -\frac{1}{|\mathcal{D}|} \sum_{(c,w)\in\mathcal{D}} \left[ y_w(c;\theta) - \log\sum_i \exp(y_i(c;\theta)) - \lambda\left(\log\sum_i \exp(y_i(c;\theta))\right)^2 \right].$$
(3.20)

The difference between the VR loss and the standard cross-entropy is that it includes a third term of  $\lambda \left( \log \sum_{i} \exp(y_i(c;\theta)) \right)^2$ . The term  $\left( \log \sum_{i} \exp(y_i(c;\theta)) \right)^2$  would penalize the loss if the normalization term is either greater or less than 1 and  $\lambda$  is a hyper-parameter in order to weight the penalty term.

In the equation, the summation term  $\sum_{i} \exp(y_i(c;\theta))$  appear twice, for which we could use importance-sampling to compute. For ease of implementation and efficiency, the two terms could share the same samples for computation.

To the best of our knowledge, there are no attempts in literature in adopting sampling to compute the *Variance Regularization* (VR) loss [62]. One possible reason is that VR loss directly regulates the summation term, and thus using sampling to compute the sum might affect the interpretability of the method. Nevertheless, here we also propose to apply the sampling-based techniques to the VR loss. It could also be seen as combining the sampled softmax and VR.

### 3.4 Chapter Summary

This chapter gave an overview of sampling for training neural-network models and introduced some of the commonly used techniques, including noisecontrastive estimation and sampled softmax. We propose to apply importancesampling to training with linear loss, as well as the variance-regularization technique. We also present some theoretical analysis on deriving the sampling distribution that minimizes the variance of the estimator from the sampling procedure, laying the foundation for sampling distribution selection, which we investigate in detail in Chapter 6. In the next chapter, we will present our experiments comparing all those techniques in terms of model performance measured in perplexities on development datasets and speech recognition accuracy.

## Chapter 4

# **Evaluation of Linear Loss**

This chapter evaluates the linear loss and compares it with some of the commonly used method used loss functions for RNNLM training. Those methods that we compare against either utilize sampling-based techniques in order to speed up training and/or train the model to self normalize, including *noisecontrastive estimation, variance regularization* and *sampled softmax*. All experiments reported in this chapter use a PyTorch implementation of RNNLMs based on [47]. We implemented on top of the code different types of training loss functions, and we have uploaded all the changes to [65] for interested readers to reproduce those experiments. In all experiments, unless otherwise specified, we use a 2-layer LSTM model with a dropout rate of 0.4 for training a full-vocabulary RNNLM on the AMI corpus. For all the sampling-based experiments, we set the sampling distribution as the unigram distribution precomputed based on the training data and use a sampling-with-replacement<sup>1</sup> procedure unless otherwise specified. For the perplexity numbers, we take a random subset of 10000 sentences from the original "train" set of AMI as the development set to report perplexities and the rest of the training sentences.

We also evaluate different language models in speech recognition tasks. Instead of directly decoding with neural language models, we use the 2-pass *n*best rescoring method where in the first pass, we generate the top 50 hypotheses with a 3-gram language model, and in the second pass, we recompute the language model weights from RNNLMs for those hypotheses. The scores from RNNLMs are linearly interpolated with the 3-gram scores, with an RNNLM weight of 0.8. ASR evaluation is performed on the AMI-SDM testset, following the standard recipe provided by Kaldi. The acoustic model is a TDNN-LSTM [66, 67], and uses the *semi-orthogonal low-rank matrix factorization* [68] techniques during training; in compiling the initial decoding graph, we include explicit pronunciation and word-dependent silence probability modeling [69]. We use *exact lattice generation* [41] to generate lattices, from which we extract the *n*-best list.

loss	Dataset	perplexity	mean	variance	stddev/mean
	train	50.78	0.2131	0.0196	0.6566
CE	dev	91.77	0.2209	0.0203	0.6451
Lincor y 10	train	49.56	1.0572	0.0326	0.1707
Linear, $x_0 = 1.0$	dev	90.20	1.0623	0.0331	0.1713
$\lim_{x \to \infty} x = 0.5$	train	48.26	0.5358	0.0080	0.1673
Linear, $x_0 = 0.5$	dev	89.57	0.5419	0.0086	0.1711
Linear, $x_0 = 2.0$	train	49.95	2.1340	0.1274	0.1673
Linear, $x_0 = 2.0$	dev	89.21	2.1441	0.1241	0.1643
VR, $\lambda = 0.01$	train	47.81	0.6520	0.0806	0.4355
$VK, \Lambda = 0.01$	dev	90.34	0.6727	0.0811	0.4233
VR, $\lambda = 0.05$	train	48.15	0.9810	0.0914	0.3081
$V R, \Lambda = 0.03$	dev	90.35	0.9957	0.0903	0.3017
VR, $\lambda = 0.1$	train	48.29	1.0074	0.0714	0.2652
$V R, \Lambda = 0.1$	dev	90.16	1.0216	0.0705	0.2598
VR, $\lambda = 0.2$	train	48.68	1.0278	0.0504	0.2184
$V R, \Lambda = 0.2$	dev	90.34	1.0396	0.0501	0.2152
VR, $\lambda = 0.5$	train	49.32	1.0244	0.0290	0.1662
$V R, \Lambda = 0.5$	dev	90.21	1.0321	0.0288	0.1643
VR, $\lambda = 1.0$	train	48.97	1.0279	0.0180	0.1306
$V R, \Lambda = 1.0$	dev	90.01	1.0341	0.0114	0.1302
VR, $\lambda = 2.0$	train	48.22	1.0301	0.0110	0.1038
v n, n - 2.0	dev	89.87	1.0350	0.0114	0.1034
VR, $\lambda = 5.0$	train	49.22	1.0398	0.0064	0.0768
$\mathbf{v}\mathbf{R}, \Lambda = 0.0$	dev	90.13	1.0426	0.0065	0.0772
VR, $\lambda = 10.0$	train	51.33	1.0391	0.0029	0.0515
v IX, /1 — 10.0	dev	90.85	1.0411	0.0029	0.0521

**Table 4.1:** Model Stats on AMI Corpus Showing Perplexity, Mean, Variance and Ratio between Standard Deviation and Mean for the Normalization Term for the Output of Models Trained with Different Loss Functions.

### 4.1 Comparison with Variance Regularization

We compare linear loss with cross-entropy loss and variance regularization (VR) loss in Table 4.1, where we report perplexity, mean, variance, and standard deviation (stddev) to mean ratio on training (train) and development (dev) sets. For the variance regularization systems, we perform a sweep over the values for  $\lambda$  to control the weight of the penalty term; for the linear loss function, we report systems trained with  $x_0$  values 0.5, 1.0, and 2.0. From Table 4.1, we have the following observations:

- for reasonably-chosen hyper-parameters, the perplexity performance of both linear and variance regularization systems can surpass that of the cross-entropy system;
- both the linear loss and variance regularization loss are effective in limiting the normalization terms;
- 3. VR systems have an advantage of controlling the variance of the normalization term by controlling the *k* hyper-parameter, although this choice also affects the perplexity performance;
- 4. For well-chosen hyper-parameters, the linear systems can outperform the variance regularization systems as measured by the perplexity of development data.

<sup>&</sup>lt;sup>1</sup>This means we allow the same word to appear in the same sample set more than once. This is the easiest sampling method to implement and is the default setting when people use methods like NCE.

We report the ASR performance of variance regularization and other RNNLMs in Table 4.2. We see that although those systems give noticeably different performances measured in perplexity on dev data, when used for rescoring *n*-best lists for ASR systems, they give very similar performances.

rescoring LM	dev	eval
no rescoring	35.9	39.8
cross-entropy	34.8	38.2
linear	34.8	38.2
$VR, \lambda = 1.0$ $VR, \lambda = 2.0$ $VR, \lambda = 5.0$	34.8 34.8 34.8	38.2 38.2 38.4

 Table 4.2: Comparisons of WER% on AMI-SDM Corpus, Linear Loss VS Variance

 Regularization

### 4.2 Comparison with Noise-contrastive Estimation

We compare the linear loss with NCE in Table 4.3 on language modeling performance as measured by perplexity on the development data.

Note from Table 4.3, when using the same number of samples during training, the linear loss systems give better performance than NCE systems and start to outperform the cross-entropy baseline when sample sizes exceed 512; we notice that NCE would need  $\geq$  1024 samples in order to beat the cross-entropy baseline. In terms of enforcing the normalization term to be close to 1.0, we notice that NCE seems slightly better at pushing its average closer to 1.0, while the linear loss would result in less variance, as shown in the variance

num-samples	Dataset	perplexity	mean	variance	stddev/mean
	train	50.78	0.2131	0.0196	0.6566
cross entropy	dev	91.77	0.2209	0.0203	0.6451
linear loss	train	49.56	1.0572	0.0326	0.1707
linear loss	dev	90.20	1.0623	0.0331	0.1713
64, NCE	train	119.41	0.7407	0.0631	0.3390
04, INCE	dev	114.55	0.7471	0.0592	0.3256
100 NICE	train	90.87	0.8311	0.0501	0.2693
128, NCE	dev	120.47	0.8383	0.0448	0.2524
DEC NICE	train	73.53	0.8956	0.0560	0.2642
256, NCE	dev	105.99	0.9084	0.0500	0.2461
E12 NICE	train	60.68	0.9243	0.0444	0.2281
512, NCE	dev	96.03	0.9387	0.0394	0.2115
1004 NICE	train	55.72	1.0143	0.0420	0.2020
1024, NCE	dev	92.77	1.0268	0.0383	0.1907
64 Lincor	train	70.80	0.9020	0.0332	0.2021
64, Linear	dev	109.63	0.9064	0.0316	0.1960
100 Lincor	train	65.73	0.9560	0.0313	0.1850
128, Linear	dev	100.55	0.9573	0.0312	0.1844
DEC Lincor	train	58.22	1.0397	0.0318	0.1717
256, Linear	dev	93.79	1.0419	0.0311	0.1694
E10 Linear	train	54.48	1.0778	0.0311	0.1638
512, Linear	dev	91.64	1.0802	0.0296	0.1592
10 <b>0</b> 4 I :	train	51.39	1.1178	0.0312	0.1581
1024, Linear	dev	89.47	1.1225	0.0310	0.1568

**Table 4.3:** Mean and Variance of Unnormalized Outputs in AMI with Sampling-based Training, Linear VS NCE, Sampling with Replacement

and the stddev/mean columns.

sample size	loss type	dev	eval
n/a	no rescoring	35.9	39.8
n/a	cross-entropy	34.8	38.2
64	NCE	36.3	40.1
128	NCE	35.9	39.6
256	NCE	35.2	38.6
512	NCE	35.0	38.5
1024	NCE	35.0	38.4
complete sum	Linear	34.8	38.2
64	Linear	35.3	38.8
128	Linear	35.1	38.6
256	Linear	35.0	38.4
512	Linear	34.9	38.4
1024	Linear	34.8	38.4

Table 4.4: Comparison of WER of RNNLMs Trained with Linear Loss VS NCE

Table 4.4 compares the ASR performance between NCE and linear loss. We see that with the same number of samples, the linear loss system would always outperform the NCE systems. When compared to the cross-entropy systems, although neither linear loss nor NCE could match the cross-entropy baseline, we see that the linear loss system could match the performance on the dev set and slightly worse on eval, while the NCE is inferior to cross-entropy in both datasets.

### 4.3 **Comparison with Sampled Softmax**

Sampled softmax is sometimes referred to as the "importance-sampling" method for RNNLM – it is similar to the linear method in that it uses importancesampling to compute the summation, but sampled softmax does not take out the log operation, resulting in a biased estimator for the loss. As we have proven on page 53, this estimator is always less than the actual value, i.e., it is an *under-estimator* for the true cross-entropy loss.

sample size	loss type	dev ppl	dev	eval
n/a	no LM rescoring	-	35.9	39.8
64	Sampled Softmax	137.81	36.3	39.7
128	Sampled Softmax	116.78	35.9	39.5
256	Sampled Softmax	102.10	35.5	38.8
512	Sampled Softmax	95.82	35.6	38.8
1024	Sampled Softmax	92.87	35.6	38.9
complete sum	Sampled Softmax	91.77	34.8	38.2
64	Linear	109.63	35.3	38.8
128	Linear	99.63	35.1	38.6
256	Linear	93.79	35.0	38.4
512	Linear	91.64	34.9	38.4
1024	Linear	89.47	34.8	38.4
complete sum	Linear	90.20	34.8	38.2

**Table 4.5:** Comparison of WER of RNNLMs Trained with Linear Loss VS Sampled

 Softmax

We compare linear loss with sampled softmax in Table 4.5, where we report both dev perplexities and ASR performance. We see that while the sampled softmax method could potentially speed up training, it does not match performance with models trained with the cross-entropy loss; it also does not train the RNNLMs to self-normalize and therefore does not help in speeding up the inference computation.

	Datacot	nomelovity		varianco	atdday /maan
num-samples	Dataset	perplexity	mean	variance	stddev/mean
anoss options	train	50.78	0.2131	0.0196	0.6566
cross entropy	dev	91.77	0.2209	0.0203	0.6451
linear loss	train	49.56	1.0572	0.0326	0.1707
linear ioss	dev	90.20	1.0623	0.0331	0.1713
64, S-VR	train	75.89	0.9103	0.0260	0.1770
04, <i>5</i> - v K	dev	109.97	0.9163	0.0243	0.1700
128, S-VR	train	60.77	0.9519	0.0315	0.1865
120, <b>5</b> - V K	dev	98.83	0.9559	0.0299	0.1808
DEC S VD	train	58.43	1.0301	0.0245	0.1519
256 <i>,</i> S-VR	dev	93.86	1.0325	0.0234	0.1483
E10 C VD	train	54.71	1.0561	0.0226	0.1424
512 <i>,</i> S-VR	dev	91.53	1.0593	0.0212	0.1373
10 <b>0</b> 4 C VD	train	52.11	1.0935	0.0204	0.1306
1024, S-VR	dev	89.83	1.0978	0.0197	0.1277
64, Linear	train	70.80	0.9020	0.0332	0.2021
04, Linear	dev	109.63	0.9064	0.0316	0.1960
179 Lincor	train	65.73	0.9560	0.0313	0.1850
128, Linear	dev	100.55	0.9573	0.0312	0.1844
DEC Lincor	train	58.22	1.0397	0.0318	0.1717
256, Linear	dev	93.79	1.0419	0.0311	0.1694
512 Lincer	train	54.48	1.0778	0.0311	0.1638
512, Linear	dev	91.64	1.0802	0.0296	0.1592
1004 Linger	train	51.39	1.1178	0.0312	0.1581
1024, Linear	dev	89.47	1.1225	0.0310	0.1568

# 4.4 Comparison with Sampled Variance Regularization

**Table 4.6:** Mean and Variance of Unnormalized Outputs in AMI with Sampling-based Training, Sampled VR VS Linear, Sampling with Replacement

In Section 3.3, we proposed to use importance-sampling on VR loss. We report the experiments in Tables 4.6 and 4.7. We use hyper-parameter  $\lambda = 2.0$  in all the sampled VR experiments.

From Table 4.6, we see that in most cases, the linear loss achieves comparable but slightly worse perplexities than sampled VR, except the case with 1024 samples where the linear system slightly outperforms the corresponding sampled VR system. Sample VR systems also seem to constrain the average normalization terms closer to 1.0 and have smaller variance, but this depends on the  $\lambda$  hyper-parameter we choose.

sample size	loss type	dev	eval
n/a	no LM rescoring	35.9	39.8
n/a	cross-entropy	34.8	38.2
64	S-VR	35.4	38.9
128	S-VR	35.1	38.8
256	S-VR	35.1	38.7
512	S-VR	35.0	38.5
1024	S-VR	34.8	38.4
64	Linear	35.3	38.8
128	Linear	35.1	38.6
256	Linear	35.0	38.4
512	Linear	34.9	38.4
1024	Linear	34.8	38.4
complete sum	Linear	34.8	38.2

Table 4.7: Comparison of WER of RNNLMs Trained with Linear Loss VS Sampled VR

From Table 4.7, we see that again, the linear loss and the sampled VR loss give very similar performances when used in the ASR task on the AMI-SDM datasets.

### 4.5 Chapter Summary

In this chapter, we compared the linear loss proposed in Chapter 2 against several commonly used loss functions in language modeling, both in terms of perplexity on development data and word error rates in speech recognition tasks. We have shown the superiority of the linear loss in both measures. We also show that using sampling during training allows us to not compute the whole summation term in the loss function, which can save much computation.

The sampling used in those experiments uses unigram distribution over the vocabulary, and we allow the same word to be sampled multiple times during each training step. In the next chapter, we perform a more comprehensive study on sampling for language model training. We investigate the impact of sampling specifications and propose more sophisticated sampling schemes that improve the models' performance.

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### Chapter 5

# Impact of Sampling Algorithm on Language Model Training

In Chapter 3, we described a training procedure for RNNLMs that uses importancesampling to speed up its computation. While sampling-based training could save much neural-network computation, the sampling procedure adds computational overheads, especially if it is not implemented efficiently. So far, in all the reported experiments in previous chapters, we use a simple "sampling with replacement" as the implementation. This is in order to get an initial idea of the application of sampling in model training. In this chapter, we conduct a study into the details of sampling algorithms and their impact on language modeling, in performance and training speed.

### 5.1 Sampling with Replacement

In previous chapters, we have so far only considered the case of "sampling with replacement". This is perhaps the easiest sampling procedure to implement, for it can be implemented as sampling one word from a unigram distribution k times, where k is the desired number of samples. Algorithm 1 shows the basic steps.

Algorithm 1: Algorithm for Sampling with Replacement
<b>Input:</b> k, p[1,,n]
<b>Result:</b> A Set S of size k
$1 S = \{\}$
2 while $ S  < k$ do
s = sample a word according to p
$4  S = S \cup \{s\}$
5 end
6 return S

On line 3 of Algorithm 1, instead of naively performing a linear search through the vocabulary, a more efficient binary-search version could be implemented. With binary-search, this step requires a run-time complexity of  $O(\log(V))$  instead of  $O(V)^1$ , with *V* being the size of the vocabulary. Since line 3 will be performed precisely *k* times, this algorithm runs with a  $O(k \log V)$  complexity when line 3 uses a binary search algorithm. Moreover, if we could afford more complex pre-processing of the sampling distribution, then a highly efficient *alias method* [70] could be adopted in this sampling scheme, which

<sup>&</sup>lt;sup>1</sup>The binary-search method also requires a one-time pre-processing of complexity O(V) to compute a vector of accumulative sums of *p*.

runs in O(1) to generate one sample on line 3, and the run-time for generating k samples would be O(k).

However, a problem with sampling with replacement is that it performs duplicated computations if the same word appears more than once in the sample. This almost always happens in real applications of this method. Take English, for example – the most frequent word in English is "the" in most domains; according to Google's N-gram Viewer<sup>2</sup>, the unigram probability for "the" has always been greater than 4.5% in the past 200 years. Hence, if we train an English language model with a sample size greater than 44, we would expect to see the word "the" in the sample at least twice. In our previously reported experiments, we have shown that a sample size around 512 is needed to achieve the best language modeling performance, and this would mean we would expect to see the word "the" appear more than 23 times in the sample! Furthermore, with the number of samples getting large, we would expect to see duplications of many other less frequent words. Even in the case of sampling as many words as the vocabulary-size, if we adopt the sampling with replacement scheme, we will likely not see all words represented in the sample, but instead, we will see frequent words appear multiple times in the sample and very few infrequent words. This would result in wasted computation, not to mention possible unnecessary estimation issues.

<sup>&</sup>lt;sup>2</sup>https://books.google.com/ngrams

### 5.2 Sampling without Replacement

We propose to use sampling without replacement as the sampling method. By definition, this would solve the duplication issue and ensure that a word appears at most once in any sample. However, by making this change, we now have to be careful and redefine terms like the "sampling distribution". Let us take a look at one example to show how this could be an issue.

Again, we use the case of the word "the" mentioned above as an example – let us assume that it has a unigram probability of 0.05, and we specify a sample size of 512 words. We have seen that in the case of sampling with replacement, it is expected that "the" would appear multiple times in the sample, since  $0.05 \times 512 > 1$ . However, in the case of sampling without replacement, we need to define the algorithm's behavior to handles words like "the". For example, should we ensure that "the" will be sampled, i.e., with a sampling probability of 1.0? Or should we assign a very high but still less-than-one probability for it to be sampled? If we need to lower the probability for the, how do we adjust the probabilities of other words? Those are non-trivial questions but are essential in the design of the algorithm.

Sampling without replacement is a commonly used method for sampling in statistics. This problem is formulated in two different ways, with very different mathematical properties of the samples (except when the sampling distribution is uniform, in which case the two formulations are equivalent). While we will describe the details in later sections, we state here, with loosely defined terms, that those two formulations differ because, due to the nature of sampling without replacement, if we "sample from" a distribution, the "distribution of the resulting samples" usually does not match the one from which we sample. Given this distinction, some of the prior work, for example, [71], [72] focus on sampling from a pre-defined distribution, while other works, such as [73], [74] and [75], focuses on making sure the output samples match a pre-defined distribution. Those works give an excellent analysis of the sampling methods' statistical properties, but in general, are not focused on applying the methods in a specific task, for example, in language modeling.

Following our previous results in Equation (3.19), we take the second approach of the interpretation for the problem, since in Equation (3.19),  $p_i$  represents the probability that a word is included in the sample, which we refer to as its *inclusion probability*. Say the vocabulary is *V* and we want to sample *k* words. It is easy to see that, the inclusion probabilities for all the words in *V* should sum up to *k*, i.e.

$$\sum_{w \in V} P(w \in \text{Sample}) = k \tag{5.1}$$

So the sampling procedure runs as follows:

 we work out the "inclusion probabilities" of all words based on their unigram probabilities and the number of samples we want; 2. we sample a set of words based on their inclusion probabilities.

For ease of presentation, in the following sections of this chapter, we first visit the second step and then go back to the first.

### 5.3 Sampling without Replacement: Algorithm

In this section, we study one problem: given a positive integer k, a vocabulary  $V = \{1, ..., n\}$ , and a list of inclusion probabilities for each word  $\{p_1, p_2, ..., p_n\}$ , such that

$$\sum_{i=1}^{n} p_i = k$$

how should one sample a set  $S \subset V$  of distinct words, such that

$$|S| = k$$

and

$$\forall i \in \{1, 2, ..., n\}, P(i \in S) = p_i.$$

### 5.3.1 An Obvious (and Wrong) Approach

Most researchers, including the author of this dissertation, came up with a very straight-forward algorithm for this problem, which unfortunately was not correct. Since it is a common misconception, we demonstrate why it is not correct in this section. The algorithm takes a simple "iteratively sample and re-normalize" method, as shown in Algorithm 2.

Algorithm 2: An Incorrect Sampling-without-replacement Algorithm
<b>Input:</b> k, p[1,,n]
<b>Result:</b> A Set S of size k
$1 S = \{\}$
<sup>2</sup> while $ S  < k$ do
3 $Z = \sum_{i \notin S} p[i]$
4 re-normalize p by Z
5 s = sample a word from p
$6  S = S \cup \{s\}$
<b>7</b>   p[s] = 0
8 end
9 return S

This algorithm runs reasonably efficient and would generate a set of distinct k samples for every run; however, if we carry out the algorithm, then the actual inclusion probabilities of words would not be equal to the specified ones. We demonstrate that with a simple example of selecting two samples from a vocabulary of size 3.

Say we have

$$(p_1, p_2, p_3) = (0.5, 0.5, 1) \tag{5.2}$$

Then the desired probabilities for selected subsets of words should be

$$p(\{1,2\}) = 0$$
  

$$p(\{1,3\}) = 0.5$$
(5.3)  

$$p(\{2,3\}) = 0.5$$

However, we can easily see that if we follow Algorithm 2, then in the first

iteration of the while loop, there is a probability of 0.5 that word 1 is sampled; then in the 2nd iteration, there is a probability of  $\frac{1}{3}$  that word 2 get sampled. This would give

$$p(\{1,2\}) > \frac{1}{6} > 0$$

which does not equal the probabilities we specified. Note that this is a simple example to demonstrate the algorithm is incorrect by devising a case where certain combinations of words are not possible. However, this algorithm is wrong even in the most general cases, even when all combinations allowed in the sample.

### 5.3.2 Reservoir Sampling Algorithm

[75] proposed an algorithm that correctly solves the problem, which is commonly referred to as the *reservoir sampling* algorithm. It is a rather complex algorithm, and we only show its main structure in Algorithm 3 (upon which we make improvements in later sections) and omit some of the details.

Algorithm 3 works by first picking the first k words in the set and then iterates over all the remaining words. When seeing a new word, it first makes a random decision whether to add this new word in the set. If not, it goes on to decide for the next word; otherwise, it randomly picks an existing word to be replaced by the new. We omit the details of computing the vector R, representing the replacement probability for each existing word. The detail of computing the vector R is presented in [75]. The key is setting them so that at

Algorithm 3: Reservoir Sampling Algorithm (incomplete)
<b>Input:</b> k, p[1,,n]
<b>Result:</b> A List S of size k
$S = [1, 2,, k]$ for $i \leftarrow k + 1$ to $n$ do
<sup>2</sup> decide with probability $p[i]$ whether to include i in S
3 if decided to add i then
4 compute $R[1,, k]$ (refer to paper for details)
5 randomly pick $j \in \{1,, k\}$ according to R
$[6 \qquad S[j] = i$
7 end
8 end
9 return S

the end of each iteration of the for loop, the inclusion probability of any word w up to the current index i is proportional to its desired inclusion probability in p[w] so that after the last iteration, the inclusion probability of all words would correspond to the values specified in p.

An analysis of the algorithm reveals that its worst-case time complexity is O(nk), n for iterating over all words, and k for computing vector R of length k. We notice that the computation of R is only needed if we decide to add the new word to the set, and the probability of that depends on the new word's inclusion probability. We can take advantage of that and make the algorithm faster (in terms of average time complexity) by arranging the word orders such that the p[i] is descending for i. Of course, sorting would add complexity of  $O(n \log n)$  to the algorithm, but in the case where we pre-process once and then sample multiple times, this one-time cost will be amortized, leading to significant speed-ups.

### 5.3.3 2-stage Reservoir Sampling Algorithm

One benefit of the reservoir sampling algorithm is that it introduces minimum dependencies among different words. Note that, since we are sampling a fixsized set, certain dependencies between words are inevitable, and one example would be the previous example shown in Equation (5.2), wherein the solution Equation (5.3), we see that 1 and 2 can never be sampled simultaneously. This type of dependency is inevitable because of the nature of the "*n* choose k" problems. However, it can be shown that this algorithm adds no other dependencies in the output than the problem imposes; again, interested readers should refer to [75] for details.

A significant problem of the algorithm is the *n* term in its complexity. This is because, in practice, *n* could be in the hundreds of thousands or even millions. We now propose a modification that does not need to iterate over all words. We achieve this by adding more dependencies in the output, which we call a *2-stage sampling algorithm*.

Here is how the 2-stage sampling algorithm works: we first divide the vocabulary  $\{1, 2, ..., n\}$  into *m* disjoint subsets  $s_1, s_2, ..., s_m$ , where

$$p(s_i) \stackrel{\text{\tiny def}}{=} \sum_{j \in s_i} p(j) \leq 1.0, \forall i \in \{1, 2, ..., m\}$$

The grouping operation could be simply implemented with a greedy algorithm, as shown in Algorithm 4. To simplify the procedure and improve efficiency, we only group words with consecutive indices, which is usually

arbitrarily chosen, and does not affect the correctness of the algorithm.

Algorithm 4: Placing Words into Groups for 2-stage Sampling
<b>Input:</b> p[1,,n]
<b>Result:</b> g[1,2,,m], each element is a group represented as a set
1 idx = 0
$2 g_i dx = 0$
$3 \text{ g_probs} = [0.0]$
4 g = []
5 while $idx < n$ do
6 <b>if</b> $g_probs[g_idx] + p[idx] \le 1.0$ <b>then</b>
7 $g[g_idx].insert(idx)$
$s \qquad g_probs[g_idx] = g_probs[g_idx] + p[idx]$
9 else
10 $g_idx++;$
11 $g[g_idx].insert(idx) g_probs[g_idx] = p[idx]$
12 end
13 end
14 return g

After the grouping, we do the sampling. In the first stage, we sample k groups according to group inclusion probabilities p(s); in the second stage, we pick precisely one element from each group, with probabilities proportional to word inclusion probabilities in this group.

It is not hard to see that this algorithm is "correct" in that, the probability for any word w being sampled is p(w): first we need to sample the group swhere  $w \in s$ , with probability  $p(s) = \sum_{i \in s} p(i)$ ; then we need to pick w from the group, with probability  $\frac{p(w)}{p(s)}$ , and the algorithm is correct because,

$$p(s) \cdot \frac{p(w)}{p(s)} = p(w).$$

However, we also see that this modification adds unnecessary dependencies between words – for example, two words in the same group would never be simultaneously sampled.

Now we analyze the complexity when the grouping  $\{s_1, s_2, ..., s_m\}$  is already given. In the first stage, we sample *k* groups from *m* groups, and so it has a complexity of O(km). It is hard to perform a rigorous analysis in the second stage because we do not know the grouping information. Nevertheless, sampling one from multiple words could easily be implemented with a binary search algorithm<sup>3</sup>, which is logarithmic w.r.t. group sizes. Suppose the grouping is close to evenly, then each group would have  $\frac{n}{m}$  elements, and then the complexity for the second stage is

$$O(k\log\frac{n}{m})$$

This will give a total complexity of

$$O(km) + O(k\log\frac{n}{m}) = O(k(m + \frac{n}{m})) = O(k\max(m, \frac{n}{m}))$$

This analysis under simple assumptions also shows that a good choice of the number of groups to use would be around  $\sqrt{n}$ . We can generalize this method to 3-stage (or even 4 or above), and the optimum number of groups

<sup>&</sup>lt;sup>3</sup>Although the alias method gives better performance than binary search, as in later chapters, we will not always sample from the same distribution; therefore the alias method is not suited for our experiments.

would then be around  $n^{\frac{1}{3}}$ , but the efficiency improvement would be marginal compared to the computation overhead.

### 5.3.4 Systematic Sampling Algorithm

Another sampling method with added dependencies is the *systematic sampling algorithm* method proposed in [76]. This algorithm is best explained with a visual aid shown in Figure 5.1. Figure 5.1 shows a 2-D space with X and Y axes. For each word *i* with inclusion probability  $p_i$ , there is a rectangle of size  $[1 \times p_i]$ , where 1 is the width on the x-axis, and  $p_i$  is the height on the y-axis. In the figure, we have assigned different words with different colors. Because we have

$$\sum_{i} p_i = k$$

We can now put the rectangles for all words at the corner of the 1st quadrant, and by possibly cutting rectangles for certain words horizontally, arrange all of them fit in the shape of a larger rectangle  $[k \times 1]$ . In the example in Figure 5.1, we cut the rectangles with the colors orange, dark blue, and purple, and all of those rectangles fit in the space of  $[6 \times 1]$ .

Now we uniformly choose a random number *r* between 0 and 1 and draw a line y = r on the rearranged rectangles, as shown with the dotted line in Figure 5.1. This line would touch exactly *k* rectangles <sup>4</sup>. The words corresponding to

<sup>&</sup>lt;sup>4</sup>In order for this to be well-defined, we shall also define what "touch" means. It might vary with implementation, but an easy way would be to consider each rectangle as taking a left-close and right-open space, except the top-rightest one, which is close for both left and

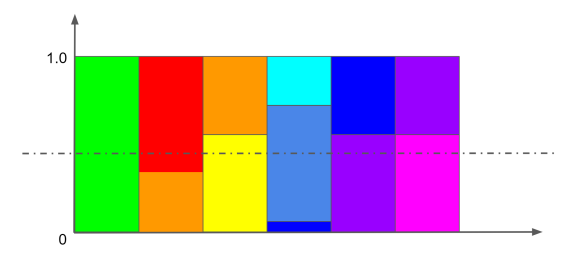


Figure 5.1: Visual Aid to Help Understand the Systematic Sampling Algorithm.

those rectangles are then the sampled words. Since we draw a random number from a uniform distribution between 0 and 1, the probability of sampling a word is proportional to its "height" in the diagram, and thus we guarantee that this method also satisfies the inclusion probability conditions. Also, it introduces strong dependencies in the distribution of the sampled words, and the relative position of rectangles determines this dependency after the rearrangement. For example, if the arrangement is like that in Figure 5.1, then we know that if the word represented by the light-blue color is sampled, then we know for sure that the words represented by red will be sampled, and the word represented by yellow will not be sampled.

Now the complexity: This algorithm requires us to search for precisely one rectangle at *k* different height levels, and each of those searches could be

right. The exact choice does not matter theoretically since the probability of cutting "exactly on the boundary" is always zero; however, in practice where numbers are represented in approximations, e.g., the floating-point number representation, this might make some edge cases more complicated than in theory.

implemented as a binary search. On average, each height level would have  $\frac{n}{k} + 1$  rectangles<sup>5</sup>. So the complexity would be

$$O(k\log(\frac{n}{k}+1))$$

Note this method is a continuous (as opposed to discrete) version of the 2stage method – because we can divide the probability mass w.r.t. one word into different parts, the algorithm can start with exactly *k* "groups". The difference is, in order to avoid picking the same word in 2 different groups (since a word can now be in two groups), the algorithm does not independently sample from each group but jointly does so, avoiding the possibility of sampling one word twice.

### 5.3.5 2-stage systematic sampling

We can also apply the 2-stage idea for the systematic sampling algorithm, where we use systematic sampling in the first stage to sample groups. However, a simple analysis of this method would reveal that this does not change the Big-O complexity of the algorithm – since in both stages, the search could be efficiently implemented as a binary search, if we adopt a 2-stage sampling method.

However, there is still merit in performing 2-stage sampling with systematic

 $<sup>^{5}</sup>$ we have +1 because, with high probability, each original rectangle would be cut in 2, though each level would on average have one more rectangle

sampling. The reason is, so far, we only sample from a unigram distribution, which allows the opportunity to pre-process the distribution – computing a vector storing the cumulative sums of the unigram probabilities to perform an efficient binary search. However, if we want to repeatedly sample from a (slightly) different distribution every time, then we could not afford to perform the pre-processing every time before sampling. However, using the 2-stage sampling method could alleviate this issue. More details are shown in Chapter 6.

### 5.4 Computing Inclusion Probabilities

In the previous sections, we assumed that the inclusion probabilities are already given and studied algorithms to sample from such distributions. In this section, we study how to generate inclusion probabilities for a vocabulary, given their unigram probabilities and the sample-size we want. Note that the algorithm we propose could be used to compute inclusion probabilities for not just unigrams but higher-order models as well, and we pick unigrams to make it easy to describe the method.

Say we have a vocabulary  $V = \{1, 2, ..., n\}$ , and a unigram distribution u(.) on V. Our task is to compute a vector of inclusion probabilities  $p_i$ , where  $p_i$  represents the probability of selecting word i in the sample. In the case of sampling k words from a vocabulary of size n, the  $p_i$ 's need to somehow

"reflect" the unigram probabilities, while satisfying the following constraints,

$$\sum_{i=1}^{n} p_i = k \tag{5.4}$$

and

$$0 \le p_i \le 1, \forall i = 1, ..., n.$$
 (5.5)

This problem is an open-ended one, to which there is not one single "correct" solution because the inclusion probabilities could "reflect" the unigram distribution in different ways. One primary reason that we want the sample to reflect the real distribution is in Equation 3.19, where we show that we could minimize the variance in the sampled estimator if the sampling distribution is proportional to the actual distribution. In our case, we use the unigram distribution as a proxy for the actual distribution, and thus an ideal setup for the inclusion probabilities is

$$p_i = k u_i, \forall i.$$

This guarantees that for every word in the vocabulary, the probability of it getting sampled is proportional to its unigram probability; however, in real languages, the unigram probabilities for different words vary by a lot, and setting the inclusion probability to be proportional to unigrams, the most frequent words in the language would usually have an inclusion probability of greater than one, which violates Equation 5.5, and the  $p_i$ 's do not represent valid probabilities any more. The statistics on the word "the" in English described in

Section 5.2 is an excellent example of this issue.

The solution we propose is to distribute the excess probability mass of frequent words to less frequent words proportionally to their unigram probabilities. In this procedure, we first compute a  $P = [p_1, p_2, ..., p_n]$ , where  $\forall i \in V, p_i = u_i \cdot k$ , so some of the numbers might be larger than one. We perform the normalization in Algorithm 5.

Algorithm 5: Inclusion Probability Normalization Algorithm
<b>Input:</b> k, p[1,,n]
<b>Result:</b> normalized p[1,,n]
1 $m = \operatorname{argmax}_i p[i]$
2 while $p[m] > 1.0$ do
d = p[m] - 1.0
4 $p[m] = 1.0$
5 $s = k - p[m]$
6 for $i \leftarrow 1$ to $n$ do
7   if $i \neq m$ then
$ \mathbf{s}    p[i] = p[i] + d \times p[i]/s$
9 end
10 end
11 $m = \operatorname{argmax}_i p[i]$
12 end
13 return p

Algorithm 5 works by picking words whose inclusion probabilities are greater than one and proportionally distribute the extra probability mass to all other words, proportional to their unigram probabilities. This procedure needs to be carried out iteratively until there are no words with larger-than-one inclusion probabilities. We need such an iterative algorithm because even if a word initially has an inclusion probability smaller than 1.0, it might end up larger than one after acquiring some of the probability mass from other words. Fortunately, this algorithm is guaranteed to terminate – a simple way to see that is, this algorithm guarantees that the difference between the largest and the smallest p[i]'s is decreasing after each iteration, and the worst-case scenario would be the "*n* choose *n*" case, where the final inclusion probability vector contains all 1's no matter what the initial values are<sup>6</sup>.

### 5.5 Evaluation of Different Sampling Methods

In this section, we compare the different approached mentioned in this chapter in terms of efficiency.

We compare the speed of running different sampling algorithms by sampling from a unigram distribution in Table 5.1. In those experiments, we randomly generate unigram probability distributions for the specified vocabulary and then normalize the generated distribution according to Algorithm 5. For 2-stage sampling algorithms, additional processing is performed to break the vocabulary into groups. This is done by following the greedy algorithm represented in Algorithm 4. For the "sorted" experiments, we sort the vocabulary once according to their inclusion probabilities in a descending way. Since we only sample from the same unigram distribution, the pre-processing steps only need to be done once in the beginning. We report the run-time, in seconds, of running the same sampling procedure of "sampling *k* words from a vocabulary

<sup>&</sup>lt;sup>6</sup>We point out here that, again, this is the result of purely theoretical analysis. In the case of floating-point numbers, depending on the details of implementation, it is possible that this "n choose n" would never terminate, and special attention is needed in the implementation of the algorithm to avoid such issues.

of size *n*" exactly *t* times.

n, k, t		reservoir	systematic		
	std	sorted	2-stage	std	2-stage
100000, 50, 100	0.256	0.288	0.032	0.016	0.008
100000, 100, 100	0.376	0.388	0.04	0.016	0.008
100000, 250, 100	1.032	0.968	0.044	0.012	0.016
100000, 500, 100	3.112	2.952	0.056	0.02	0.012
100000, 1000, 100	9.94	9.228	0.088	0.028	0.024
100000, 2000, 100	33.592	28.364	0.208	0.036	0.036
100000, 4000, 100	113.864	87.208	0.704	0.044	0.06
100000, 10000, 100	514.148	354.064	7.648	0.084	0.116
100000, 20000, 100	1473.72	867.288	86.472	0.128	0.176
100000, 40000, 100	3536.45	1488.59	761.188	0.196	0.276
10000, 50, 1000	0.488	0.468	0.056	0.008	0.012
10000, 100, 1000	1.132	1.056	0.08	0.012	0.016
10000, 250, 1000	5.076	4.316	0.128	0.02	0.028
10000, 500, 1000	16.08	12.932	0.2	0.036	0.06
10000, 1000, 1000	48.012	35.58	0.36	0.052	0.096
10000, 2000, 1000	138.936	87.74	1.132	0.104	0.14
10000, 4000, 1000	323.832	146.00	6.276	0.172	0.26

**Table 5.1:** Time (of *t* runs of *n* choose *k* in seconds) of Sampling from Unigrams

For reservoir sampling, as shown in columns 2 to 4 in Table 5.1, we have the following observations.

 sorting the vocabulary helps reduce the run-time of the algorithm for most of the experiments. However, in a few cases where the number of samples is small, it takes more time than the unsorted version, and this is due to the added cost of sorting the vocabulary – when we only want a small number of samples, this added cost is too high for the efficiency improvement to compensate. 2. while sorting brings a small improvement to the speed of the algorithm, the 2-stage method brings significant speed-up to the original algorithm; depending on the actual configuration, this speed-up ranges from one order of magnitude (e.g., the first row) to two orders of magnitudes (e.g., the second-to-last row).

Now we analyze columns 5 and 6 for the systematic sampling algorithm. We see the following,

- 1. Compared to the reservoir sampling method, the systematic algorithm is much faster. In the extreme case of a large number of samples, it is 3536.45/0.276 = 12813 times faster than the reservoir algorithm.
- 2. the 2-stage version of the algorithm slows down the computation this is no surprise, as our analysis showed that the 2-stage algorithm has the same complexity as the original 1-stage algorithm in terms of Big-O order, however, because the added overhead of the 2-stage algorithm, it would result in a larger constant in the complexity that is hidden in the Big-O notations.

### 5.6 Language Modeling Experiments

In this section, we perform a simple set of experiments investigating the difference of different sampling methods in the task of language model training. We use the aforementioned PyTorch RNNLM implementation as a starting point,

num-samples	Dataset	perplexity	mean	variance	stddev/mean
cross entropy	train	50.78	0.2131	0.0196	0.6566
	dev	91.77	0.2209	0.0203	0.6451
1:0000 1000	train	49.56	1.0572	0.0326	0.1707
linear loss	dev	90.20	1.0623	0.0331	0.1713
	train	65.03	0.9533	0.0326	0.1893
64, NR	dev	100.69	0.9525	0.0331	0.1911
100 NID	train	59.25	1.0360	0.0256	0.1544
128, NR	dev	95.32	1.0424	0.0250	0.1518
	train	54.52	1.0997	0.0346	0.1692
256, NR	dev	92.01	1.1040	0.0336	0.1659
512, NR	train	49.10	1.1361	0.0436	0.1838
	dev	89.12	1.1428	0.0429	0.1812
1024, NR	train	46.25	1.1823	0.0504	0.1899
	dev	89.26	1.1892	0.0518	0.1914
64, R	train	70.80	0.9020	0.0332	0.2021
	dev	109.63	0.9064	0.0316	0.1960
128, R	train	65.73	0.9560	0.0313	0.1850
	dev	100.55	0.9573	0.0312	0.1844
256, R	train	58.22	1.0397	0.0318	0.1717
	dev	93.79	1.0419	0.0311	0.1694
512, R	train	54.48	1.0778	0.0311	0.1638
	dev	91.64	1.0802	0.0296	0.1592
1024, R	train	51.39	1.1178	0.0312	0.1581
	dev	89.47	1.1225	0.0310	0.1568

Table 5.2: Comparison between Different Sampling Methods

and implement the different schemes described in previous sections that uses a pre-computed unigram distribution as the sampling distribution.

### 5.6.1 The Impact of Replacement

We report the training stats on the AMI corpus in Table 5.2 with two types of sampling procedures, namely sampling with replacement (R) and sampling

without replacement (NR). As described before, in the R scheme, we could sample the same word multiple times for the same batch, which makes it easier to implement the sampling algorithm, but there could be a duplicate of computation; when we sample without replacement (NR), there is no duplicate of the same word in the sampled set.

From Table 5.2, we see that, in general, the sampling-without-replacement methods yield better perplexities than sampling-with-replacement for all samplesizes, and with more significant differences for smaller sample-sizes. This is expected since, in the sampling-without-replacement approach, the generated samples have no duplicates and cover more vocabulary words. We also see that sampling-without-replacement methods result in slightly higher variances for the normalization term. This is explained by the fact that, for the samplingwithout-replacement approach, the inclusion probabilities of words are not always proportional to their unigram probabilities, thus resulting in higher variances, following our analysis done in Section 3.2.3.

### 5.7 Chapter Summary

In this chapter, we focus on the sampling algorithm used in importancesampling based language model training. We present algorithms for sampling with and without replacement and propose improvements on the samplingwithout-replacement algorithm, making it more efficient. We also present an algorithm for normalizing the inclusion probability vector used in the samplingwithout-replacement algorithm.

We present our experimental results comparing the efficiency of different sampling algorithms and comparing their performance in language modeling tasks. We show that our proposed 2-stage sampling method gives significant speed-up over the unmodified baseline algorithm and that using a sampling-without-replacement scheme outperforms the sampling-withreplacement scheme in language modeling tasks.

### Chapter 6

## Batch Training and Sampling from Longer Histories

In previous chapters, we have proposed using importance-sampling to help speed up RNNLM training with the linear loss and reported experimental results, where we compared the linear loss with some other loss functions and investigated the impact of different sampling algorithms.

Recall in Equation (3.19) on page 55, where we have shown that to minimize the variance of the loss estimator computed from sampling, the sampling distribution need be proportional to the "true" distribution of data. In all the experiments we have reported, we use simple unigram distribution as a proxy for the true distribution. While not necessarily the optimal distribution, this allows smooth interaction between the sampling-based training and batchbased training – by sampling from a unigram distribution and not considering any history information, different examples in a batch may share the same samples, which helps improve the computational efficiency of model training.

At this point, readers might not be convinced of the importance of including longer histories, especially since we have demonstrated with our previously reported experiments that reasonably good performance could be achieved with sampling from unigrams. While we will conduct detailed experiments, comparing results with different sampling distributions, here is an intuitive explanation for why the history might matter.

Let us take English as an example and consider the word "San". Say a sentence contains the phrase "San Francisco", and the model now is computing the loss for the word "Francisco". We know that words like "Jose", "Bernardino" and "Diego" all have relatively low unigram probabilities but frequently occur after "San". To accurately estimate the normalization term for history "San", it would be essential to include those words in the sample. However, sampling from a unigram distribution would make this very unlikely, making the estimated normalization term deviate from the actual value; On the other hand, if we sample from a bigram in this case, then it is much more likely that all those words get sampled, leading to a more accurate estimator for the loss function.

This chapter discusses how to incorporate longer history information in the sampling-based training scheme, for example, sampling from a bigram distribution. In the case of training on one example at a time, it is trivial to change the training scheme to support sampling from bigrams; however, it is not the case for batch-based training. This is because different examples in the same batch usually have different histories, so we can not make all examples in the batch share the same samples; it is possible to generate different samples for different examples in the same batch, adding unwanted computation overheads and is therefore not ideal.

To balance the improved computational efficiency of batch training and the more accurate estimation with history information, we propose generating the sampling distribution from the average of all the *n*-gram distributions in the batch.

### 6.1 Average *n*-gram Distribution in a Batch

Let us view all examples in a batch *B* as a set of (history, word) pairs, denoted as (h, w).

$$B = \{(h, w)\}$$

By pre-training an *n*-gram model on the training data, each *h* would correspond to a probability distribution over the vocabulary by truncating history and retaining only the last n - 1 words. We denote the distribution for history *h* as  $P_n(.|h)$ . Then, we set the sampling distribution  ${}^1 s(.)$  for this batch as,

$$s_B(w) = \frac{1}{|B|} \sum_{(h,w)\in B} P_n(w|h)$$
(6.1)

### 6.2 Sampling with Longer Histories

Previously in Chapter 5, we introduced efficient algorithms to sample from a unigram distribution, which involves a one-time pre-processing of the distribution, which the efficient sampling algorithm needs. A side effect of using longer histories as the sampling distribution is that now we sample from different distributions for different batches, so it is no longer possible only to pre-process once and reuse the output of pre-processing for all later computations. Here we propose a method that achieves the middle ground – a sampling algorithm that involves a one-time pre-processing step and supports sampling from different run-time distributions while still being efficient.

Note that the method we propose is not a general-purpose sampling scheme that supports any run-time distribution but operates under the assumption that while the run-time sampling distributions might differ, their differences are relatively small. To achieves this, we first perform a highly selective pruning procedure over the higher-order *n*-grams. The pruning works by removing certain higher-order *n*-grams if they do not meet the specified criteria, and

<sup>&</sup>lt;sup>1</sup>Readers are reminded here that *sampling distribution* is not the distribution we sample words from, but rather a starting point to generate the *inclusion probability* distribution, which the sampling algorithms use. More details are in Section 5.4.

we use their back-off lower-order *n*-gram probabilities instead. For example, if bigram p(A|B) does not meet our criteria and is pruned away, then the sampling probability of *A* with history *B* will be its unigram probability P(A); if trigram P(A|BC) is pruned away but P(A|C) is not pruned, then the sampling probability of *A* with history *BC* will be its bigram probability P(A|C).

The pruning criteria is the following,

1. we prune away  $p(w|w_1, ..., w_{n-1})$  if the ratio

$$\frac{p(w|w_1,...,w_{n-1})}{p(w)}$$

is smaller than a threshold, and we set this threshold to be 100;

for n > 2, we prune away p(w|w<sub>1</sub>,..., w<sub>n-1</sub>) if its ratio to its immediate backoff *n*-gram, i.e.

$$\frac{p(w|w_1,...,w_{n-1})}{p(w|w_2,...,w_{n-1})}$$

is smaller than a threshold, which we set as 2.

The intuition behind the pruning idea is, given a history, if the higher-order *n*-gram probability of a word is significantly larger than its back-off *n*-gram or unigram, then we need to raise the sampling probability of this word by keeping its higher-order *n*-gram. Otherwise, we would not likely sample this word and significantly underestimate the normalization term in our estimation.

After the pruning is performed, for any history h, most of the words w

in the vocabulary would rely on backing off to their unigram probabilities in generating their sampling probabilities. By backing off, we do not mean directly using their unigram probabilities, but instead, take the unigram probabilities and multiply by a factor to make sure the sum of probabilities for all words add up to one. This is part of the standard methods with a back-off *n*-gram model, and readers could refer to [77] for more details.

Note that in previous chapters, we pre-process the unigram probability vector to compute a vector of accumulative sums, on which we perform a binary search in run-time to save computation. Now, because we might have a different distribution at run-time, such an accumulative sum vector is different each time. We use the 2-*stage sampling* method described in Chapter 5 for sampling and propose the following method to avoid adding substantial computational overheads for higher-order *n*-grams. Here is an outline of the 2-stage sampling method,

- Pre-processing
  - 1. Generating grouping structures
  - 2. Compute accumulative sum vector for groups
  - 3. Compute accumulative sum vector for within each group
- Run-time
  - 1. pick groups according to group accumulative sum vector

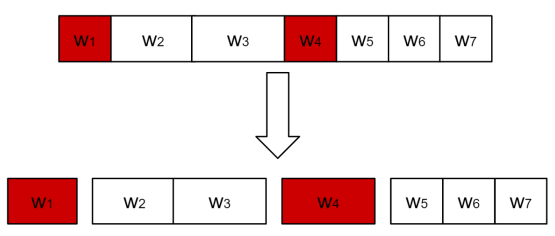
 pick a word from each selected group using its accumulative sum vector

For sampling with *n*-grams, its outline is slightly different,

- Pre-processing
  - 1. Generating grouping structures
  - 2. Compute accumulative sum vector for within each group
- Run-time
  - 1. Re-grouping
  - 2. pick groups according to the output from step 1
  - pick a word from each selected group using its accumulative sum vector

The pre-processing step is almost identical to the unigram case, and the only difference is that we do not compute the accumulative sum vector for groups since it is not needed.

At run time, the first step is to perform a procedure that we call re-grouping. Say the current history is h, and we first identifies all words w following this history whose probabilities do not back off to unigrams. We find the groups they belong to for all those words and change the structures for those groups by further dividing them into subgroups. The result of the re-grouping is that each resulted group would contain only a single word that does not back off



**Figure 6.1:** Changing group assignments for words when changing the sampling distribution from unigrams to higher-order *n*-grams. The group containing words  $w_1, ..., w_7$  is divided into 4 groups since  $w_1$  and  $w_4$  have higher-order *n*-gram probabilities.

to unigram, or a consecutive block of words, all of which back off to unigram. Note we do not change the relative ordering of the words in this group.

An example of this is shown in Figure 6.1. In this example, we have a group with words  $w_1, ..., w_7$ , and the words marked in red, namely  $w_1$  and  $w_4$ , do not back off to unigram. We then further divide the group into four subgroups:  $w_1$  and  $w_4$  both become single-word groups. Two more groups are formed that consist of back-off words, one for  $\{w_2, w_3\}$ , and the other for  $\{w_5, w_6, w_7\}$ .

The second step is to pick *k* groups from all groups. For this step, we need to generate the accumulative sum vector for the groups from scratch each time. This was carried out in pre-processing steps for unigram 2-stage sampling since the group probabilities do not change. This step is the primary overhead that this algorithm introduces. Also, we note that the number of groups now is larger than the unigram sampling case because step 1 of the algorithm would increase the number of groups. A careful analysis shows that the increase for

the number of groups is bounded by 2d, where d is the number of words that do not back off to unigram. This is because, in the re-grouping step, a word that does not back off to unigram adds at most two additional groups in the algorithm. Therefore, as long as we prune the *n*-gram language well, this step does not add substantial overhead to our computation.

The last step is to pick precisely one word from each sampled group. Because in the grouping restructure step, we do not change the relative ordering of the words, we could reuse the pre-computed accumulative sum vector for computation.

### 6.3 Experiments

In this section, we report experiments that compare sampling from a unigram distribution versus from distribution with histories, including bigram and trigram histories. The experiments reported in this section are conducted with Kaldi, and the RNNLM implementation uses Kaldi-RNNLM [78].

We first report experiments on the AMI dataset. The data preparation of this dataset follows the provided script [79] in Kaldi. Note that this script has its way of dividing the training and development data, and thus the reported perplexities on the development set are not comparable with the previously reported AMI numbers. The results are in Table 6.1.

We also report the rate of convergence comparing unigram, bigram, and

num-samples	<i>n</i> -gram order	perplexity	mean
128	1	58.3	1.13
128	2	53.6	1.09
128	3	51.9	1.06
256	1	51.4	1.09
256	2	49.7	1.08
256	3	49.1	1.07
512	1	49.4	1.08
512	2	48.4	1.08
512	3	48.0	1.07

Table 6.1: Effect of Sampling From Longer History in Switchboard

trigram-based sampling, for the setup using 256 samples in Figure 6.2, where we plot the cross-entropy loss on the development set.

Combining the information from Table 6.1 and Figure 6.2, we have the following observation,

- 1. Including history information for sampling helps improve language modeling performance as measured by dev perplexity.
- 2. Including history information also helps bring the mean of the normalization term closer to one; this effect is more pronounced for models with fewer samples.
- 3. History information also helps improve the convergence rate during training.
- 4. For all the observation above, we have seen that using a bigram language model is better than unigram, and trigram would be better than bigram.

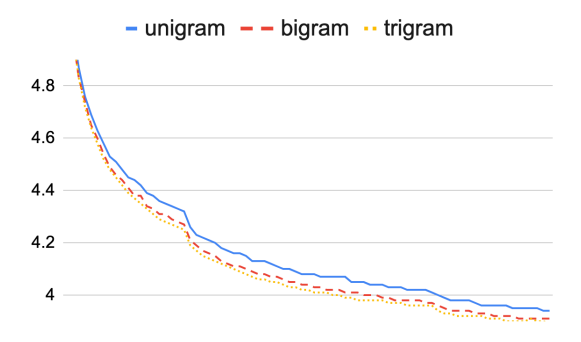


Figure 6.2: Convergence Rate VS Sampling Distribution

### 6.4 Chapter Summary

This chapter identifies the potential problem of the sampling-based method introduced in previous chapters, which uses a unigram probability distribution for sampling. We propose to include history to improve the sampling-based RNNLM training and sample from an *n*-gram distribution based on the context. We propose an efficient method that computes the average *n*-gram distribution based on all histories in the batch and makes the examples in the same batch share the same samples. This makes the computation tractable, and experiments show that including history information in the sampling step leads to better language models' performance measured by perplexity on the development set. It also pushes the average normalization terms closer to one.

This concludes the first part of this thesis, which is primarily on improving the computational efficiency of RNNLMs, both for training and inference computation. In part II, we introduce methods to improve the computational efficiency of applying RNNLMs in lattice rescoring for speech recognition.

## PART II:

# IMPROVING THE COMPUTATIONAL EFFICIENCY OF RNNLM LATTICE RESCORING

### Part II Outline

Various techniques to make the RNNLM computation more efficient both in training and inference were studied in Part I. We focus in Part II on applying RNNLMs in speech recognition tasks. We study the commonly used lattice-rescoring method for applying an RNNLM and propose improvements to the algorithm, making it more computationally efficient. This part includes Chapters 7 and 8. We first give a gentle introduction to finite-state transducers in Chapter 7, which lays the foundation of the lattice-rescoring algorithm on which we focus. We then introduce the standard lattice-rescoring algorithm using the FST framework and the commonly used *n*-gram approximation method for applying an RNNLM in lattice rescoring, which serves as the baseline method upon which we try to improve. We then introduce the pruning algorithm for lattice-rescoring and report our experimental results.

### Chapter 7

## Lattice Rescoring in the FST Framework

We begin with a brief introduction to *Finite-state transducers* (FST)<sup>1</sup>, with which lattice rescoring methods are usually implemented. FSTs are commonly used in most major hybrid speech recognition systems. They elegantly unify Hidden-Markov Models, the use of context-dependent phones, pronunciation lexicon, and an *n*-gram language model into one decoding graph, each of which may be represented using an FST, and may be amalgamated into a single FST, using the *FST-composition* operation, while preserving all the information of the individual FSTs. Although the FST framework is not necessary for building an ASR system, and it does not add any additional modeling power, it dramatically

<sup>&</sup>lt;sup>1</sup>We follow the convention in the speech community that when we say FST, it usually means *weighted finite-state transducer* or WFST, except when we make clear we are talking about unweighted FSTs.

reduces the amount of work by researchers by providing a level of abstraction and makes the system easier to understand and implement.

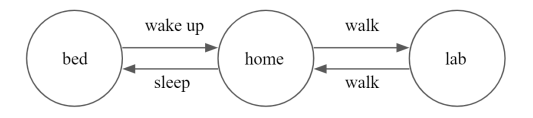
This chapter is not a comprehensive description of FSTs and FST operations but a gentle introduction to the necessary concepts to understand our proposed algorithms in the next chapter. Interested readers are referred to [23, 80] for details on FSTs. In this chapter, we first introduce a *finite-state automaton*, which may be viewed as a special (and simple) version of an FST; then, we talk about FSTs and the *composition* operation, which serves as the foundation of the standard lattice-rescoring algorithm and the pruned improvement that we propose in Chapter 8.

### 7.1 Finite-state Automaton

In plain natural language, a *finite-state automaton* represents a graph with a finite number of nodes connected with directed arcs. Let us take a look at a real-life example in Figure 7.1. Although simple, this diagram is a surprisingly realistic depiction of all those years of working on a Ph.D. experienced by the author.

We can see the following from the diagram.

- It has three states;
- It has four arcs connecting from a state to another;
- On each arc, there is an associated label.



**Figure 7.1:** Diagram depicting the author's experience of working on a Ph.D. It consists of three states and four directed arcs connecting one state to another. This diagram does not satisfy the condition for finite-state automata because it does not have a start state and a final state.

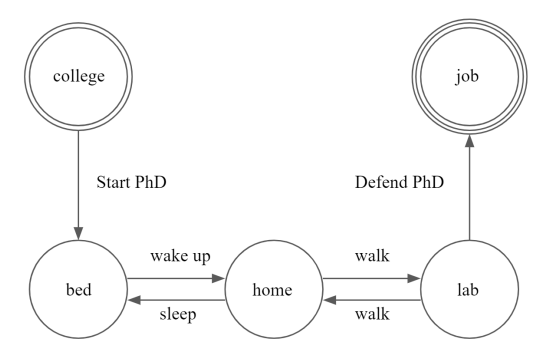
However, Figure 7.1 does not yet satisfy the condition for an FSA. Two additional requirements are needed in order to make a mathematically well-defined FSA.

- It needs to have a start state;
- It needs to have one or more final state[s];

If we take Figure 7.1 and change it slightly to Figure 7.2 by adding a start state and final state, along with necessary arcs that connect them to the original diagram, then it is a valid FSA.

Mathematically, an FSA is a 5-tuple,  $(Q, \Sigma, I, F, \delta)$ , where

- *Q* is a finite set of states;
- Σ is a finite alphabet set;
- $I \subseteq Q$  is the set of initial states;
- $F \subseteq Q$  is the set of final states;



**Figure 7.2:** A finite-state automaton depicting the author's experience of working on PhD. It consists of five states, and 6 directed arcs connecting one state to another. The start state is represented by the double circle and the final state is represented by the triple circle.

δ ⊆ Q × (Σ ∪ {ε}) × Q is the set of arcs. (ε represents the "empty" symbol, or lack of a symbol)

The details of the definition might vary. For example, some definitions require that the cardinality of *I* or *F* must be exactly 1. Nevertheless, those constraints do not hamper the modeling capacity of an FSA. <sup>2</sup>

We could extend the definition of an FSA to *weighted finite-state automaton* (WFSA), where associated with each arc in  $\delta$ , there is a weight. The domain of the weight is any *semi-ring*, and readers may refer to [81] for more details.

<sup>&</sup>lt;sup>2</sup>By this, we mean that if an FSA A under one definition breaks some constraint of a different definition of FSA, then we could easily reconstruct another FSA A' that is mathematically equivalent.

### 7.2 Finite-state Transducer

*Finite-state transducers* (FST) are a generalization of FSAs, where instead of just one symbol associated with an arc, now there is a separate "input symbol" and "output symbol". Note that an FSA could be seen as a special case of FST where the input and output symbols are always identical. Similarly, we expand the definition of (unweighted) FSTs to weighted finite-state transducers (WFST) by making each arc associate with a weight.

There are two common ways to view a WFST. Firstly, a WFST represents a function<sup>3</sup>

$$s_i \to (s_o, w) \tag{7.1}$$

In the second way, a WFST represents a function,

$$(s_i, s_o) \to w$$
 (7.2)

In both interpretations,  $s_i$  represents an input sequence,  $s_o$  an output sequence; and w a weight. In the first interpretation, it means that the FST takes input  $s_i$  and transduces it to  $s_o$ , with weight w; in the second interpretation, it means that the FST "accepts" the sequence pair ( $s_i$ ,  $s_o$ ) with weight w.

<sup>&</sup>lt;sup>3</sup>Strictly speaking, a function needs to be *right-unique* [82], meaning there should be a unique output for a given input; this is not always the case for general FSTs since FSTs also support one-to-many mapping; but within the scope of this thesis, the WFSTs we use are always deterministic and thus always represent a function.

### 7.3 FST Composition

*Composition* is an operation on FSTs that combines two transductions. If FST *A* transduces sequence *x* to sequence *y*, and FST *B* transduces *y* to *z*, then the Composition of *B* and *A*, represented by  $B \circ A$ , transduces *x* directly to *z*. FST composition is an important operation that combines FSTs that operate on different layers of representations. For example, in a typical speech recognition system, there are four different FSTs, namely *H*, *C*, *L*, and *G*. The four FSTs represent the HMM topology that transduces from phone-states to context-dependent phones, context-dependency that transduces context-dependent phones to words, and finally grammar, which is usually an FSA that we use to compute the weight of word equences<sup>4</sup>), respectively. By composing all four FSTs, we generate a decoding graph *HCLG*,

$$HCLG = H \circ C \circ L \circ G, \tag{7.3}$$

which transduces phone-state sequences directly to word-sequences and output the associated weights. The information provided by such a decoding graph is vital for a speech recognition system to work. Note that the composition operation is associative; therefore, no parentheses are needed in the Equation

<sup>&</sup>lt;sup>4</sup>We may think of *G* as an FSA. However, in practice, we use special symbols in the arcs, so not all arcs have the same input and output symbols. A discussion of that is out of the scope of this thesis. Interested readers may refer to [83] and search the keyword "disambiguation symbol" for details.

7.3 to specify the order of composition.

Readers should refer to [22] for details on FST composition. We give the algorithm for conventional FST-composition in Algorithm 6 as background, and for later showing the changes in algorithm for the pruned composition. We use the algorithm given in Figure 7 from [22], with slight changes for consistency with the notations used in this thesis. We follow the notation in [22] where E(q) represents the set of all arcs leaving state q, and o(e) and i(e) represent the output and input symbol on arc e, respectively; n(e) represents the "next state" of arc e.

Algorithm 6: FST Composition
<b>Input:</b> FST $T_1 = (Q_1, \Sigma_1, I_1, F_1, \delta_1), T_2 = (Q_2, \Sigma_2, I_2, F_2, \delta_2)$
<b>Result:</b> Composition of $T_1$ and $T_2$
$Q = I_1 \times I_2$
2 $\tilde{S}$ = queue storing elements in $I_1 \times I_2$
3 while $S \neq \emptyset$ do
$4   (q_1, q_2) = S.pop()$
5 $if(q_1, q_2) \in I_1 \times I_2$ then
$\begin{bmatrix} 6 \\ I \end{bmatrix} = I \cup \{(q_1, q_2)\}$
7 end
s <b>if</b> $(q_1, q_2) \in F_1 \times F_2$ then
9   $F = F \cup \{(q_1, q_2)\}$
10 end
<b>for</b> $(e_1, e_2) \in E[q_1] \times E[q_2]$ <i>s.t.</i> $o[e_1] = i[e_2]$ <b>do</b>
12 <b>if</b> $(n[e_1], n[e_2]) \notin Q$ <b>then</b>
13 $Q = Q \cup \{(n[e_1], n[e_2])\}$
14 $S.push((n[e_1], n[e_2]))$
15 end
<b>16</b> $E = E \cup \{((q_1, q_2), i[e_1], o[e_2], w[e_1] \otimes w[e_2], (n[e_1], n[e_2]))\}$
17 end
18 end
19 return $T = (Q, \Sigma, I, F, \delta)$

From Algorithm 6, we see that it uses a queue *S* to store pairs of (input-state,

output-state)'s to process and terminates when the queue is eventually empty. Each time we take an element  $(q_1, q_2)$  from the queue, we find all pairs of  $(e_1, e_2)$  such that  $e_1$  leaves from  $q_1$  and  $e_2$  leaves from  $q_2$ , and the output symbol on  $e_1$  is the same as the input symbol on  $e_2$ , and merge those arcs and add a new state and a new arc to the output FST, with appropriate weights. We then add the new states to the queue so they will be processed later.

### 7.4 FST Representation of Lattices

Lattice-rescoring is a standard method for utilizing a recurrent language model in ASR. A *lattice* is a way of representing a set of hypotheses in ASR tasks. Depending on the exact usage scenario, phone-state lattices, phone lattices, and word lattices are used in speech recognition. A lattice is made up of states and arcs and is commonly represented as an FST in speech recognition systems. To differentiate the scores computed from the acoustic model and the language model, when we use FSTs to represent lattices, we usually store a pair of weights on each arc, one representing the acoustic model scores and the other the language model. Given a lattice represented as an FST, if we start from the start-state and follow any arcs sequence to the final-state, then the concatenation of all the symbols on the visited arcs forms a hypothesis. The weight of that hypothesis may be computed as the semiring product  $\otimes$  of the weights on its arcs. One example of a word lattice is shown in Figure 1.2 on page 16. We copy that figure here in Figure 7.3 for easy reference.

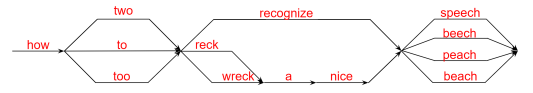


Figure 7.3: Copy of the Word Lattice Example First Shown in Figure 1.2

In this example, the start-state is at the left-most and the final-state the rightmost. We print out the symbols on the arcs but omit the weights to simplify the illustration. In this case, if we follow all the upper-arcs in the figure, we get the hypothesis, "how two recognize speech"; if we follow all the lower-arcs, we get "how to wreck a nice beach". We note that the two sentences sound somewhat similar in terms of pronunciation, and this is no surprise for lattices generated from a speech recognition system because the way to generate the lattice, in plain words, is to find word sequences that "match" the acoustic input (as measured by an acoustic model), that also "makes sense" themselves (as measured by a language model). However, we point out that this is an elementary example of a lattice to illustrate the concept. In practice, depending on the acoustic model, decoding language model, and the beam-size used, the actual lattices generated from ASR systems are usually much more complicated than this.

Table 7.1 shows some statistics on the lattices generate by the standard decoding procedure in Kaldi for different datasets. We report the stats on Switchboard (SWBD), Wall Street Journal (WSJ), and AMI datasets. For WSJ, we report the stats on both Dev93 and Eval92 testsets, and for AMI, we report both Dev and Eval testsets under the single microphone condition. We choose

the default lattice-free MMI model provided in Kaldi and decode with each dataset's default parameters. We report the total number of utterances (#utts), the total number of arcs (#arcs), and the average number of arcs per utterance (#arcs per utt) and the average number of arcs per word (#arcs per word ) across all lattices for those datasets.

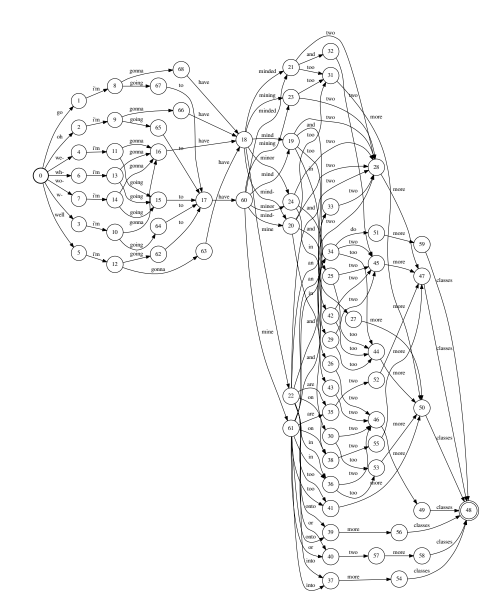
dataset	#utts	#arcs	#arcs per utt	#arcs per word
SWBD-Eval2000	4458	10,593,049	2,376.2	248.0
WSJ-Dev93	503	425,191	845.3	51.0
WSJ-Eval92	333	198 <i>,</i> 530	596.2	34.8
AMI-SDM-Dev	13059	181,935,163	13,931.8	1916.8
AMI-SDM-Eval	12612	208,486,336	16,530.8	2325.9

Table 7.1: Statistics on Kaldi Generated Lattices for Different Datasets

For a visual demonstration, we include an example of a real lattice generated from the SWBD-Eval2000 corpus, with utterance-id en\_4156-A\_030470-030672, as shown in Figure 7.4. The reference text for this utterance is "well I am going to have mine in two more classes". The lattice has 127 arcs, which is around 5% of the average size of lattices for the SWBD-Eval2000 dataset. Nevertheless, we still see that it is highly complex, with many possible paths from the start-state 0 (at the left) to the final state 48 (at the bottom right).

### 7.5 Lattice-rescoring with FST Composition

Lattice-rescoring uses the score computed from an external language model to replace the language model weights on the lattice. As it is usually the case that



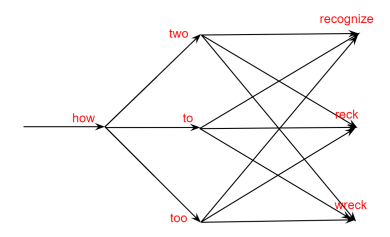
**Figure 7.4:** A Real Lattice for SWBD-Eval2000 Data Generated by Kaldi, Utterance ID: en\_4156-A\_030470-030672. The reference for the utterance is, "well i am going to have mine in two more classes." The lattice has a start-state 0 at the left and a final state 48 at the bottom right, with 127 arcs. The word labels on each arc is shown, but we omit the weights.

the external language model is of higher quality than the one used to generate the initial lattice, this process improves the quality of the lattice and helps reduce the error rates of the hypotheses computed from the lattice. In lattice-rescoring, although the typical use case is when the original lattice is generated with an n-gram language model, and the external language model is a neural language model, this need not be the case – the language model used in the original lattice and the external language model both may take any form. However, in order to see an improvement in the ASR performance, the external language model needs to be "better" than the lattice language model – in practice, besides rescoring with RNNLMs, we sometimes also use a higher-order n-gram model to rescore lattices generated with a lower-order n-gram. For ease of description, we refer to the language model that is used to generate the original lattice as "lattice LM".

There are two ways in which we could understand rescoring, both of which are equivalent.

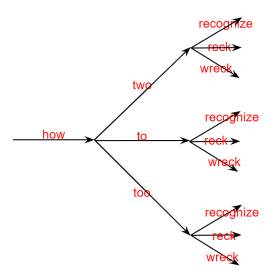
- 1. For all sequences in the lattice, we first remove its lattice LM weights and add new weights computed on-the-fly from the external LM.
- 2. For all sequences in the lattice, we compute its weight difference between the external LM and the lattice LM and add that difference to the lattice.

Because the external language model is usually "better" than the lattice language model at capturing long contexts, the rescoring procedure usually needs to change the lattice's topology. Let us look at the lattice in Figure 7.3 as an example, which we rescore with a *n*-gram language model. Note that in the original lattice, the arcs associated with words "two", "to" and "two" point to the same state; therefore, this state represents three different bi-gram histories. This state then has out-going arcs associated with words "recognize", "wreck" and "reck". If rescored with a bi-gram model, then at this state, nine bi-gram probabilities need to be included in the lattice, and the same word might have different probabilities following different histories.



**Figure 7.5:** Topology of lattice from Figure 7.3 if rescored by a bi-gram model. Note this lattice is partial, and only shows the part that is close to the start-state of the lattice.

Figure 7.5 shows how the lattice topology changes if rescored with a bigram language model. Note it only shows the arcs near the start-state of the FST. We notice that there are significantly more states and more arcs than the corresponding part from the original lattice; now each state corresponds to a distinct bi-gram history, e.g., unlike in Figure 7.3, where the arcs corresponding to words "two", "to" and "too" merge back into the same state, here since their subsequent arcs need to reflect bi-gram probabilities based on different histories, we can not merge those states anymore. We also notice that we still merge states in the rescored lattice, e.g., at the right side of the lattice, all arcs associated with the word "recognize" are pointing to the same state. This is



**Figure 7.6:** Topology of lattice from Figure 7.3 if rescored by an RNNLM. Note this lattice is partial, and only shows the part that is close to the start-state of the lattice.

because, although they associate with different histories ("how two recognize", "how to recognize" and "how too recognize"), a bi-gram model only looks at the last word in the history, and therefore those histories are all equivalent in the bi-gram sense.

Since RNNLMs do not use the *n*-gram assumption, it does not allow any state merging if an RNNLM rescores a lattice. Figure 7.6 shows the lattice topology if we rescore the lattice from Figure 7.3 with an RNNLM. Note that since no state merging is performed, the rescored lattice is expanded like a tree. Note that the example in Figure 7.6 is the result of performing *exact* lattice rescoring; in practice, in order to save computational cost, it is common to use an inexact version of lattice rescoring – we still have an *n*-gram assumption with n = 4,5 or 6, and merge states that represent histories that are equivalent under the *n*-gram assumption.

#### 7.5.1 Exact Lattice Rescoring

Now we introduce the algorithm for performing an exact version of latticerescoring. A naive and straightforward implementation would enumerate all possible sequences in the rescored lattice and individually update the weights on those sequences with an external language model. We notice a lot of repeated computation for different sequences in this implementation for different sequences, e.g., if two or more hypotheses have a common prefix of several words, the computation of the prefix could be re-used. A much more efficient rescoring mechanism could be easily explained using the composition operation of FSTs. The algorithm is shown in Algorithm 7.

Algorithm 7: Performing Lattice Rescoring with FST Composition	
Input: Lattice L, n-gram FST A, RNNLM B	
<b>Result:</b> Rescored Lattice L'	
1 B' = FstWrapper(B)	
$_{2}$ L' = Compose(L, -A, B')	
3 return L'	

On line 2 of Algorithm 7, it uses a composition operation to subtract the lattice language model weights from the original lattice and then add new weights computed from an RNNLM. To add the weights from an RNNLM, we need to perform a composition between an FST and an RNNLM. This is achieved by viewing an RNNLM as a dynamically growing "finite"-state transducer, which in the algorithm we refer to as a "FstWrapper"<sup>5</sup>. This wrapper

<sup>&</sup>lt;sup>5</sup>We emphasize that we can never convert an RNNLM to an actual FST, herefore we added the quotation mark. However, to rescore a lattice, we only need to perform a finite number

has all the same interfaces as a standard FST; the difference is that it takes an RNNLM as input, generates the FST on-the-fly, and only generates the states and arcs needed during the composition process. During the composition process, the algorithm frequently "queries" the FSTWrapper for a score P(w|h) for different (w, h) pairs, and the wrapper computes those scores with the RNNLM on-the-fly.

Note back in Chapter 1 on Page 11, we mentioned that an RNNLM defines two functions, i) a function  $s_t = \delta(s_{t-1}, w_t)$  that defines state-transition of the RNN, and ii) a function  $p(w|s_t) = f(s_t, w)$  that generates a probability distribution over words based on the RNN hidden state. We also mentioned that an initial-state  $s_0$  needs to be associated with an RNNLM as well. The initial-state, along with the two functions above, uniquely defines an RNNLM. Given this background, we give the algorithm that wraps an RNNLM to an FST in Algorithm 8. Note that this algorithm is not a typical algorithm that takes a single input and terminates after computing an output, but rather a procedure that is always running and handling new input queries while updating its internal structures. A major part of the "input" to the algorithm is the new query (h, w) that the algorithm handles in the while loop from line 4 to line 10. To make the algorithm description more illustrative, we do not include this type of input in the "Input" section of the algorithm.

We emphasize here that Algorithm 8 is not a general-purpose wrapper

of computations on the RNNLM to compute language model weights for a finite number of [history, word] pairs, and could indeed convert those scores into an FST for composition.

for an RNNLM, but one that only works in the context of the composition operation defined in Algorithm 7. If this wrapper is used in the composition context described in Algorithm 7, then the map query m[h] on line 5 of the algorithm would always return a valid hidden representation that is added previously on line 8. However, if we use this wrapper in other applications, there is no guarantee that the key h exists in the map m.

**Algorithm 8:** Wrapping an RNNLM to an Dynamically Growing FST **Input:** An RNNLM defined by functions  $\delta$ , f and initial state  $s_0$ . **Output:** Output  $p(w \mid h)$  whenever there is a query. 1 state-map  $m = \{\}$ 2  $s_{bos} = \delta(s_0, <s>)$  $m["<s>"] = s_{bos}$ 4 while receives a query (h, w) do s = m[h]5  $p(w \mid h) = f(s, w)$ 6 new-h = concat(h, w)7 m[new-h] =  $\delta(h, w)$ 8 Create New State new-h 9 Create New Arc (h, new-h, w,  $\log p(w \mid h)$ ) 10 11 **end** 

### 7.5.2 Lattice Rescoring with *n*-gram Approximations

An exact lattice-rescoring algorithm in practice is not feasible because the resulting lattice grows exponentially to the length<sup>6</sup> of the original lattice. Instead of exact lattice-rescoring, usually, an inexact version of lattice-rescoring that adopts an *n*-gram approximation algorithm is used to limit the research space. The algorithm works by not differentiating between history states in the

 $<sup>^{6}</sup>$ By "length" of a lattice, we mean the average distance in terms of the number of arcs from the start state to a final state in the lattice.

RNNLM that are the same in the (n - 1) most recent words.

**Algorithm 9:** Wrapping an RNNLM to an FST with *n*-gram Approximation **Input:** an RNNLM defined by functions  $\delta$ , *f* and initial state  $s_0$ . **Output:** computed  $p(w \mid h)$  whenever there is a query. 1 state-map  $m = \{\}$ 2  $s_{bos} = \delta(s_0, <s>)$  $m["<s>"] = s_{bos}$ 4 while a new query (h, w) do **if** *len*(*h*) > *n* - 1 **then** 5 h = Last-[n-1]-WordsOf(h)6 end 7 new-h = concat(h, w)8 **if** *len(new-h)* > *n* - 1 **then** 9 new-h = Last-[n-1]-WordsOf(new-h) 10 end 11 if *!m.has\_key(new-h)* then 12 CreateState new-h 13 m[new-h] =  $\delta(h, w)$ 14 end 15 if lexist an arc from h to new-h then 16 s = m[h]17  $p(w \mid h) = f(s, w)$ 18 CreateArc (h, new-h, w,  $\log p(w \mid h)$ ) 19 end 20 21 end

The outline for the *n*-gram approximation algorithm is the same as the exact rescoring algorithm, as shown in Algorithm 7. The only difference is in how we wrap the RNNLM into an FST. Now because we no long differentiate all histories for RNNLMs, a more sophisticated wrapper than the one described in Algorithm 8 is needed and shown in Algorithm 9.

Let us first look at exact lattice-rescoring with a simple example. Say the original lattice is the one shown in Figure 7.7 (a), and we perform exact lattice-rescoring, then the FST wrapper result of the RNNLM will eventually have the

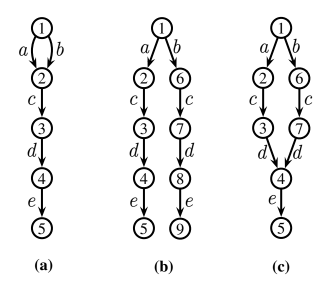


Figure 7.7: Examples of Lattices

topology shown in (b). The states in this wrapped FST are added in Algorithm 8 at line 9, and the arcs are added on line 10. In this case, since all different histories are differentiated, whenever a query (h, w) arrives, the algorithm concatenates h and w, and creates a new state for concat(h, w), and create an arc from state h to concat(h, w). The weights on the arcs accurately reflect the scores computed from RNNLM for the corresponding (h, w) pair.

Figure 7.7 (c) shows the wrapped RNNLM, if we use 3-gram approximation in lattice-rescoring. Compare that with Figure 7.7 (b), we see state 4 and 8 from (b) are merged into a single state 4 (because they share the 2-gram history (*c*, *d*)). This is accomplished in Algorithm 9 through multiple steps. First, whenever a new query (*h*, *w*) arrives, it truncates both histories *h* and *new-h* = *concat*(*h*, *w*), by only retaining last (*n* – 1) words in the histories. Then it checks if a state corresponding to history *new-h* is already created. If not, the algorithm creates such a state and an arc from *h* to *new-h*; otherwise, the algorithm does not create new states or arcs, but "borrow" the weight on the arc from *h* to *new-h* that already exists in the FST. For example, say the algorithm first queries (h = (a, c), w = d), then after truncation, h = (a, c) and *new-h* = (c, d), and state 4 is created, with the weight on arc from state 3 to 4 storing weight  $-\log P_{RNNLM}(d|a,c)$ ; later when the algorithm queries (h = (b,c), w = d), then after truncation, we have h = (b,c) and *new-h* = (c,d). We note that a state corresponding to history (c,d) had already been created (state 4), but an arc from *h* to *new-h* had not, so we skip the state creation, and create a new arc pointing from state 7 to state 4, and the weight on this arc also reflects the correct weight  $-\log P_{RNNLM}(d|b,c)$ .

At this point, if the composition continues, then two more queries,  $(h_1 = (a, c, d), w_1 = e)$  and  $(h_2 = (a, c, d), w_2 = e)$  will be made. Without loss of generality, let's assume that  $(h_1 = (a, c, d), w_1 = e)$  comes first, which will make the algorithm create state 5, as well as an arc from state 4 to state 5, storing weight  $-\log P_{RNNLM}(e|a, c, d)$ ; then when the query  $(h_2 = (a, c, d), w_2 = e)$  is made, with truncation, it becomes a query from history (c, d) to (d, e), with label e. This corresponds to the exact same arc we just created, and thus here, we skip both state and arc creation and borrow the previously computed results. In this case, we are using the previously computed score  $-\log P_{RNNLM}(e|a, c, d)$ to approximate  $-\log P_{RNNLM}(e|b, c, d)$ . Note that if previously, the query  $(h_2 = (a, c, d), w_2 = e)$  comes earlier than  $(h_1 = (a, c, d), w_1 = e)$ , then the opposite result happens, where we will be using score  $-\log P_{RNNLM}(e|b, c, d)$ to approximate  $-\log P_{RNNLM}(e|a, c, d)$ . In the actual composition algorithm, the order in which arcs are processed is arbitrarily chosen, and we see that this could potentially affect the computation of p(e|c, d).

From this analysis, we see that the *n*-gram approximation method is more computationally efficient for it creates fewer states and fewer arcs in the FST wrapper for RNNLM; however, this is achieved at the cost of accuracy – longer-than-*n* history words would not be correctly utilized in RNNLM, and this may result in degradation of performance.

How can we minimize the negative impact of performing the *n*-gram approximation? Note that the goal of lattice-rescoring is to make the lattice represent more accurate scores so that it is more likely that the gold sequence<sup>7</sup> will have better scores than competing hypotheses. Therefore, we would prefer that the golden sequence have all correct scores, and only the competing incorrect hypotheses have inexact scores because of the approximation. Of course, there is no way to know the gold sequence during rescoring, but we may use useful heuristics in the computation to achieve better results than arbitrarily guessing, which is the main topic for the next chapter.

<sup>&</sup>lt;sup>7</sup>By gold sequence, we mean that the "oracle" or correct sequence that the speech audio represents.

## 7.6 Chapter Summary

In this chapter, we gave a brief introduction to finite-state automata (FSA), finite-state transducers (FST), weighted finite-state transducers (WFST), and the composition operations defined on WFSTs, which lays the foundation for the lattice-rescoring algorithm. We analyzed the exact lattice-rescoring algorithm as well as the inexact version with *n*-gram approximations. We use an example to illustrate the potential drawback of the *n*-gram approximation method, which we address in the next chapter.

# Chapter 8

# **Pruned Lattice Rescoring**

In Chapter 7, we introduced basic concepts in FST and used the FST framework in describing the basic methods for lattice-rescoring. Besides the exact version of lattice-rescoring, we have also introduced a commonly-used *n*-gram approximation method for lattice-rescoring in order to speed up the computation. We showed that the *n*-gram approximation method is more computationally efficient than the exact algorithm, but the approximation could degrade latticerescoring performance.

This chapter proposes a pruned version of lattice-rescoring that further improves efficiency over the *n*-gram approximation method. The basic idea of pruning is to use a heuristic score that reflects how "promising" an arc is and always expand the most promising arcs first while completely discarding the least promising arcs. The high-level algorithm is shown in Algorithm 10.

The difference between Algorithm 10 with the non-pruned Algorithm 7 is

Algorithm 10: Lattice Rescoring with FST Pruned Composition
<b>Input:</b> Lattice L, n-gram FST A, RNNLM B
<b>Result:</b> Rescored Lattice L'
$_{1}$ B' = RnnlmWrap(B)
<sup>2</sup> L' = PrunedCompose(L, Compose(-A, B'))
3 return L'

in Line 2, where a special *PrunedCompose* composition is used for the pruned algorithm. Note that in Chapter 7, we mentioned that composition is associative; therefore, no parentheses are needed in Algorithm 7 to specify the order of operation in composition; however, in Algorithm 10, we need to specify the order of operations, where we first perform an on-the-fly composition between the negated lattice LM – *A* and the RNNLMWrapper *B*' and generate a "difference FST". Then we compose the original lattice with this "difference FST" for rescoring. Note that this order of operation only defines the computational dependency, in that during *PrunedCompose*, every bit of information about the difference FST needs to be computed with the *Compose* operation first before being used in *PrunedCompose*; but this does not mean that we first compute DiffFST = Compose(-A, B') and when it terminates, we then perform PrunedCompose(L, DiffFST). The computation of those two composition operations alternates, and both B' and DiffFST grow on-the-fly.

### 8.1 **Pruned composition**

In Algorithm 11, we introduce *pruned composition*. Compared to the standard composition algorithm shown in Algorithm 6, this pruned composition that

Algorithm 11: Pruned FST Composition

**Input:** FST  $T_1$ ,  $T_2$ , beam **Result:** Composition of  $T_1$  and  $T_2$  $1 Q = I_1 \times I_2$ <sup>2</sup> *S* = priority queue storing elements in  $I_1 \times I_2 \times \{0.0\}$ <sup>3</sup> while  $S \neq \emptyset$  do  $(q_1, q_2, p) = S.pop()$ 4 if *p* > *beam* then 5 break 6 end 7 if  $(q_1, q_2) \in I_1 \times I_2$  then 8  $I = I \cup \{(q_1, q_2)\}$ 9  $\lambda(q_1, q_2) = \lambda_1(q_1) \otimes \lambda_2(q_2)$ 10 end 11 if  $(q_1, q_2) \in F_1 \times F_2$  then 12  $F = F \cup \{(q_1, q_2)\}$ 13  $\rho(q_1, q_2) = \rho_1(q_1) \otimes \rho_2(q_2)$ 14 end 15 for  $(e_1, e_2) \in E[q_1] \times E[q_2]$  s.t.  $o[e_1] = i[e_2]$  do 16 if  $(n[e_1], n[e_2]) \notin Q$  then 17  $Q = Q \cup \{ (n[e_1], n[e_2]) \}$ 18  $p = ComputePriority((n[e_1], n[e_2]))$ 19  $S.push((n[e_1], n[e_2]), p)$ 20 end 21  $E = E \cup \{((q_1, q_2), i[e_1], o[e_2], w[e_1] \otimes w[e_2], (n[e_1], n[e_2]))\}$ 22 end 23 24 end 25 return T

we propose is "partial", where not all possible arcs and states are processed, and the composition result is only a sub-graph of the result of the standard composition. In the composition process, we discard certain states and arcs in the output, and the decision for whether to retain an arc depends on a heuristic function we define. Compared with Algorithm 6, which uses a queue to store arc pairs to process, with the pruned algorithm, we define a priority score for elements in the queue, and prioritize arcs with better scores and discard those that are "very bad". This is implemented with a priority queue. The algorithm requires an external function *ComputePriority()*, whose details will be presented in the next section. Assuming the priority score is good, meaning it gives better scores to arcs in the gold sequence, we see the following from the algorithm,

- 1. In lines 5 to 7, by discarding certain arcs, the algorithm can terminate early, reducing the run-time of the composition operation, making lattice-rescoring run faster;
- 2. By adopting a priority queue, the algorithm always prioritizes "more promising" arcs. In the context of lattice-rescoring, using an *n*-gram approximation where state merging is required means that when merging two states, the algorithm picks a "better" history state for the merged state, and thus minimizing the potential negative effects of the approximation in preserving the scores for the best path in the lattice. <sup>1</sup>

## 8.2 Heuristics

From the previous section's analysis, a good heuristics function for arcs in the FST is needed for the pruned rescoring algorithm to work. The ideal heuristic would give good scores to arcs in the gold sequence and bad scores to arcs that are not. However, there is no way to know the gold sequence, and we propose

<sup>&</sup>lt;sup>1</sup>Readers can refer back to the analysis of the *n*-gram approximation example on Page 130, where we show that when merging two states, it is the history that was first computed with the RNNLM that gets stored in the FSTWrapper for RNNLM, and impacts all future computation

to use a proxy for that instead: we assume if we rescore the lattice with an RNNLM without approximation, then the best path in the rescored lattice is most likely to be the gold sequence, and we set up our heuristics in order to preserve this sequence.

### 8.2.1 Assumption

We propose the heuristics based on an assumption we make on the scoring output from a language model. We first present a crucial theoretical result here as the foundation for the heuristic. Let  $M_1$ ,  $M_2$  be two probabilistic language models, i.e. let *S* be the set of all possible sequences, and

$$\sum_{W \in S} P_{M_1}(W) = \sum_{W \in S} P_{M_2}(W) = 1.$$
(8.1)

Following Equation (8.1), we expect that, for any sequence *W*, the difference of probability scores given by the two models for *W* is 0, i.e.

$$\forall W \in S, \mathbb{E}[P_{M_1}(W) - P_{M_2}(W)] = 0.$$
 (8.2)

In the pruning algorithm, the actual assumption we use is,

$$\forall W \in S, \mathbb{E}[\log P_{M_1}(W) - \log P_{M_2}(W)] \approx 0.$$
(8.3)

Note that while we can easily prove Equation (8.2) following (8.1), we

have to use "approximately equal" in Equation (8.3) because mathematical expectations do not carry through a non-linear log function; however we have empirically found that Equation (8.3) is a good approximation and is useful in practice. The consequences of the assumption in Equation (8.3) is, the expected difference of total weight of the original *n*-gram model and the external RNNLM is 0, hence when a lattice is rescored with the external RNNLM, we would see some paths get their weights increased, and others decreased, but the overall changes in weights from the to the original lattice is close to 0.

### 8.2.2 Background: $\alpha$ and $\beta$ Scores

Before introducing the algorithm, we first need to define some terminology. Note some of the terminologies are very similar to HMM terminologies, and here we borrow them in the context of a word-lattice. For simplicity, in this section, we assume we use the tropical-semiring [84] in all the FSTs <sup>2</sup>. Under this semiring, when we say a weight  $w_1$  is better than  $w_2$ , we mean  $w_1 < w_2$ ; the "best" path of a lattice is the path that has the lowest weight among all possible paths in the lattice.

#### **Forward Weights**

For each state, we define its *forward weight*, which we call  $\alpha$  score as "the negative log-probability of the best path from a start-state to this state". It is

<sup>&</sup>lt;sup>2</sup>In particular, we use the *min tropical semiring* where weights are interpreted as negative log-probabilities.

defined recursively as,

$$\alpha(s) = \begin{cases} 0, & \text{if } s \text{ is a start state;} \\ \\ \min_{[i,s] \in A} \Big[ \alpha(i) + w([i,s]) \Big], & \text{otherwise.} \end{cases}$$
(8.4)

where [i, s] represents an arc from state i to s, A the set of all arcs and w([i, s]) represents the weight of the arc [i, s]. Since we can topologically sort the states in an FST-represented lattice, Equation (8.4) can be implemented by iterating the sorted list and for any state, we need only check its antecedent states to iterate all its incoming arcs.

#### **Backward Weights**

We define the *backward weight* for a state, which we call a  $\beta$  score as "the negative log-probability of the best path from this state to any final-state". Just like the forward weights, it is also defined recursively,

$$\beta(s) = \begin{cases} 0, & \text{if } s \text{ is a final state;} \\ +\infty, & \text{if a final state is not yet reachable from } s \\ \min_{[s,i] \in A} \left[ \beta(i) + w([s,i]) \right], & \text{otherwise.} \end{cases}$$

$$(8.5)$$

Similarly, we can also compute this quantity efficiently by taking advantage of the topologically sorting of the FST states.

#### **Properties of the Forward and Backward Weights**

For any arc with starting-state  $s_1$  and destination state  $s_2$ , and the weight on the arc w, if we compute the following,

$$\alpha(s_1) + w + \beta(s_2). \tag{8.6}$$

Since  $\alpha(s_1)$  is the weight of the best path from a start state to  $s_1$ , and  $\beta(s_2)$  is the weight of the best path from  $s_1$  to a final state, it naturally follows that the quantity in Equation (8.6) stores the weight of the best path from a state to a final state, with the constraint that this path passes through the arc from  $s_1$  to  $s_2$ . Therefore, if we compute  $\alpha(s_1) + w + \beta(s_2)$  for any arc on the best-path of a lattice, they are all equal to the weight of the best path; and the quantity for any other arc in the lattice would be worse (larger) than the weight of the best path.

### 8.2.3 Heuristic

Before talking about the heuristic, let us introduce some terminology in describing the rescoring process. Note that we are simultaneously dealing with two lattices during the rescoring process, i.e., the original lattice whose arcs store lattice language model weights and a rescored lattice which grows in the rescoring process. We call the original lattice *A* and the rescored lattice *C* in this section's description. Note that as *C* grows during rescoring, each state  $c \in C$  corresponds to a state  $a \in A$ , which we call *c*'s "source state". Similarly, an arc  $(s_1, s_2, x, w) \in C$  also corresponds to a "source arc" in A as well<sup>3</sup>.

The heuristic we propose is computed for each arc  $c \in C$ . Say arc c is from  $s_1$  to  $s_2$  with weight w, then the heuristic tries to approximate the value  $\alpha_C(s_1) + w + \beta_C(s_2)$ , where the smaller the value, the higher its priority.

The first thing we notice is that, while all the forward scores for each lattice state is easy to compute while the lattice grows, it is not the case for backward scores. From the definition, the backward score for a state *s* is only well-defined when a final state is reachable from *s* or mathematically when there exists a sequence of arcs that connect *s* to a final state. However, in the earlier stages of rescoring a lattice, a final state is usually not reachable from any state in the output lattice, and therefore we need to handle this case and estimate the backward score.

We propose to use the assumption in Equation (8.3) in order to approximate backward scores for all states. Let's first re-write  $\beta_C(c)$  as,

$$\beta_C(c) = \beta_A(a) + \delta(c),$$

where *a* is the state in *A* corresponding to *c*, and

$$\delta(c) = \beta_C(c) - \beta_A(a).$$

<sup>&</sup>lt;sup>3</sup>Here we slightly abuse the meaning of  $\in$ . When we say a state is "in" an FST *A*, we mean it is in the set of states associated with *A*; similarly, when we say an arc is "in" an FST, we mean it is in the set of arcs associated with the FST.

In other words,  $\delta(c)$  is the difference between the two backward scores.

Now, we propose to use the  $\tilde{\delta}(c)$  defined in Equation (8.7) to recursively estimate the value of  $\delta(c)$ , using approximations when necessary.

$$\tilde{\delta}(c) = \begin{cases} \beta_{C}(c) - \beta_{A}(a), & \text{if } \beta_{C}(c) < +\infty; \\ 0, & \text{if } \beta_{C}(c) = +\infty \text{ and } c \text{ is a start state;} \\ \tilde{\delta}(\text{prev}(c)), & \text{otherwise.} \end{cases}$$
(8.7)

We note that in order to compute the  $\delta$ , there are three cases to consider. In the first case, we have a well-defined  $\beta_C(c)$ , and therefore we compute its exact value and there is no need to perform any approximation. In the second case,  $\beta_C(c)$  is infinity and c is a start-state. According to the assumption from Equation (8.3), we estimate the difference between the two beta scores as 0, since no information indicates either one language model gives better scores than the other; in the third case, we have an infinite  $\beta_C(c)$  again, but c is not a start state. In this case, we go back to the previous state of c and borrow the delta score from that state instead. The rationale is since prev(c) is the closest state to c, and therefore c and prev(c) should have a very similar difference in their backward scores.

Given the definition of forward and backward scores in Section 8.2.2, and the definition of  $\tilde{\delta}(c)$  in Equation 8.7, now we give the heuristic function H(.)

for a state  $c = s_1, s_2, x, w$  in the output lattice, in Equation (8.8),

$$H(c) = \alpha_{\rm C}(s_1) + w + \beta_A(s_2) + \tilde{\delta}(s_2), \tag{8.8}$$

where  $\tilde{\delta}(s_2)$  is defined in Equation (8.7).

## 8.3 Applying the Heuristics in Composition

Combining Algorithm 11 and the heuristic function defined in Equation (8.8), the pruned version of FST-based lattice-rescoring is ready to run. This section proposes several methods we use to ensure this algorithm's high performance and efficiency.

### 8.3.1 Lazy Updates of Forward/Backward Scores

We point out that to ensure the algorithm's correctness, we need to keep updating the forward and backward scores of all states in the output lattice whenever a new state is added to it. Although each update runs relatively fast (its time complexity is linear to lattice size), carrying out the computation repeatedly is still quite costly. Suppose the final rescored lattice has *n* states. Since the output lattice has to grow from size 0 and add states one-by-one, and each time a new state is added, we need to update all the forward and backward scores, the total computational cost becomes,

$$O(1) + (2) + O(3) + \dots + O(n) = O(\frac{n(n+1)}{2}) = O(n^2)$$

We propose using a workaround to reduce the computational overhead significantly: we only periodically update the forward and backward values for output lattice states. Note, the consequence of this design is that most of the time, we use stale values of these quantities that may be slightly worse than the exact value.

In designing the schedule to update those values, our goal is to limit the added overhead to linear to the final lattice size. To achieve that, we pre-define a constant  $\lambda > 1$  and follow the update schedule shown in Algorithm 12. In this algorithm, the schedule monitors the total size of the output lattices and recomputes everything if the current size is  $\lambda$  times the previous size, where  $\lambda$  is a constant greater than 1, and we choose  $\lambda = 1.25$  in practice.

Algorithm 12: Scheduling for Updating Alpha's and Beta's
<b>Input:</b> A constant $\lambda$ , and the output lattice L
1 n = NumStates(L)
2 current_limit = $\mathbf{n} \cdot \lambda$
<b>3 while</b> <i>a new state is added to output lattice L</i> <b>do</b>
4  n = NumStates(L)
5 <b>if</b> <i>n</i> > <i>current_limit</i> <b>then</b>
6 current_limit = current_limit $\cdot \lambda$
7 UpdateAlphaBetas()
8 end
9 end

To analyze this schedule's time complexity, let us assume that the final size

of the output lattice is n, then the function UpdateAlphaBetas() on line 7 will be called  $\lceil \log_{\lambda} n \rceil$  times. The size of the lattice each time the UpdateAlphaBeta() is called grows exponentially. Since the computational complexity of one call to UpdateAlphaBeta() is linear to the lattice-size at the time of call, the total cost is

$$O(1) + O(\lambda) + O(\lambda^2) + \dots + O(\lambda^{\lceil \log_{\lambda} n \rceil - 1}).$$

By using the Equation for computing summation of geometric series, the total cost equals to

$$O(\frac{\lambda^{\lceil \log_{\lambda} n \rceil - 1} - 1}{\lambda - 1}) = O(\frac{n}{\lambda - 1}) = O(n)$$

Therefore, we conclude that the schedule we propose to update all the forward and backward scores only adds a linear (to the final lattice size) computational overhead to the rescoring process.

#### 8.3.2 Initial Computation

We notice that the algorithm does not always speed up the lattice rescoring computation if the lattice is huge during our initial experiments with the pruning algorithm. Our investigation reveals that the reason is that Equation 8.3 may be interpreted as a comparison of the entropy of an *n*-gram model and an RNNLM. We have empirically observed that RNNs tend to give sharper output

distributions, while *n*-grams tend to give more smooth distributions. Therefore, the RNNLMs usually have smaller average entropy than *n*-grams. This difference, therefore, accumulates on longer sequences and leads to a poorer heuristic. We work around this problem by changing the initial computation of the algorithm. We first always go along the original lattice's best path and replace the *n*-gram scores with RNNLM scores. Once this path is visited, the states along this best path all have well-defined backward scores, and the states visited afterward have a better estimate of their backward scores by using case 3 of Equation 8.7.

## 8.4 Experiments

In this section, we report experimental results of running the pruned latticerescoring. We compare it with the standard *n*-gram approximation algorithm without pruning, both in terms of ASR performance as well as rescoring speed. All experiments are conducted using the open-source speech recognition toolkit Kaldi.

### 8.4.1 Rescoring Speed and Output Lattice Size

We first report how much the pruning algorithm could reduce the computation of the rescoring procedure. We first show the rescoring speed for different rescoring algorithms. We pick the AMI datasets for experiments and compare the algorithms' speed; we report the average time (in seconds) it takes to rescore all lattice generated for the development set, divided into 30 roughly equal-sized jobs.

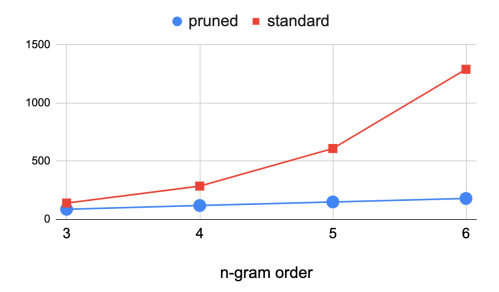


Figure 8.1: Average run-time (in seconds) of lattice-rescoring, AMI-DEV

We plot the run-time and output lattice size of different rescoring algorithms in Figures 8.1 and 8.2, where for the pruned algorithm, we use beam-size = 6. We use *n*-gram approximation to limit the search space for both the pruned and unpruned version of the rescoring algorithms. In both figures, we use blue lines with square-shaped dots representing the algorithm with lattice pruning and red lines with circled dots to represent the standard algorithm. We see from both of those figures that, for the standard rescoring algorithm, the running time and output lattice size grow exponentially to *n*-gram order (roughly doubling whenever the *n*-gram order is incremented by one), while the growth seems linear for the pruned algorithm. As a result, the gap between

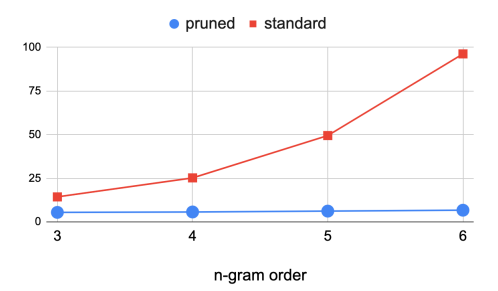


Figure 8.2: Average number of arcs per frame of rescored lattices, AMI-DEV

the two lines in both figures grows larger for larger *n*-gram orders.

While Figures 8.1 and 8.2 only show the experimental results with beamsize = 6, we report number for different beam-sizes in Table 8.1. We experiment with different beam-sizes for the pruned algorithm, including 2, 4, 6, and 8, and use *n*-gram approximation to limit the search space, with *n* being 3, 4, 5, 6. We also report numbers when we do not use *n*-gram approximation ( $n = +\infty$ ) and the standard rescoring algorithm (no pruning). In the case of no pruning and not using any *n*-gram approximation, the experiment was unable to finish due to memory exhaustion, so that cell in the table is deliberately left blank.

From Table 8.1, we have the following observations,

1. Using an *n*-gram approximation significantly reduces the running time of the rescoring procedure;

n-gram order	beam=2	beam=4	beam=6	beam=8	no prune
3	29.06	49.96	86.92	111.96	140.42
4	31.76	62.64	119.26	168.9	286.05
5	34.48	72.18	149.9	230.22	609.52
6	37.54	85.46	179.9	302.86	1290.78
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	170.32	660.22	1840.2	3575.92	-

Table 8.1: Speed (seconds) Comparison of Lattice-rescoring, AMI-DEV

2. Using a lower beam-size also has a noticeable effect in reducing the running time. In particular, when running a 5-gram approximation, using a beam-size = 4 gives almost 10X speed-up for the algorithm, compared to the standard rescoring without using pruning; and when using a 6-gram approximation, this speed-up is 15X.

n-gram order	beam=2	beam=4	beam=6	beam=8	no prune
3	1.44	2.70	5.44	8.11	14.32
4	1.45	2.74	5.68	8.97	25.22
5	1.48	2.89	6.17	10.4	49.48
6	1.52	3.04	6.69	11.79	96.30
$\infty$	1.78	6.13	19.99	43.19	-

Table 8.2: Output Lattice-size Comparison of Lattice-rescoring, AMI-DEV

We also report the size of the output lattice after the rescoring procedure in Table 8.2. Note that since the pruned rescoring algorithm is mathematically equivalent to incomplete/partial composition of FSTs, which discards arcs when processing based on the beam, this makes the output lattice smaller than rescoring without pruning. For the number reported here, the experiment setting is the same as Table 8.1. We measure the size of a lattice by its average number of arcs per frame, computed by the lattice-depth-per-frame binary of Kaldi. We also see very similar trends in the impact on lattice-size compared to rescoring speed. We notice that the reduction of size with beam-size 4, for 5-gram and 6-gram approximation is 17X and 31X, compared to 10X and 15X in speed. This discrepancy can be explained by the computational overhead of the pruned algorithm required to compute all the forward and backward scores. We see that this overhead is outweighed by the pruning algorithm's speed improvement, consistently for all configurations from the reported results.

#### 8.4.2 WER performances

In the previous section, we have shown that the pruned rescoring brings speedup to the rescoring procedure. However, does this speed-up come free, or do we have to sacrifice the performance in order for the algorithm to run faster? In this section, we answer that question by reporting experimental results on several datasets comparing the pruned rescoring with the standard rescoring algorithm in terms of ASR performance.

In Table 8.3, we compare the WERs of different rescoring results as well as the un-rescored baseline. The baseline is decoded from a 3-gram LM. During lattice-rescoring, the 3-gram model weight is interpolated with the computed RNNLM weight, and the interpolation weight is optimized for each corpus. As we can see, in all recipes, our pruning method achieves better WERs than the standard algorithm for all *n*-gram orders.

Corpus	Test set	<i>n</i> -gram only baseline	RNNLM with <i>n-</i> gr 3-gram standard pruned		ram approximation 4-gram standard pruned	
AMI	dev	24.2	23.7	23.4	23.4	23.3
	eval	25.4	24.6	24.4	24.3	24.2
SWBD	swbd	8.1	7.4	7.2	7.2	7.1
	eval2000	12.4	11.7	11.5	11.5	11.3
WSJ	dev93	7.6	6.4	6.2	6.4	6.2
	eval92	5.1	4.1	3.9	3.9	3.8
LIB	test-clean	6.0	4.9	4.8	4.8	4.7
	test-other	15.0	12.7	12.4	12.4	12.3
	dev-clean	5.7	4.4	4.3	<b>4.3</b>	4.3
	dev-other	14.5	12.3	12.0	11.9	11.7

 Table 8.3: WER of Lattice-rescoring of Different RNNLMs in Different Datasets

Combining the results in Table 8.3, Figures 8.1 and 8.2, we conclude that the proposed pruned lattice-rescoring algorithm runs much faster, greatly reduces the output lattice size, and improves ASR performances compared to the standard algorithm.

## 8.5 Chapter Summary

This chapter proposed a pruned version of lattice-rescoring, implemented as a pruned composition for FSTs. We compute the forward and backward scores for lattices and propose a heuristic score that uses those scores. With the heuristics defined, we implement the composition algorithm that prioritizes more promising arcs while discarding arcs that are not promising. We carefully design the update schedule for the forward and backward scores so that it only adds a linear computational overhead to the rescoring process. We show from experiments that the proposed algorithm makes the rescoring process more efficient and improves the quality of the rescored lattice and helps reduce word-error-rates on several datasets.

# Chapter 9

# Conclusion

This dissertation investigates improving neural language models in the context of automatic speech recognition, focusing on speeding up training, inference, and the application of neural LM in ASR, without degradation in ASR performance.

We propose a new training loss function, which we call *linear loss*, which is inspired by taking a first-order Taylor expansion of the cross-entropy objective. We have shown that the linear loss slightly outperforms the standard cross-entropy loss in terms of model performance; it also trains the network to self-normalize, which alleviates the need for normalization during inference. Experiments also show that the linear loss outperforms the varianceregularization loss, another commonly used self-normalizing loss for training language models.

We propose to use sampling-based techniques to speed up linear-loss LM

training. We have theoretically shown that our proposed linear loss function, combined with importance-sampling techniques, yields an unbiased estimator of the actual loss. We compare with noise-contrastive estimation and show that our method is superior.

We study the impact of sampling algorithms on LM training. The usual choice of such a sampling algorithm is a *sampling with replacement* procedure, which could have duplicates of samples and waste some of the computation. We propose to use a *sampling without replacement* procedure for LM training, ensuring there are no duplicates in the sample, and so we can fully utilize the computational resources. To make the method mathematically sound, we propose a method to normalize a probability vector for a set of vocabulary to an "inclusion probability vector" to deal with words with very high probabilities. We have shown with experiments on multiple datasets that *sampling without replacement* consistently gives better results.

When dealing with a large vocabulary (of up to hundreds of thousands of words), the sampling algorithm might become a non-negligible computational overhead during the LM training, so we study ways to speed up the algorithm. We study two types of implementations of the sampling without replacement procedure and propose a novel 2-*stage* sampling algorithm that dramatically reduces the run-time of the sampling algorithm, as shown in experiments. We also show in experiments that the 2-stage sampling achieves speed-up at no cost of training performance.

We study an important method to utilize neural language models for ASR – lattice rescoring. We show that even with the commonly used *n*-gram approximation techniques, the lattice rescoring procedure still is computationally inefficient, and the approximation would result in degradation in performance; we propose a *pruned lattice rescoring* method, which uses a heuristic to guide the search, and discards non-promising paths during the rescoring process, resulting in an algorithm that is significantly faster than the *n*-gram approximation baselines but also achieves better performance.

To sum up, here are the major contributions of this dissertation.

- 1. We propose a new loss function for neural language model training, which we call *linear loss*;
- 2. We show that the linear loss could greatly reduce inference computation for not having to normalize the output;
- 3. We show that the linear loss slightly outperforms the standard crossentropy loss in experiments;
- 4. We propose an importance-sampling scheme that works with the linear loss, and show the superiority of this method over the commonly used alternative method noise-contrastive estimation;
- 5. We propose several improvements over the sampling-based training methods, including using the sampling without replacement scheme, utilizing longer histories in sampling, and 2-stage sampling. These methods

either improve the performance of trained models or improve computational efficiency during model training.

6. We propose a pruned version of lattice-rescoring that utilizes a neural language model. We show that the pruned algorithm runs significantly faster than the standard (non-pruned) version and brings gains in ASR performance measured by word-error-rates.

These claims have been validated by theoretical analysis where appropriate and by empirical results on several data sets commonly used in ASR. The implementation of these algorithms and procedures is made open-source.

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# Vita

Hainan Xu received his bachelor's degree in Software Engineering from Shanghai Jiaotong University, in the beautiful city of Shanghai, 300 miles from his hometown in Lianyungang, Jiangsu, China. Inspired by Prof. Kai Yu at the Speech Group at Shanghai Jiaotong University, Hainan began a Ph.D. in Computer Science at Johns Hopkins University in 2013, advised by Prof. Daniel Povey and Prof. Sanjeev Khudanpur, working on speech recognition. Hainan is a contributor to the open-source ASR Kaldi toolkit. He has worked on acoustic modeling, pronunciation and silence modeling, and various aspects of language modeling with speech recognition applications. Hainan has also worked on end-to-end sequence modeling for speech recognition and machine translation, advised by Prof. Shinji Watanabe and Prof. Phillip Koehn. He is a contributor to the open-source end-to-end ASR Espresso toolkit and the main contributor for the machine translation data-cleaning toolkit Zipporah. In 2015, he interned at the speech team at Google in New York City, supervised by Dr. Cyril Allauzen and Dr. Michael Riley. In September 2019, Hainan became a full-time software engineer at the speech team at Google in NYC, working

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