ABSTRACT

Semi-supervised learning has demonstrated promising results in automatic speech recognition (ASR) by self-training using a seed ASR model with pseudo-labels generated for unlabeled data. The effectiveness of this approach largely relies on the pseudo-label accuracy, for which typically only the 1-best ASR hypothesis is used. However, alternative ASR hypotheses of an N-best list can provide more accurate labels for an unlabeled speech utterance and also reflect uncertainties of the seed ASR model. In this paper, we propose a generalized form of the connectionist temporal classification (CTC) objective that accepts a graph representation of the training targets. The newly proposed graph-based temporal classification (GTC) objective is applied for self-training with WFST-based supervision, which is generated from an N-best list of pseudo-labels. In this setup, GTC is used to learn not only a temporal alignment, similarly to CTC, but also a label alignment to obtain the optimal pseudo-label sequence from the weighted graph. Results show that this approach can effectively exploit an N-best list of pseudo-labels with associated scores, outperforming standard pseudo-labeling by a large margin, with ASR results close to an oracle experiment in which the best hypotheses of the N-best lists are selected manually.

Index Terms — graph-based temporal classification, semi-supervised ASR, pseudo-labeling, self-training, WFST

1. INTRODUCTION

Over the last decade, automatic speech recognition (ASR) technologies have progressed to a point where ASR has become a central user interface for various electronic devices. This progress can largely be dedicated to improved acoustic models using more sophisticated neural network architectures, as well as to improved sequence-to-sequence modeling approaches, known as end-to-end ASR, which have greatly contributed to optimizing the training and decoding process of ASR systems [1-3]. However, to achieve state-of-the-art performance, end-to-end ASR models are generally more data hungry compared to traditional hybrid HMM-DNN ASR solutions [4]. Although the available amount of manually transcribed training data has grown over the years, a lack of training data still exists, especially for low-resource domains and languages. To alleviate these problems, data augmentation as well as self- and semi-supervised learning can be applied in order to utilize unlabeled data for training. In self-supervised learning, typically a latent representation of speech is learned from unlabeled data to pre-train a neural network model for a following supervised or semi-supervised learning step [5-7]. Semi-supervised learning approaches exploit some amount of labeled data to support learning from unlabeled data [8-9]. Self-training uses a seed ASR model, trained from transcribed data, to generate pseudo-labels for unlabeled data for further training [10-13]. This approach has recently become increasingly popular in end-to-end ASR as it has show promising results [14-16]. While most pseudo-labeling approaches only exploit the best ASR hypothesis for an unlabeled data set, alternative ASR hypotheses can encode information about uncertainties, which can be useful [13], as well as pseudo-labels with potentially fewer errors. In [17], an N-best list of ASR hypotheses is used by summing over the weighted losses of multiple pseudo-labels for a single speech utterance, where weights are estimated from scores of a strong language model (LM). In [18], multiple pseudo-labels are generated for each unlabeled speech utterance using different dropout settings, which are used for self-training with the purpose of capturing ASR uncertainties.

In this work, we propose a new objective function termed graph-based temporal classification (GTC), which generalizes the popular connectionist temporal classification (CTC) loss function [19] to accept a graph representation as an input for modeling user-defined output symbol structures, including the possibility to assign probabilities to the generated output symbol sequences through transition weights. The proposed GTC loss function can be used to model all possible variations of CTC, including Gram-CTC [20], the automatic segmentation criterion (ASG) [21], and other not yet explored structures that can be modeled using weighted finite automata.

We apply the GTC loss to the semi-supervised learning problem, where we generate a WFST-based graph from an N-best list of pseudo-labels in order to leverage the information contained in alternative hypotheses as well as in their ASR scores. In such a setup, GTC is expected to find not only the best temporal alignments, similar to CTC, but also an optimal label sequence encoded in such a graph. Note that self-training with lattice-based supervision was also proposed in [22] using a hybrid ASR system and the LF-MMI objective in order to incorporate frame-level confidence scores and alternate word pronunciations for the 1-best word sequence. However, we here consider the more general case of alternative transcriptions using a GTC-based end-to-end ASR system.

ASR results demonstrate that the proposed GTC-based semi-supervised learning approach can effectively utilize an N-best list of ASR hypotheses for self-training by achieving substantial improvements over the commonly used 1-best pseudo-labeling approach.

2. GRAPH-BASED TEMPORAL CLASSIFICATION

Let us consider a feature sequence $X$ of length $T'$ derived from a speech utterance, processed by a neural network to output a sequence $Y$ of length $T$ potentially different from $T'$ due to downsampling, and an associated graph $G$ that is used to form the output target during training. The graph-based temporal classification (GTC) objective function marginalizes over all possible node sequences that can be output by graph $G$, which includes all valid node patterns as well as all valid temporal alignment paths, the latter being known from CTC [19]. Thus, the conditional probability for a given graph $G$ is defined by the sum over all node sequences in $G$, which can be writ-
We denote by \(g\) can be computed for any node sequences of length \(T + 2\) (including non-emitting start and end nodes), \(\pi\) denotes a single node sequence and alignment path, and \(p(\pi|X)\) is the posterior probability for the path \(\pi\) given feature sequence \(X\).

We introduce a few more notations that will be useful to derive \(p(G|X)\). We assume that an acoustic model, e.g., a neural network model, transforms the feature sequence \(X\) into a posterior probability distribution sequence \(Y = (y^1, \ldots, y^T)\), where \(y^t\) denotes the vector of posterior probabilities and \(y_k^t\) the posterior probability for output symbol \(k\) at time \(t\). We index the nodes using

\[\pi_{t:t'} = (\pi_t, \ldots, \pi_{t'})\]  

the node sub-sequence of \(\pi\) from \(t\) to \(t'\). Note that \(\pi_0\) and \(\pi_{T+1}\) correspond to the non-emitting start and end nodes 0 and \(G + 1\).

In CTC, the conditional probabilities \(p(l|X)\) for a given label sequence \(l\) are computed efficiently by a dynamic programming algorithm, which is based on computing the forward and backward variables and stitching both together to compute \(p(l|X)\) at any given time index \(t\). In a similar fashion, the GTC forward probability can be computed for \(g = 1, \ldots, G\) using

\[
\alpha_t(g) = \sum_{\pi \in \mathcal{S}(G, T) : \pi_0 = g} \sum_{\tau = 1}^T W(\pi_{\tau-1}, \pi_{\tau}) y_k^\tau(\pi_{\tau}),
\]

where \(G_{0,g}\) denotes the sub-graph of \(G\) starting at node 0 and terminating at node \(g\). The sum is taken over all possible \(\pi\) whose sub-sequence up to time index \(t\) can be generated in \(t\) steps from the sub-graph \(G_{0,g}\). The backward variable \(\beta\) is computed similarly for \(g = 1, \ldots, G\) using

\[
\beta_t(g) = \sum_{\pi \in \mathcal{S}(G, T) : \pi_T + 1 = g} \sum_{\tau = T}^T W(\pi_T, \pi_{\tau+1}) y_k^{\tau}(\pi_{\tau+1}),
\]

where \(G_{G+1, G+1}\) denotes the sub-graph of \(G\) starting at node \(g\) and terminating at node \(G + 1\). By using the forward and backward variables, the probability function \(p(G|X)\) can be computed for any \(G\) by summing over all \(g\):

\[
p(G|X) = \sum_{g \in G} \alpha_t(g) \beta_t(g) y_k^{\tau}(g) / y_l^{\tau}(g).
\]

For gradient descent training, the loss function

\[
\mathcal{L} = -\ln p(G|X)
\]

must be differentiated with respect to the network outputs, which can be written as:

\[
-\frac{\partial \ln p(G|X)}{\partial y_k^t} = -\frac{1}{p(G|X)} \frac{\partial p(G|X)}{\partial y_k^t},
\]

for any symbol \(k \in \mathcal{U}\), where \(\mathcal{U}\) denotes a set of all possible output symbols.

Because \(\alpha_t(g) \beta_t(g) / y_l^{\tau}(g)\) is proportional to \(y_k^{\tau}(g)\),

\[
\frac{\partial \ln p(G|X)}{\partial y_k^t} = \frac{\alpha_t(g) \beta_t(g)}{y_l^{\tau}(g)^2},
\]

and from (4), we can derive

\[
\frac{\partial p(G|X)}{\partial y_k^t} = \frac{1}{y_k^t} \sum_{g \in \Psi(G, k)} \alpha_t(g) \beta_t(g),
\]

where \(\Psi(G, k) = \{g \in G : l(g) = k\}\) denotes the set of nodes in \(G\) at which symbol \(k\) is observed.

To backpropagate the gradients through the softmax function, we need the derivative with respect to the unnormalized network outputs \(u_k^t\) before the softmax is applied, which is

\[
-\frac{\partial \ln p(G|X)}{\partial u_k^t} = -\sum_{k' \in \mathcal{U}} \frac{\partial \ln p(G|X)}{\partial y_k^t} \frac{\partial y_k^t}{\partial u_k^t}.
\]

By substituting (3) and the derivative of the softmax function \(\partial y_k^t / \partial u_k^t = y_k^t - y_k^t / y_l^{\tau}(g)\) into (5), we finally derive

\[
\frac{\partial \ln p(G|X)}{\partial u_k^t} = 1 - \frac{1}{y_k^t p(G|X)} \sum_{g \in \Psi(G, k)} \alpha_t(g) \beta_t(g),
\]

where we used the fact that

\[
\sum_{k' \in \mathcal{U}} \frac{1}{y_k^t} \sum_{g \in \Psi(G, k')} \alpha_t(g) \beta_t(g) y_k^t / y_l^{\tau}(g) = \sum_{g \in \Psi(G, k)} \alpha_t(g) \beta_t(g) / y_l^{\tau}(g) = p(G|X),
\]

and

\[
\sum_{k' \in \mathcal{U}} \frac{\partial \ln p(G|X)}{\partial u_k^t} y_k^t / y_l^{\tau}(g) = 1 / p(G|X) p(G|X) y_k^t = y_k^t.
\]

For efficiency reasons, we implemented the GTC objective in CUDA as an extension for PyTorch.

3. Graph Generation for Self-Training

In the supervised case, GTC can be used to train an ASR system similarly to CTC by setting the graph to insert blank symbols between the labels of the ground truth transcription and include CTC-like transition rules [19]. However, we here wish to consider more general cases in which there may be multiple transcriptions obtained from a seed ASR system for an unlabeled utterance. The proposed GTC loss makes it possible to learn the model parameters with label information in a graph format. In a semi-supervised ASR scenario, we can use \(N\)-best hypotheses or word/token lattices as pseudo labels, which are typically given by a baseline system trained with a small amount of labeled data. As an example, a graph representation of an \(N\)-best hypotheses list is shown in Fig 1. For computational reasons, it is preferable to make the graph compact while retaining correct predictions as much as possible.

In this work, we generate a compact CTC-like graph from \(N\)-best hypotheses according to the following steps:

1. Convert the \(N\)-best hypotheses to a sausage-form confusion network (CN) using minimum Bayes risk decoding [23].
In Step 1, a scaling factor $\mu$ can be applied to the ASR score (log probability) of each hypothesis, where the scaling factor indicates the degree to which the ASR scores are reflected in the label probabilities in the CN: $\mu = 1$ means the ASR scores are used without alteration, and $\mu = 0$ means the hypotheses are treated equally without considering the ASR scores. We optionally add pruning steps after steps 1 and 2 to reduce the size of the CTC graph, which eliminate arcs if the assigned probabilities are less than a threshold $\eta$.

4. EXPERIMENTS

4.1. Dataset

We use as the ASR benchmark the LibriSpeech corpus of read English audio books, which provides about 960 hours of training data, 10.7 hours of development data, and 10.5 hours of test data. The development and test data sets are both split into approximately two halves named “clean” and “other”. This separation is based on the quality of the recorded speech utterances, which was assessed using an ASR system. The training data is also split into three subsets: “clean” 100 hours, “clean” 360 hours, and “other” 500 hours. We use the “clean” 100 hours subset for supervised training and consider the remaining 860 hours as unlabeled data.

4.2. ASR System

Figure 2 illustrates the ASR system used in this work, a transformer-based neural network architecture that employs the proposed GTC loss function of Section 2 for training. We use 80-dimensional log-mel spectral energies plus 3 extra features for pitch information as acoustic features input to the neural network. The neural network architecture consists of a two-layer convolutional neural network (CNN) module followed by a stack of $F = 12$ transformer-based encoder layers with a projection layer plus softmax function at the end to map the neural network output to a posterior probability distribution. Each layer of the 2-layer CNN module is used as a stride of 2, a kernel size of $3 \times 3, 320$ channels, and a rectified linear unit (ReLU) activation function. In addition, a linear neural network layer is applied to the output of the last CNN layer. Sinusoidal positional encodings are added to the 2-layer CNN module output before feeding it to the transformer-based encoder. Each transformer layer employs 320-dimensional self-attention layers with 4 attention heads, layer normalization, and a feed-forward neural network module of inner dimension 1540. Residual connections are applied to the self-attention and feed-forward module outputs. Dropout with a probability of 0.1 is used after self-attention and after the feed-forward module as well as for the inner dimension of the feed-forward module. In addition, SpecAugment-based data augmentation is utilized for training. ASR output symbols consist of a blank symbol plus 5000 subwords obtained by the SentencePiece method, which we generated from the transcripts of the “clean” 100th LibriSpeech training data subset only. The ASR model is trained for 100 epochs using the Adam optimizer with $\beta_1 = 0.9, \beta_2 = 0.98, \epsilon = 10^{-5}$, and learning rate scheduling similar to with 25000 warmup steps and an initial learning rate factor of 5.0.

A language model (LM) is employed via shallow fusion at inference time, which consists of 2 long short-term memory (LSTM) neural network layers with 1024 units each trained using stochastic gradient descent and the official LM training text data of LibriSpeech, where we excluded sentences that occur in the 860h training data subsets. ASR decoding is based on a time-synchronous
Table 1. Oracle label error rates (LER) [%] for the 860 hours “unlabeled” training data subsets “clean” and “other” using different pseudo-label representations. CN$^{200}$ denotes a confusion network generated from the 20-best ASR hypotheses for each utterance, where low and high indicate low and high pruning settings.

| data set | 1-best | 10-best | 20-best | CN$^{200}$ | CN$^{\text{low}}$ | CN$^{\text{high}}$ |
|----------|--------|---------|---------|-----------|----------------|-----------------|
| clean 360h | 11.3   | 9.0     | 8.7     | 8.3       | 8.6           | 8.9             |
| other 500h | 18.3   | 15.7    | 15.3    | 14.5      | 14.8          | 15.3            |

prefix beam search algorithm similar to \cite{39}. We use a decoding beam size of 30, a score-based pruning threshold of 14.0, an LM-weight of 0.8, and an insertion bonus factor of 2.0.

4.3. Graph Analysis

A seed ASR model is trained using the 100h “clean” training data set of LibriSpeech to generate decoding results for the remaining 860h LibriSpeech training data and to obtain an N-best list of ASR hypotheses used as pseudo-labels for each utterance. Each N-best list of pseudo-labels is used to generate a CTC-style confusion network (CN) as discussed in Section 3 with different settings for pruning. We compare “no”, “low”, and “high” pruning settings in our experiments, where we use a scaling factor of $\mu = 0.6$ and a threshold of $\eta = 0.02$ for “low” and $\eta = 0.05$ for “high” pruning. The pruning settings are determined based on experiments using the development test data sets of LibriSpeech, where selected parameters resulted in a good trade-off between the oracle label error rates (LERs) and the graph densities, which ideally should both be small. Pruning reduces the size and density of a graph, as can be identified by the ratio of the number of non-blank nodes in the graph to the number of labels in a reference sequence, the ground-truth transcription:

- The average graph densities for “no”, “low”, and “high” pruning are 1.510, 1.233, and 1.119 for the “clean” 360h training data set, and 1.545, 1.275, and 1.147 for the “other” 500h training data set. Table 1 shows the oracle LERs for N-best lists of different sizes N as well as for CNs that are generated from the 20-best ASR hypotheses for each utterance. Oracle LERs of an N-best list are obtained by selecting the best pseudo-label sequence from that list, i.e., the sequence that has the smallest edit distance compared to the ground-truth transcription. Oracle LERs of a graph correspond to the minimum edit distance between an FST and a reference sequence, the ground-truth transcription, which is computed by a composition operation between an acoustic FST and an edit-distance FST, followed by a single-source shortest-path algorithm \cite{32}. We can see from Table 1 that an N-best list contains ASR hypotheses with much lower error rates compared to the 1-best hypothesis. For example, selecting the oracle hypothesis from the 20-best list reduces the average LER by 2.5% (clean 360h) and 3.0% (other 500h) on an absolute scale. Using an oracle pseudo-label computed from an N-best list in a graph format reduces the LER even further, since a graph representation of an N-best list allows for more flexible label combinations, as illustrated in Fig. 1.

| N/A | pseudo-labels | W pruning | clean | other | test |
|-----|---------------|-----------|-------|-------|------|
| 1-best | CN$^{200}$ | 1.0 | no     | 5.9    | 15.2  | 6.1 | 15.8 |
| CN$^{20}$ | p | low      | 5.7    | 14.7   | 5.9  | 15.1 |
| CN$^{20}$ | p | high     | 5.6    | 14.2   | 5.9  | 15.0 |
| oracle 20-best |                 |     | 5.2    | 13.9   | 5.5  | 14.5 |
| ground-truth |                 |     | 3.2    | 8.3    | 3.5  | 8.4  |

4.4. ASR Results

GTC-based results for different pseudo-label representations are shown in Table 2. “N/A” denotes results of the seed ASR model, which is trained using the 100h of labeled clean LibriSpeech training data only. CN$^{200}$ indicates the use of CTC-style confusion networks that are generated from the 20-best ASR hypotheses obtained from the seed model for each utterance of the 860h of unlabeled training data. Three different CN setups are compared: a CN without pruning and with all transition weights set to 1.0, a CN with probabilistic transition weights and the low pruning settings of Section 4.3, as well as a CN with probabilistic transition weights and high pruning. In addition, ASR results obtained when training on the best pseudo-label sequence manually selected from the 20-best list are shown as “oracle 20-best”, and results for supervised ASR using the ground-truth transcriptions as “ground-truth”. Table 2 shows that 1-best pseudo-labeling improves the word error rates (WERs) of the LibriSpeech test data sets by a large margin, e.g., from 21.2% to 15.8% for test-other. Training on multiple pseudo-label sequences encoded in graph format further improves the WERs, whereby the best results are obtained when using a CN with high pruning settings and probabilistic transitions weights. Although an unpruned graph is more likely to contain the correct transcription, we suppose the large variance in such a graph makes it harder to learn the best label sequence due to more label noise. Therefore, pruning and the use of transition weights can guide self-training to find pseudo-label sequences of lower error rates. In the best setting, the proposed GTC-based semi-supervised learning approach achieves 1.0% and 0.8% lower WERs for the dev-other and test-other test sets of LibriSpeech, only 0.3% and 0.5% higher compared to the “oracle 20-best” ASR results, which correspond to a lower-bound of the WERs for training on multiple pseudo-labels obtained from an N-best list.

5. CONCLUSIONS

We proposed a new objective function that generalizes the popular CTC loss function to accept weighted finite automata in order to model the training targets using user-defined transition rules and transition weights. The proposed graph-based temporal classification (GTC) loss is applied to a semi-supervised ASR problem in order to leverage a confusion network with a CTC-like structure generated from an N-best list of pseudo-labels for self-training. We demonstrate that GTC-based self-training improves ASR results compared to 1-best pseudo-labeling. The use of pruned graphs and probabilistic transition weights further helps GTC to learn better pseudo-labels from such a graph with improved ASR results. In our experiments, GTC-based self-training has achieved up to 1.0% better WERs compared to the commonly used 1-best pseudo-labeling approach with results close to an oracle experiment, where the best pseudo-label sequence was selected manually from the 20-best list of ASR hypotheses.
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