Parallelizing Linear Recurrent Neural Nets Over Sequence Length

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Abstract

Recurrent neural networks (RNNs) are widely used to model sequential data but their non-linear dependencies between sequence elements prevent parallelizing training over sequence length. We show the training of RNNs with only linear sequential dependencies can be parallelized over the sequence length using the parallel scan algorithm, leading to rapid training on long sequences with small minibatch size. We develop a parallel linear recurrence CUDA kernel and show that it can be applied to immediately speed up several RNN architectures by up to a factor of 9. We abstract recent work on linear RNNs into a new framework of linear surrogate RNNs and develop a linear surrogate for the long short-term memory unit, the GILR-LSTM, which is able to achieve throughput 30 times greater than the currently fastest-available LSTM implementation. We show that the GILR-LSTM is able to perform well on real-world problems,

Thought: we could rephrase this as allowing us to break the necessity of using large batches to train fast

1 Introduction

Recurrent neural networks (RNNs) are widely used for sequence modelling tasks in domains such as natural language processing [18], speech recognition [2], and reinforcement learning [10]. Most RNNs, including popular variants such as long short-term memories (LSTMs) [11] and gated recurrent units (GRUs) [6], contain a non-linear dependency between sequential inputs. These non-linear dependencies create a very flexible class of models but limit the feasibility of training RNNs on long sequences as each sequence element must be processed sequentially. Modelling sequences of thousands to millions of elements is important to domains such as robotics, remote sensing, control systems, speech recognition, medicine, and finance.

The RNN serial evaluation inefficiency problem is usually mitigated by parallelizing the forward and backward pass over a minibatch of inputs. Without minibatches, RNN evaluation is a sequence of matrix-vector multiplications. Minibatches transform RNN computation into a sequence of more efficient matrix-matrix multiplications, but this speed-up brings several disadvantages. RNN model size is often limited by GPU memory size, and running a forward and backward pass on a minibatch requires memory linear in the minibatch size. Grouping data into minibatches increases the latency of each pass and reduces the rate of optimization steps. Finally, training with larger minibatches damages generalization ability [13]. Given these effects, it is desirable to obtain high training throughput with small minibatches. Persistent RNNs [7] use a novel implementation that can achieve high GPU utilization with very small minibatch sizes when the recurrent state is larger than 500 elements, but even persistent RNNs become limited by the serial evaluation inefficiency at smaller hidden sizes.

Numerous prior works have shown strong performance from neural sequential models with only linear dependence on earlier sequence elements. Balduzzi and Ghifary [3] investigated RNNs with only elementwise linear recurrence relations $h_t = \alpha_t \odot h_{t-1} + (1-\alpha_t) \odot x_t$ and developed linear variants of LSTM and GRU that perform similarly to standard non-linear RNNs on text generation tasks. Bradbury et al. [5], Kalchbrenner et al. [12], Gehring et al. [8], and van den Oord et al. [19] have successfully applied networks of convolutions over sequences for tasks such as machine translation, language modelling, and audio generation. These works have observed up to an order of magnitude

increase in training throughput compared to RNN alternatives. Convolutional sequence models typically rely on either an attention mechanism or a (possibly linear) recurrent layer to integrate information at scales larger than the filter width. Introduction of a recurrent layer prevents full parallelization over the sequence length while attention mechanisms are expensive to apply on long sequences in online inference use cases.

A linear recurrence is a specific instance of a general form of computation known as a scan. Scans and reductions are computations involving repeated application of a binary operator \oplus over an array of data. Computing the sum or maximum of an array is an example of a reduction, while a cumulative sum is a common example of a scan operation. Throughout this work, the scan of \oplus with initial value b is defined as

$$SCAN(\oplus, [a_1, a_2, ..., a_n], b) = [(a_1 \oplus b), (a_2 \oplus a_1 \oplus b), ..., (a_n \oplus a_{n-1} ... \oplus a_1 \oplus b)]$$

The reduction of \oplus over array A and initial value b is denoted REDUCE(\oplus , A, b) and is the final element of SCAN(\oplus , A, b). Despite their dependent computation graph, algorithms exist to parallelize scans and reductions when \oplus is associative [15].

Blelloch [4] shows that first order recurrences of the form $h_t = (\Lambda_t \otimes h_{t-1}) \oplus x_t$ can be parallelized with the parallel scan algorithm if three conditions are met:

- 1. \oplus is associative: $(a \oplus b) \oplus c = a \oplus (b \oplus c)$
- 2. \otimes is semiassociative: there exists a binary associative operator \odot such that $a\otimes(b\otimes c)=(a\odot b)\otimes c$
- 3. \otimes distributes over \oplus : $a \otimes (b \oplus c) = (a \otimes b) \oplus (a \otimes c)$

Considering the familiar operations in linear algebra, we see that the associative operation of vector addition $(x \oplus y = x + y)$, the semiassociative operation of matrix-vector multiplication $(A \otimes x = Ax)$ and the associative operation of matrix-matrix multiplication $(A \odot B = AB)$ satisfy Blelloch's three conditions, allowing $h_t = \Lambda_t h_{t-1} + x_t$ to be evaluated in parallel over time steps t for vectors x_t and square matrices Λ_t .

We investigate this idea further and deliver the following contributions:

- We classify RNNs which satisfy the conditions above, and show that many RNNs used in practice such as Quasi-RNNs (QRNNs) [5] are contained in this class
- We provide an implementation of the parallel linear recurrence algorithm as a CUDA kernel, and show that it speeds up training of QRNN and Simple Recurrent Unit (SRU) [20] architectures by factors of up to 9x
- We describe how several recent linear RNNs can be described as linear surrogates for non-linear architectures. We introduce a linear surrogate for the LSTM and show that we are able to train it with a speedup of 5-10x compared to the CuDNN-LSTM when we use the parallel linear recurrence algorithm.

2 Parallel linear recurrence

As the method is essential to this work, Algorithm 1 presents the parallel linear recurrence algorithm for the interested reader.

2.1 Theoretical performance

The cost of a serial scan over a sequence of length T is $C_{\text{sscan}} \in \mathcal{O}((C_{\otimes} + C_{\oplus})T)$, compared to the parallel scan cost $C_{\text{pscan}} \in \mathcal{O}(2(C_{\odot} + C_{\otimes} + C_{\oplus})(T/p + \lg p))$ [4] on p processors. If h_t is a vector of dimension n then $C_{\odot} \in \mathcal{O}(n^3), C_{\otimes} \in \mathcal{O}(n^2), C_{\oplus} \in \mathcal{O}(n)$ giving $C_{\text{pscan}} \in \mathcal{O}(2(n^3 + n^2 + n)(T/p + \lg p))$ and $C_{\text{sscan}} \in \mathcal{O}((n^2 + n)T)$. The $\mathcal{O}(n^3)$ cost of the matrix multiplication in the parallel algorithm can counter-act any parallel speedups for sufficiently large hidden states and lead to a slower algorithm overall.

To avoid this problem, we will only consider diagonal matrices Λ_t , in which case both matrix-matrix and matrix-vector multiplication have cost proportional to n and $C_{\text{pscan}} \in \mathcal{O}(6n(T/p + \lg p))$

Algorithm 1 Parallel linear recurrence on p processors

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1: Let y = [(\Lambda_1, x_1), (\Lambda_2, x_2), ..., (\Lambda_T, x_T)]

2: Let binary operator • act as (\Lambda, x) • h = \Lambda h + x

3: Let S_0 = 1, S_i < E_i, E_i + 1 = S_{i+1}, E_{p-1} = T for i in 0, p-1

4:

5: parfor i \leftarrow 0, p-1 do

6: P_i = \text{REDUCE}(\odot, \Lambda_{S_i:E_i}, I)

7: R_i = \text{REDUCE}(\bullet, y_{S_i:E_i}, 0)

8: end parfor

9:

10: Let z = [(P_0, R_0), (P_1, R_1), ..., (P_p, R_p)].

11: C = \text{SCAN}(\bullet, z, h_0) \triangleright compute C_i = P_i C_{i-1} + R_i with C_{-1} = h_0

12:

13: parfor i \leftarrow 0, p-1 do

14: h_{S_i:E_i} = \text{SCAN}(\bullet, y_{S_i:E_i}, C_{i-1})

15: end parfor

16: return h
```

and $C_{\text{sscan}} \in \mathcal{O}(2nT)$. This gives a parallel speedup factor of $\frac{pT}{3(T+\lg p)}$. Assuming $p \ll T$, then $C_{\text{pscan}} \leq C_{\text{sscan}}$ when $p \geq 3$.

As we are only considering diagonal matrices, we write the linear recurrence as $h_t = \lambda_t \odot h_{t-1} + x_t$ where \odot indicates elementwise multiplication.

Limiting Λ_t to be diagonal may seem like a severe constraint but there are several reasons to do so beyond the unfavorable parallelization performance. Relatively few neural network models use separate recurrent matrices for each sequence element and using these separate matrices would require potentially prohibitive n^2T memory. Applying the same matrix Λ to each sequence element is also unappealing considering that a matrix multiplication can be thought of as a rotation and a scaling. The same rotation at every element seems unlikely to be useful, and the scaling is exactly what's captured in diagonal vectors λ_t . Recurrent coefficient vectors λ_t provide enough flexibility to implement schemes such as exponential moving averages or a gating mechanism.

2.2 Backpropagation

$$\begin{split} \nabla_{h_T} L &= \frac{\partial L}{\partial h_T} \\ \nabla_{h_t} L &= \frac{\partial h_{t+1}}{\partial h_t} \odot \nabla_{h_{t+1}} L + \frac{\partial L}{\partial h_t} \\ &= \lambda_{t+1} \odot \nabla_{h_{t+1}} L + \frac{\partial L}{\partial h_t} \\ \nabla_{\lambda_t} L &= \frac{\partial h_t}{\partial \lambda_t} \odot \nabla_{h_t} L = h_{t-1} \odot \nabla_{h_t} L \\ \nabla_{x_t} L &= \nabla_{h_t} L \\ \nabla_{h_0} L &= \frac{\partial h_1}{\partial h_0} \odot \nabla_{h_1} L = \lambda_1 \odot \nabla_{h_1} L \end{split}$$

The backpropagation equations center around a linear recurrence over $\frac{\partial L}{\partial h_t}$ in the reverse order of the original sequence. This allows for parallelizing both the forwards and backwards pass of a linear RNN over the sequence length.

2.3 Implementation

GPUs commonly used for deep learning in 2017 consist of between 640 and 3200 parallel processors known as warps. Each warp operates on 32 single precision floating point numbers in parallel.

This work implemented parallel linear recurrence as a CUDA kernel with bindings into the TensorFlow [1] framework. Each warp acts as a processor, which means the algorithmic p is up to 3200 and the theoretical parallelization speedup factor is up to several hundred. The 32 lanes of each warp work on different elements of the recurrence vector in parallel. These implementation details mean that peak performance is only obtained on sequences of at least several thousand steps on at least a 32 element vector.

3 Models

Parallel linear recurrence can be used to construct a wide variety of differentiable modules that can be evaluated in parallel. Common applications of linear recurrence include gating schemes and exponential moving averages. Although linear recurrence values can depend only linearly on previous elements, the stacking of linear recurrent layers separated by non-linearities allows for a non-linear dependence on the past. In this sense the non-linear depth of a linear recurrent network is the number of layers and not the sequence length.

3.1 Gated impulse linear recurrent layer

A gated impulse linear recurrent (GILR) layer transforms its m dimensional inputs x_t into a sequence of n dimensional hidden states h_t :

$$g_t = \sigma(Ux_t + b_g)$$

$$i_t = \tau(Vx_t + b_z)$$

$$h_t = g_t \odot h_{t-1} + (1 - g_t) \odot i_t$$

A GILR layer applies the same non-linear transform to each sequence element and then accumulates the sequence elements with a non-linear gating mechanism. Gate g_t uses the sigmoid activation function to give values in [0,1] for reasonable gating semantics, while impulse i_t can use any activation function τ . Stacking GILR layers allows for rich non-linear dependence on previous events while still taking advantage of fast parallel sequence evaluation.

3.1.1 Impact on effective "batch size"

Consider evaluating an RNN with recurrence $h_t = \sigma(Uh_{t-1} + Vx_t + b)$ from m inputs to n hidden units on a sequence of length T with minibatch size b using a serial evaluation strategy. At each of T iterations, the naive approach performs two $(b,m) \times (m,n)$ matrix multiplications. Larger matrix multiplications achieve higher throughput due to less IO overhead, so the better approach computes Vx_t for all t ahead of time in a single $(bT,m) \times (m,n)$ matrix multiply. The non-linear recurrence forces even the better approach to perform T potentially small $(b,m) \times (m,n)$ matrix multiplications in serial. This makes serial RNN performance heavily dependent on minibatch size.

Now consider the GILR, noting that it has the same two matrix-vector multiplications per iteration as the above RNN. The intermediate variables g and i can be evaluated for all t with a single $(bT,m)\times(m,n)$ matrix multiplication each. Given g and i, h can be computed using a parallel linear recurrence over T vectors each of bn elements. Rather than T small operations, the GILR can be evaluated over all sequence elements with two large matrix multiplications and a parallel linear recurrence. GILR performance is much less dependent on batch size as the matrix multiplication kernel sees an "effective batch size" of bT and T is typically large.

3.2 Linear surrogate RNNs

RNNs learn a transition function $s_t = f(s_{t-1}, x_t)$ which combines previous state s_{t-1} with input x_t to compute current state s_t . Non-linear f prevents application of the parallel linear recurrence algorithm and forces slow serial evaluation. To work around this inefficiency, note that s_t serves dual purposes. In $s_t = f(s_{t-1}, x_t)$, s_{t-1} serves as an input to f summarizing the previous inputs while s_t serves as the output of f to be passed to other layers of the network. We can decouple these uses and introduce independent variables for each purpose: s_t is passed onto other layers of the network and we introduce the linear surrogate \tilde{s}_t which is passed onto the next state, with $s_t = f(\tilde{s}_{t-1}, x_t)$. We are still able to choose a nonlinear f, our only limitation being that \tilde{s}_t must be linearly computable.

We refer to this class of model as a linear surrogate RNN (LS-RNN). QRNNs [5] are LS-RNNs using $\tilde{h}_{t-1} = W_k x_{t-k} + ... W_1 x_{t-1}$ and strongly typed RNNs[3] are LS-RNNs with $\tilde{h}_t = x_{t-1}$. Although not a rule, LS-RNNs can often be parallelized over sequence length with either convolution or linear recurrence.

Consider an LSTM:

$$f_t, i_t, o_t = \sigma(U_{f,i,o}h_{t-1} + V_{f,i,o}x_t + b_{f,i,o})$$

$$z_t = \tau(U_z h_{t-1} + V_z x_t + b_z)$$

$$c_t = f_t \odot c_{t-1} + i_t \odot z_t$$

$$h_t = o_t \odot c_t$$

An LSTM has state $s_t = (h_t, c_t)$. Since c_t depends only linearly on c_{t-1} , no surrogate is needed for c_t . h_t has a non-linear dependence on h_{t-1} , so h_t needs a linear surrogate. Introducing a GILR layer as the surrogate, we obtain the GILR-LSTM:

$$\begin{split} g_t &= \sigma(V_g x_t + b_g) \\ j_t &= \tau(V_j x_t + b_j) \\ \tilde{h}_t &= g_t \odot \tilde{h}_{t-1} + (1 - g_t) \odot j_t \\ f_t, i_t, o_t &= \sigma(U_{f,i,o} \tilde{h}_{t-1} + V_{f,i,o} x_t + b_{f,i,o}) \\ z_t &= \tau(U_z \tilde{h}_{t-1} + V_z x_t + b_z) \\ c_t &= f_t \odot c_{t-1} + i_t \odot z_t \\ h_t &= o_t \odot c_t \end{split}$$

For m inputs and hidden size n, a GILR-LSTM contains 2n(n+m) more parameters than the equivalently sized LSTM to handle the mapping from x to \tilde{h} . More generally, a LS-RNN contains all of the same parameters as the underlying RNN as well as some additional parameters to compute the linear surrogate.

4 Experiments

We perform several experiments, first confirming that our implementation of the parallel linear recurrence algorithm is able to achieve higher throughput than a similarly implemented serial version of the algorithm. We show that the parallel kernel is up to 40 times faster than the serial equivalent for long sequence lengths and that this speedup translates to implementations of LS-RNNs such as QRNN (where we can get a speedup of up to 9x).

In order to illustrate that the linearization does not necessarily come at the cost of expressibility, we show that the GILR-LSTM architecture, computed with the parallel linear recurrence algorithm, is able to outperform the optimized CuDNN LSTM implementation on a pathological example from the original LSTM paper [11]. Finally we show that our GILR-LSTM is able to quickly achieve state of the art results on an existing medical data set.

I hope so...

4.1 Throughput Benchmarks

We first illustrate the throughput advantage of the parallel scan algorithm for evaluating the linear recurrence. For a minibatch comprised of b sequences of length T, we define the number of events as bT and the throughput as the number of events per second. We implement two CUDA kernels, one which evaluates the parallel linear recurrence described in algorithm 2, and one which evaluates the linear recurrence using the standard serial approach (with parallelization over features but not sequence length). This comparison is performed directly at the kernel level, avoiding any overhead from TensorFlow. We find that the parallel kernel has a distinct advantage at long sequence lengths with a speedup factor of up to 40x, as shown in table 1. At short sequence lengths, the parallel kernel does not perform well due to the overhead of multiple passes over data and communication between processors.

Table 1: Speedup of the parallel kernel compared to the serial kernel at various values of (minibatch size \mathbf{x} features) = m. Performance was directly measured without any overhead from TensorFlow.

Sequence Length	m=4	m = 32	m = 128
16	0.06	0.06	0.05
256	0.22	0.22	0.86
4,096	1.02	2.94	3.36
65,536	38.5	41.8	17.5

Sequence Length	GILR-LSTM	SRU	QRNN(2)	QRNN(10)
16	0.61	0.28	0.38	0.78
256	0.91	0.84	0.86	0.99
4,096	0.98	1.38	1.18	1.05
65,536	1.41	9.21	6.68	2.05

Figure 1: Speedup obtained when using parallel linear recurrence compared to serial linear recurrence. We analyse this for several recent architectures: the Simple Recurrent Unit from [20], the Quasi-RNN from [5], with a convolutional filter of size 2 and 10, and the GILR-LSTM which we introduce in section 3.2. To give a fair model of a typical use-case maximising the memory usage of the GPU, we set batch size b such that Tb=65536 for a sequence length T.

Many recently introduced RNNs can be accelerated with the parallel linear recurrence algorithm. Table 4.1 shows the throughput advantage by using a parallel linear recurrence when compared to a serial linear recurrence. Since PLR and SLR are simply different methods for computing the recurrence, there is no penalty incurred for this speedup - we obtain exactly the same output.

Typically experiments will aim to maximize the memory utilisation of the GPU, so we simulate this by choosing the product of the batch size and the sequence length to be constant. We see that the architectures which have fewer terms (for which the linear recurrence is a higher proportion of the computational burden) are more effected by the switch to PLR. This is particularly clear in the case of the QRNN, where including a wider convolutional window results in less time spent on the recurrence and less to gain from the parallelization.

Finally, we compare the throughput of the linear surrogate LSTM to the current fastest available implementation of the LSTM, the CuDNN-LSTM. Figure 2 shows that our GILR-LSTM has higher throughput for most of the low-batch-size regime, with a 30x speedup at a sequence length of 65,536. Recent work [13] has suggested that small mini-batches may aid generalization by encouraging the discovery of shallow minima. The GILR-LSTM allows us to achieve high throughput for all choices of minibatch size and sequence length.

Unlike the previous case, these speedups do come at a cost. The GILR-LSTM architecture is different to the canonical LSTM, and it's possible that the loss of model capacity outweighs the speedup in training. The following experiment shows that the GILR-LSTM is able to outperform the LSTM on an example originally designed to illustrate the advantages of the LSTM.

4.2 Synthetic Example

In order to demonstrate how PLR may be used to speed up tasks that the LSTM is well-suited for, we tackle a pathological inference problem. Originally introduced as example 2b from Hochreiter and Schmidhtbuer's introduction of the LSTM, [11], it involves storing a number for a very long sequence of time steps. We have an alphabet of size p with each character represented as a particular one-hot vector in this space. We input a sequence of p of these vectors, chosen at random (with replacement). The first vector in the sequence is always the same, up to the sign of the component (i.e. it is $\pm p_0$.

The two signs on the first component separate the sequences into two sets. The whole sequence is fed into the LSTM and we aim to learn to classify the sequences. This requires remembering the first element over the length of the sequence. Due to the large throughput of the GILR-LSTM, we would expect that it would do better when the sequence length is large. In the original formulation of the

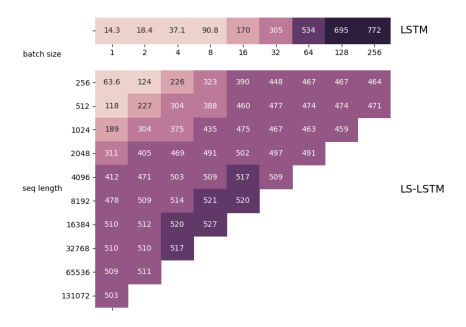


Table 2: Throughput comparison between LSTM-256-256 with GILR-LSTM-256-256, with a 32-dimensional input and a 2-dimensional output, for various batch sizes and sequence lengths. The LSTM only has a single row of data because its throughput is independent of sequence length. Entries are missing from the GILR-LSTM table because there was not enough memory on the GPU to handle such large batch sizes and sequences.

problem (dealing in the regime with around one hundred timesteps), p is set equal to n. Since this would make the size of the input data grow impractically large as $\mathcal{O}(n^2)$ for long sequences, we fix p=128 and vary n.

We generated three sets of data: for n equal to 1,024, 8,192, and 1,048,576. For each of these we compared a two-layer GILR-LSTM with 512 hidden units to a canonical LSTM network with roughly the same number of parameters. We could imagine using a two-layer LSTM with 512 hidden units in each layer, or a one-layer LSTM with 1024 hidden units.

Initial experiments showed that the two-layer LSTM was much quicker to converge than the one-layer LSTM, so here we compare the GILR-LSTM to the two-layer CudnnLSTM. We ran all experiments on a NVIDIA K80 GPU, choosing the largest minibatch size which fit into the GPU memory, with five runs per configuration allowing us to find the average and standard deviation of the time and number of iterations to convergence. For all CUDA runs, a brief search over learning rate and batch size was carried out to find the parameters which allow the network to converge most rapidly. The criterion for convergence was five consecutive minibatches giving 100% accuracy. As can be seen from the learning curves in figure 2, this was a reasonable criterion. For the longest sequence-length, we didn't observe the CuDNN-LSTM converging.

The results illustrate that the GILR-LSTM is able to robustly outperform the CuDNN-LSTM, which is currently the fastest LSTM implementation available. This is somewhat surprising, as we might expect that the LSTM would do particularly well at an example that was specifically constructed for it. We can conclude that the nonlinearities in the LSTM's design are not vital for its performance given that the linear GILR-LSTM is able to converge, and that if these nonlinearities are not completely necessary, they entail a large performance cost by diallowing the plr algorithm from being used. Furthermore, the sequence length of over one million is the largest sequence length we have seen seriously discussed. The GILR handles it straightforwardly.

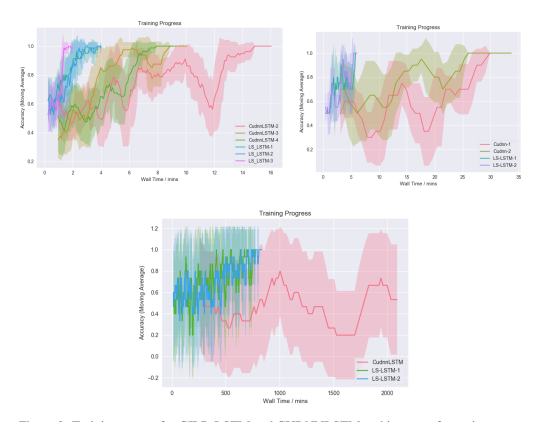


Figure 2: Training curves for GILR-LSTM and CUDNNLSTM architectures for various sequence lengths. (Clockwise from top-left: 1024, 8192, 1048576)

Table 3: Performance of the GILR-LSTM compared to CUDA-optimised CudnnLSTM implementation on problem 2b from (cite '97). We see that the GILR-LSTM has a clear advantage in training speed as the sequence length increases. We did not observe the Cudnn-LSTM converging for the longest task, even after several days. *For the longest sequence length, the number of hidden units was decreased to 64 for both architectures so that the net could fit in memory.

Sequence Length	1,024		8,192		1,048,576*	
	CuDNN	GILR	CuDNN	GILR	CuDNN	GILR
Iterations (1000s)	1.0 ± 0.4	0.55 ± 0.04	0.44 ± 0.05	0.56 ± 0.16	-	14 ± 3
Wall Clock time (hours)	0.28 ± 0.08	0.031 ± 0.002	0.58 ± 0.06	0.10 ± 0.03	-	9.7 ± 1.7

4.3 The Medical Setting

A common application of RNNs is in medicine. In order to illustrate the increased performance that the plr algorithm allows, we implement an GILR-LSTM on the problem presented in , the 2016 PhysioNet/Computing in Cardiology Challenge 2016. The aim of the Classification of Normal/Abnormal Heart Sound Recordings challenge is to determine which of a set of EEGs are produced by people with abnormal hearts. The data is recorded from a variety of sources and split into a training and testing set. The testing set is not publicly available, but can be evaluated through a web portal. The top results cover a large range of techniques, but generally rely on featurization of the raw EEG records. We want to see if we are able to leverage our advantage at high sequence-lenghts to train directly on the raw EEG. The EEGs are sampled at 2,000 Hz and range in length from several seconds to more than one hundred seconds, giving a max sequence length of over 200,000. Since the testing server is not equipped with a GPU, we cannot run PLR in the inference. We resample the records to 500 Hz so that we are able to evaluate the test results in a reasonable amount of time, padding all records that are shorter than the longest.

The training set of data is unbalanced, with 75% of the data corresponding to normal hearts. We rebalance the data by oversampling the abnormal data by a factor of three, including each abnormal heart three times. We train a network consisting of three stacked GIRL-LSTM layers, each with 16 hidden units. Although reasonably small, since the input is only one-dimensional 16 hidden units allows us to develop representations. PLR allows us to train the network very rapidly over a long sequence length, so we don't have to downsample and lose information or spend lots of time developing methods to form features. We were able to fit exactly to the training set after a few hours of training, and on the unseen test data we obtained a score of PLACEHOLDER.

5 Conclusion

Parallel linear recurrence is an extremely powerful algorithm and the GILR-LSTM is just one of many possible models that can be built with it. While we were previously forced to use large minibatches in order to obtain reasonable training speed, with PLR in many cases we can train rapidly on small minibatch sizes, freeing us to utilise extremely long sequence lengths and more complex models. Our CUDA kernel is up to 40 times faster than a serial kernel on sequence lengths of 65,536, and this benefit transfers to implemented models, speeding up the SRU [20] by a factor of 9. The impressive results in [20] and other recent work comprehensively show that linearly-recurrent RNNs are, in many respects, equally as powerful as nonlinear LSTMs. Our results on the synthetic example continue this trend, showing that the GILR-LSTM is able to handle sequence length regimes that are longer than any we have seen seriously suggested in the literature. We feel that the framework of describing linear RNNs as linear surrogates for nonlinear RNNs is a fruitful approach, and that this, paired with the PLR algorithm, will substantially advance the ability to train RNNs, and to understand timeseries data.

We intend to expand upon the experiments section and open-source the parallel linear recurrence kernel in the near future.

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If we get good results (doesn't have to be best ever, but can be on par), then keep

If we don't get good results, or don't have results, we will need to kick this section

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