ELIGIBILITY PROPAGATION IN MULTI-LAYER RECURRENT SPIKING NEURAL NETWORKS

WERNER VAN DER VEEN

Faculty of Science and Engineering University of Groningen

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INTRODUCTION

The human brain is one of the most complex systems in the universe. Its approximately 86 billion neurons (Azevedo et al., 2009) and 100–500 trillion synapses (Drachman, 2005) are capable of abstract reasoning, pattern recognition, memorization, and sensory experience—while consuming only about 20 W of power (Drubach, 2000; Sokoloff, 1960).

An early goal of artificial intelligence has been to construct systems that exhibit similar intelligent traits (Turing, 1948). One of the proposed methods was to emulate the network of biological neurons in the human brain using simple units called perceptrons (Rosenblatt, 1958). These perceptrons were inspired by Hebbian learning (Hebb, 1949), which was a new (and later corroborated) theory on biological learning. This proved to be more difficult than initially hoped, partly due to ostensibly insurmountable problems such as the inability of perceptrons to learn linearly inseparable tasks (Minsky and Papert, 1969). As a consequence, excitement waned, funding dried up (Crevier, 1993), and the field began focusing on more practical applications, such as expert systems and symbolic reasoning (Haugeland, 1985). After the popularization in the 1980s of trainable Hopfield networks (Hopfield, 1982) and backpropagation (Rumelhart, G. E. Hinton, and Williams, 1986), which enabled learning linearly inseparable tasks, artificial neural networks (ANNs) and the connectionist approach were embraced with a new appreciation.

These ANNs are networks of small computational units that can be trained to perform specific pattern recognition tasks. Backpropagation has proven to work well in training ANNs with multiple layers, most popularly in the field of deep learning (DL), which has become a dominant field in artificial intelligence. This popularity is partly due to exponentially increasing computing power and data storage capabilities, as well as the rise of the Internet, which has also provided ample training data. Some variations on ANNs have shown to improve learning performance, such as using convolutional (CNN) and recurrent (RNN) neural networks, both of which are, like the perceptron, inspired by the architecture of the human brain (Fukushima and Miyake, 1982; LeCun, Bengio, et al., 1995; Lukoševičius and Jaeger, 2009). These types of networks approach or exceed human level performance in some areas (Schmidhuber, 2015).

ENERGY LIMITS However, DL-based methods are starting to show diminishing returns; training some state-of-the-art models can require so much data and computing power that only a small number of organizations has the resources to train and deploy them. The computational processes of self-driving cars, for example, consume on the order of a thousand watts. One of the current top submissions of the "Labeled Faces in the Wild" face verification task is a deep ANN by Paravision

that was trained using a dataset of 10 million face images of 100 thousand individuals¹. Beside very large datasets, deep ANNs also require a significant amount of power to train. For instance, ResNet (Kaiming He et al., 2016) has been trained for 3 weeks on a 8-GPU server consuming about 1 GWh. A more extreme case is the 11-billion parameter version of Google's T5 model (Raffel et al., 2019), which is estimated to cost more than \$1.3 million per training run (Sharir, Peleg, and Shoham, 2020). This high power consumption precludes computations in mobile low-power or small-scale devices, which now require at least a connection to a cloud computing server.

The energy consumption of DL contrasts strongly with that of the human brain, which can learn patterns using far less energy and data. This is because despite the biologically inspired foundation, deep ANNs are fundamentally different from the brain, which is an inherently time-dependent dynamical system (Sacramento et al., 2018; Woźniak et al., 2020) that relies on biophysical processes, recurrence, and feedback of its physical substrate for computation (Bhalla, 2014; Sterling and Laughlin, 2015). Deep ANNs are implemented on von Neumann architectures (Von Neumann, 1993), i. e., a system with a central processing unit (CPU) and separate memory, which are significantly different from the working model of the brain (Schuman et al., 2017).

One reason for the inefficiency of deep ANNs is that they suffer from the von Neumann bottleneck (Zenke and E. O. Neftci, 2021), which involves a limited throughput between the CPU and memory—a data operation cannot physically co-occur with fetching instructions to process that data because they share the same communication system. Parallelization on GPUs has alleviated this bottleneck to some extent, but the human brain is more efficient as it is embedded in a physical substrate whose neurons operate fully in parallel (A Pastur-Romay et al., 2017) using sparsely occurring postsynaptic potentials (or *spikes*) (Bear, Connors, and Paradiso, 2020), and where no explicit data processing instructions exist. A spike can be represented as a binary value which causes the synapse to increase the membrane potential in the efferent neuron to change by a fixed value (Bear, Connors, and Paradiso, 2020). Connections in ANNs are represented abstractly by large weight matrices, which are all multiplied with neuron activation values at every propagation cycle. In the brain, a synapse spikes sparsely and thereby saves energy while conveniently including an informative temporal component.

A second reason is that backpropagation requires two passes over the ANN: the first to compute the network output given an input, and the second to propagate the output error back into the network to move the weights between neurons in the direction of the negative gradient. Backpropagation in RNNs is often performed by unrolling the network in a feedforward ANN in a process called backpropagation through time (BPTT). The human brain, in contrast, is unlikely to use

¹ See http://vis-www.cs.umass.edu/lfw/results.html#everai. Last accessed January 2021.

backpropagation, BPTT, or gradients of the output error (Lillicrap and Santoro, 2019).

SPIKING NEURAL NETWORKS Spiking neural networks (SNNs) (Gerstner and Kistler, 2002; Maass, 1997) are another step towards biological plausibility of connectionist models. The concept dates back to the 1980s (Hopfield, 1982); nevertheless, they are sometimes considered as the third generation of ANNs (Maass, 1997). SNNs use neurons that do not relay continuous activation values at every propagation cycle, but spike once when they reach a threshold value. SNNs are competitive to ANNs in terms of accuracy and computational power, as well in their ability to display precise spike timings (Lobo et al., 2020). Their sparse firing regimes also offer improved interpretability of their behavior as compared to traditional ANNs (Soltic and Kasabov, 2010), which is desired in areas such as medicine or aviation.

However, SNNs have not been as popular as ANNs. One reason for this is that spike-based activation is not differentiable. As a consequence, backpropagation cannot be directly used to move in the negative direction of the error gradient, although some attempts have been made to bridge this divide (Bellec, Scherr, Hajek, et al., 2019; Bohte, Kok, and La Poutre, 2002; Hong et al., 2010; J. H. Lee, Delbruck, and Pfeiffer, 2016; Ourdighi and Benyettou, 2016; Sacramento et al., 2018; Whittington and Bogacz, 2019; Y. Xu et al., 2013) and to make backpropagation more biologically plausible.

Similarly, it has been demonstrated that approximations of BPTT can be applied to recurrent SNNs (RSNNs) (Bellec, Salaj, et al., 2018; Huh and Sejnowski, 2017). Other RSNN training methods rely on control theory to train a chaotic reservoir of neurons (Gilra and Gerstner, 2017; Thalmeier et al., 2016). Information locality can be preserved in some of these methods (Alemi et al., 2018), i. e., a neuron or synapse can only access information of itself, or the communication of synapses or neurons (resp.) with which it is directly connected. The FORCE training method (Nicola and Clopath, 2017), in contrast, can also train RSNNs but is nonlocal and therefore not suited for biologically plausible implementations. This body of research led to increased understanding of training SNNs and how to obtain better learning performances. For instance, both single- and multi-layer SNNs have shown good performance in visual processing (Escobar et al., 2009; Kheradpisheh et al., 2018; D. Liu and Yue, 2017) and speech recognition (Dong, Xuhui Huang, and B. Xu, 2018; Tavanaei and Maida, 2017). However, none of these algorithms are biologically plausible. While DL was rapidly becoming popular during the 2010s, there was no clear learning algorithm for SNNs that could compete with ANNs. A second reason for the relative unpopularity of SNNs is that they are generally emulated in von Neumann architectures, undermining their energy efficiency advantages.

NEUROMORPHIC COMPUTING SNN learning algorithms are particularly useful in the upcoming field of neuromorphic computing (NC) (Mitra, Fusi, and Indiveri, 2008), in which analog very-large-scale in-

tegration (VLSI) systems are used to implement neural systems. On the surface, it can be understood as running neural networks not abstracted in a digital system, but physically embedded in a dedicated analog medium. A central advantage of NC is energy efficiency (Hasler and Akers, 1990; J.-C. Lee and Sheu, 1990; Tarassenko et al., 1990). This energy efficiency, combined with NC's massive parallelism (Monroe, 2014), makes VLSIs particularly relevant for implementing SNNs.

Like SNNs, neuromorphic systems typically use sparse, event-based communication between memristors (Maan, Jayadevi, and James, 2016) and physically colocalized memory and computation (E. O. Neftci, 2018; Sterling and Laughlin, 2015). Although colocalized memory and computation has also been implemented in von Neumann machines, such as Google's TPU², Graphcore's IPU³, or Cerebras' CS-1⁴, neuromorphic systems are more efficient for running ANNs (Merolla et al., 2014; Rajendran et al., 2019). The energy consumption of CMOS artificial neurons is several orders of magnitude lower than that of neurons in an ANN, and even 2–3 times lower than the energy consumption of biological neurons (Elbez et al., 2020). Neuromorphic systems are also more tolerant to device variation (Yu et al., 2013).

Because of this massive parallelism, high energy efficiency, good error tolerance, and good ability to implement cognitive functions, neuromorphic systems are attracting strong interest. In particular, SNNs emerged as an ideal biologically inspired NC paradigm for realizing energy-efficient on-chip intelligence hardware (Davies et al., 2018; Merolla et al., 2014), suitable for running fast and complex SNNs on low-power devices. For instance, a competitive image classification performance was reached with a 6-order of magnitude speedup in a leaky integrate-and-fire (LIF) SNN in field-programmable gate arrays, compared to digital simulations (G. Zhang et al., 2020).

BIOLOGICAL LEARNING To run an SNN on neuromorphic hardware, a local and online algorithm is needed. The precondition of locality refers to the idea that a neuron or synapse can only access information or communication with which it is physically connected. For instance, the inner state of a neuron can only be influenced by itself, or by the spikes it receives from afferent neurons. Similarly, a synapse can only spike or change its weight based on signals from the afferent and efferent neuron. This is a direct consequence of the colocalization of processing and memory. The precondition of online can be regarded as temporal locality—neurons and synapses can only access information that physically exists at the same point in time. They cannot access future information, nor past information if it was not explicitly retained.

The brain also adheres to these two constraints. Most learning algorithms for SNNs are based on a form of Hebbian learning, which is a major factor in biological learning and memory consolidation. Classical Hebbian learning is often summarized by "cells that fire together, wire

² See https://cloud.google.com/tpu/docs/tpus. Last accessed January 2021

³ See https://www.graphcore.ai/products/ipu. Last accessed January 2021

⁴ See https://cerebras.net/product/#chip. Last accessed January 2021

together", if there is a causal relationship between these cells, such as a postsynaptic potential on a connecting synapse. Direct application of Hebbian learning in a spiking neural network will generally lead to a positive feedback loop, because "wiring cells together", or increasing the synaptic strength, will in turn increase the likelihood that they also fire together (Zenke, Gerstner, and Ganguli, 2017). Furthermore, classical Hebbian learning describes no way for a synapse to weaken.

Spike-timing-dependent plasticity (STDP) (Abbott and Nelson, 2000; Caporale and Dan, 2008) is a type of Hebbian learning that incorporates temporal causality on a synapse from neuron A to neuron B: if B spikes right after neuron A, then the synapse is strengthened, but if B spikes right before A, it is weakened. However, this too leads to runaway activity through the same positive feedback loop. It is widely known that STDP is a fundamental learning principle in the human brain (Caporale and Dan, 2008; Kandel et al., 2000), including perceptual systems in the sensory cortex (S. Huang et al., 2014). STDP by itself can be used as an unsupervised learning algorithm or to forming associations in classical conditioning (Diehl and Cook, 2015; C.-H. Kim et al., 2018). Furthermore, it has been demonstrated to form associations between memory traces in SNNs, which are crucial for cognitive brain function (Pokorny et al., 2020). To allow supervised learning, or operant conditioning, a learning signal is required to influence the direction of the synapse weight change: a positive learning signal will reinforce the association (long-term potentiation), and a negative learning signal weakens it (long term depression) (Lobov et al., 2020). STDP with a learning signal is known as reward-modulated STDP (R-STDP) (Legenstein, Pecevski, and Maass, 2008) in the field of SNNs and three-factor Hebbian learning in neuroscience (Frémaux and Gerstner, 2016), outperforming its classical two-factor counterpart (Porr and Wörgötter, 2007) because learning signals are crucial for maintaining long-term memory (Bailey et al., 2000) and behavioral changes.

Neurotransmitters are used to modulate the learning signal in the brain. Dopamine, for instance, which has a central behavioral and functional role in the primary motor cortex (Barnes et al., 2005; Dang et al., 2006), has been shown to modulate synapses through dendritic spine enlargement during a very narrow time window (Dang et al., 2006). It is behaviorally related to novelty and reward prediction (S. Li et al., 2003; Schultz, 2007) by gating neuroplasticity of corticostriatal (Reynolds, Hyland, and Wickens, 2001; Reynolds and Wickens, 2002) and ventral tegmental (VTA) synapses (Bao, Chan, and Merzenich, 2001). In the VTA, dopaminergic neurons respond to learning signals in a highly localized manner that is specific for local populations of neurons (Engelhard et al., 2019). This is also the case in other areas of the midbrain (Roeper, 2013). Acetylcholine is another example of a neuromodulator that gates synaptic plasticity in the cortex and enables state-dependent learning, in which memories are recalled better if the sensory context is similar to that during the memory encoding (Shulz et al., 2000).

However, R-STDP by itself does not solve the credit assignment problem, which relates to neuromodulation of synapses after a learning signal is presented with some delay. In that case, when the learning signal is presented, the neurons have long spiked, and it is not clear which synapses elicited the behavior that is rewarded or punished. Recent research suggests that the brain uses *eligibility traces* (Florian, 2007; Izhikevich, 2007) to solve the credit assignment problem (Gerstner, Lehmann, et al., 2018; Stolyarova, 2018). In particular, the synaptic CaMKII protein complex is activated during the induction of long-term potentiation (LTP) of biological synapses if the presynaptic neuron spikes shortly before a postsynaptic neuron (Sanhueza and Lisman, 2013). This LTP is maintained over behavioral time spans, and gradually fades. When followed by a learning signal in the form of a neurotransmitter, synaptic plasticity is induced (Cassenaer and Laurent, 2012; Gerstner, Lehmann, et al., 2018; Yagishita et al., 2014).

Synaptic plasticity was demonstrated using eligibility traces in deep feedforward SNNs (Kaiser, Mostafa, and E. Neftci, 2020; E. O. Neftci et al., 2017; Zenke and Ganguli, 2018) and could be implemented in feedforward VLSIs. In Zenke and Ganguli (2018) it is asserted that these methods are also applicable for RSNNs. Eligibility traces have also been shown to solve difficult credit assignment problems in SNNs using R-STDP (Bellec, Scherr, Subramoney, et al., 2020; Legenstein, Pecevski, and Maass, 2008) and in RNNs (Kaiwen He et al., 2015), and have a predictable learning effect (Legenstein, Pecevski, and Mass, 2008). R-STDP in SNNs has been used to solve a number of tasks, including training a stable lane keeping controller (Bing, Meschede, Chen, et al., 2020), but can suffer from catastrophic forgetting and lack of policy evaluation in mapless navigation (Bing, Meschede, K. Huang, et al., 2018). More recently, Demirag et al. (2021) have shown that phase change memory (PCM) devices allow eligibility traces to linger over behavioral timescales in CMOS/memristive architectures.

Eligibility propagation (e-prop) (Bel-ELIGIBILITY PROPAGATION lec, Scherr, Subramoney, et al., 2020) is a local and online learning algorithm for RSNNs that can be mathematically derived as an approximation to BPTT (see also Section 2.1.4). It uses local learning signals and eligibility traces for any type of neuron. In e-prop, the learning signal is a local variation on random broadcast alignment, which propagates the error directly back onto the neurons with a random weight, resembling the function of a neuromodulator in the brain. This has been suggested to provide a diversity of feature detectors for task-relevant network inputs (Bellec, Scherr, Subramoney, et al., 2020). Broadcast alignment can perform as effectively as backpropagation in some tasks in feedforward ANNs (Lillicrap, Cownden, et al., 2016; Nøkland, 2016) and multi-layer SNNs (Clopath et al., 2010; Samadi, Lillicrap, and Tweed, 2017), but performs poorly in deep feedforward ANNs for complex image classification tasks (Bartunov et al., 2018).

The local and online properties of e-prop make it a biologically plausible learning algorithm that can be implemented on VLSIs. E-prop has

been demonstrated to work for a large variety of tasks, including on classifying phones (i. e., speech sounds), for which it performs competitively with LSTMs (Graves, Mohamed, and G. Hinton, 2013), a popular RNN architecture that uses BPTT.

So far, only the LIF and adaptive LIF (ALIF) neuron models have been used in e-prop. In Traub et al. (2020), a functional modification was made to the LIF model such that STDP can occur. In particular, STDP occurs when the neuron model provides a negated gradient signal in the case when a presynaptic signal arrives too late. This resembles the biological phenomenon of error-related negativity (ERN) (Nieuwenhuis et al., 2001), which is a negative brain response that immediately follows an erroneous behavioral response and peaks after 80–150 ms with an amplitude that depends on the intent and motivation of a person. Traub et al. (2020) also showed this effect for the Izhikevich neuron (Izhikevich, 2003). However, these STDP-modified neurons were shown only in a single-synapse demo to illustrate the STDP properties, not in a full learning task.

MULTI-LAYER RSNNS The human neocortex, which is responsible for processing sensory information, controlling motor output, and mediating higher-order cognitive functions, is organized into six recurrent layers of spiking neurons (Greig et al., 2013).

The discovery of backpropagation allowed gradient descent—based training of multi-layer ANNs, which significantly increased their performance. Multi-layer CNNs show higher levels of abstraction in deeper layers of the network. For instance, early convolutional filters identify lines and edges, while deeper filters identify more complex shapes. In RNNs, stacking recurrent layers results in a similar abstraction—but it is temporal instead of spatial (Gallicchio, Micheli, and Pedrelli, 2017; Hermans and Schrauwen, 2013). Deeper RNN layers exhibit slower time dynamics and longer memory spans than shallow layers (Gallicchio, 2018), suggesting that they ignore small variations in the input signal and integrate larger temporal patterns. It is unclear if these findings extrapolate to RSNNs.

THIS RESEARCH In this report, the performance of e-prop in Bellec, Scherr, Subramoney, et al. (2020) is reproduced. Whereas e-prop was implemented using automatic differentiation in that research, in this report it is done explicitly, using the e-prop learning equations. Furthermore, the two new STDP neuron models of Traub et al. (2020) are experimentally verified in the TIMIT phone classification task. One of these models, the STDP-LIF, will be modified to an adaptive version called STDP-ALIF. Finally, the effect of stacking multiple RSNN layers in an e-prop with each of the three neuron models is examined.

Chapter 2 describes the basic version of the e-prop framework. Chapter 3 describes the method used to implement the TIMIT learning task and modify the e-prop algorithm to a multi-layer framework with different neuron models. The results are presented in Chapter 4 and discussed in Chapter 5. Finally, Chapter 6 concludes this report.

2.1 ELIGIBILITY PROPAGATION

2.1.1 Network model

An eligibility propagation model \mathcal{M} is defined by a tuple $\langle M, f \rangle$, where M is a function

$$\mathbf{h}_{j}^{t} = M\left(\mathbf{h}_{j}^{t-1}, \mathbf{z}^{t-1}, \mathbf{x}^{t}, \mathbf{W}_{j}\right)$$
(2.1)

that defines the hidden state \mathbf{h}_{j}^{t} of a neuron j at a discrete time step t, where \mathbf{z}^{t-1} is the observable state of all neurons at the previous time step (i. e., the binary spike values), \mathbf{x}^{t} is the model input vector at time t, and \mathbf{W}_{j} is the weight vector of afferent synapses. The update of the observable state of a neuron j at time t is defined by

$$z_i^t = f\left(\mathbf{h}_i^t\right). \tag{2.2}$$

This formalization means that e-prop is a *local* training method, because a neuron's observable state depends only on its own hidden state, and the hidden state depends only on observable signals that are directly connected to it. E-prop is also an *online* training method, because both the hidden and observable state of a neuron depend only on variables in the previous time point.

2.1.2 Neuron models

LIF NEURON In Bellec, Scherr, Subramoney, et al. (2020), the LIF neuron model is formulated in the context of e-prop, along with a variant (viz. ALIF) that has an adaptive threshold based on the neuron's spiking frequency. The observable state of a LIF model is given by

$$z_j^t = H\left(v_j^t - v_{\text{th}}\right),\tag{2.3}$$

where H is the Heaviside step function, and $v_{\rm th}$ is the threshold constant. (Note that this and all other used constant parameters are listed in Table A.2.) Consequently, the observable state $z_j^t \in \{0,1\}$ is binary, and denotes the spike activity of a neuron. These spikes are the only communication between neurons in the model. The LIF neuron model has a single state h_j^t that contains only an activity value v_j^t and evolves over time according to the equation

$$v_j^{t+1} = \alpha v_j^t + \sum_{i \neq j} W_{ji}^{\text{rec}} z_i^t + \sum_i W_{ji}^{\text{in}} x_i^{t+1} - z_j^t v_{\text{th}}, \tag{2.4}$$

where $W_{ji}^{\rm rec}$ is a synapse weight from neuron i to neuron j, α is a constant decay factor. Whenever j spikes (i.e., $z_j^t = 1$), the activity of the neuron

is reduced to a value near 0 by the term $-z_j^t v_{\rm th}$. Furthermore, z_j^t is fixed to 0 for $T^{\rm refr}$ time steps to model neuronal refractoriness. In biological neurons, the refractory period consists of an "absolute" phase, during which evoking a new spike is impossible, and a subsequent "relative" phase where the threshold is temporarily increased (Purves et al., 2008). Clamping z_j^t to 0 emulates only this absolute phase.

ALIF NEURON The ALIF neuron introduces a threshold adaptation variable a_j^t to the hidden state of the neuron, such that $\mathbf{h}_j^t \stackrel{\mathsf{def}}{=} \left[v_j^t, a_j^t \right]$. In an ALIF neuron, the spiking threshold increases after a spike, and otherwise decreases back to a baseline threshold v_{th} .

This resembles *spike frequency adaptation* (SFA), a common feature of neocortical pyramidal neurons (Benda and Herz, 2003). SFA is a homeostatic control mechanism that affects the spiking frequency based on the recent spiking activity. Ahmed et al. (1998) found that a single time constant is a good fit to characterize the threshold's exponential decay to a steady state.

The observable state of an ALIF neuron is therefore described by

$$z_j^t = H\left(v_j^t - v_{\text{th}} - \beta a_j^t\right) \tag{2.5}$$

and

$$a_j^{t+1} = \rho a_j^t + z_j^t, (2.6)$$

where ρ is an adaptation decay constant. The interaction between the neuron activity, adaptive threshold and spiking behavior is illustrated in Figure 2.1.

In this paper, the LIF neuron is generalized as an ALIF neuron for which $\beta=0$, effectively canceling the effect of the threshold adaptation value a_j^t on the observable state z_j^t in Equation 2.5. Therefore, only the e-prop derivations for the ALIF neurons will be described in the following sections.

2.1.3 Network topology

The neurons receive inputs from the input feature vector and the observable states of the afferent neurons.

Weights are initialized by sampling them from a Gaussian distribution $\mathcal{N}\left(0,\sqrt{N}\right)$ where N is the number of afferent neurons. For instance, the weights \mathbf{W}^{in} between the input and the first layer are sampled from $\mathcal{N}\left(0,\sqrt{39}\right)$, because there are 39 input features.

A randomly selected 80% of synaptic weights is then set to a value of 0, as well as the synapses that connect a neuron to itself, rendering them ineffective.

Figure 2.2 illustrates the basic architecture of a single-layer e-prop model.

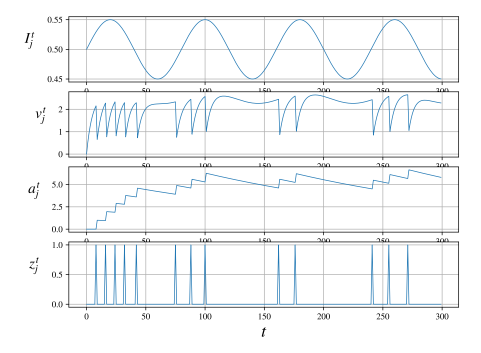


Figure 2.1: A simulated ALIF neuron j receives a sinusoidal input I for 300 time steps t. This figure illustrates the adaptive threshold a, which increases at every spike z, requiring a higher activity v for a next spike. When a spike occurs, v decreases by $v_{\rm th}$.

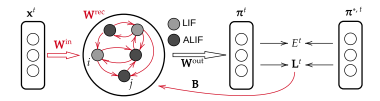


Figure 2.2: An illustration of a single-layer network architecture.

2.1.4 Deriving e-prop from RNNs

Eligibility propagation is a local and online training method that can be derived from backpropagation through time (BPTT). In BPTT, an RNN is unfolded in time, such that the backpropagation method used in feedforward neural networks can be applied to compute the gradients of the cost with respect to the network weights.

In this subsection, the main equation of e-prop

$$\frac{dE}{dW_{ji}} = \sum_{t} \frac{dE}{dz_{j}^{t}} \cdot \left[\frac{dz_{j}^{t}}{dW_{ji}} \right]_{\text{local}}$$
(2.7)

is derived from the classical factorization of the loss gradients in an unfolded RNN:

$$\frac{dE}{dW_{ji}} = \sum_{t'} \frac{dE}{d\mathbf{h}_{j}^{t'}} \cdot \frac{\partial \mathbf{h}_{j}^{t'}}{\partial W_{ji}},\tag{2.8}$$

where summation indicates that weights are shared.

We now decompose the first term into a series of learning signals $L_j^t = \frac{dE}{dz_j^t}$ and local factors $\frac{\partial \mathbf{h}_j^{t-t'}}{\partial \mathbf{h}_j^t}$ for all t after the event horizon t':

$$\frac{dE}{d\mathbf{h}_{j}^{t'}} = \underbrace{\frac{dE}{dz_{j}^{t'}}}_{L_{i}^{t'}} \frac{\partial z_{j}^{t'}}{\partial \mathbf{h}_{j}^{t'}} + \frac{dE}{d\mathbf{h}_{j}^{t'+1}} \frac{\partial \mathbf{h}_{j}^{t'+1}}{\partial \mathbf{h}_{j}^{t'}}.$$
(2.9)

Note that this equation is recursive. If we substitute Equation 2.9 into the classical factorization (Equation 2.8), we obtain a recursive expansion that has $\frac{dE}{d\mathbf{h}_{i}^{T+1}}$ as its terminating case:

$$\frac{dE}{dW_{ji}} = \sum_{t'} \left(L_j^{t'} \frac{\partial z_j^{t'}}{\partial \mathbf{h}_j^{t'}} + \frac{dE}{d\mathbf{h}_j^{t'+1}} \frac{\partial \mathbf{h}_j^{t'+1}}{\partial \mathbf{h}_j^{t'}} \right) \cdot \frac{\partial \mathbf{h}_j^{t'}}{\partial W_{ji}} \tag{2.10}$$

$$= \sum_{t'} \left(L_j^{t'} \frac{\partial z_j^{t'}}{\partial \mathbf{h}_j^{t'}} + \left(L_j^{t'+1} \frac{\partial z_j^{t'+1}}{\partial \mathbf{h}_j^{t'+1}} + (\cdots) \frac{\partial \mathbf{h}_j^{t'+2}}{\partial \mathbf{h}_j^{t'+1}} \right) \frac{\partial \mathbf{h}_j^{t'+1}}{\partial \mathbf{h}_j^{t'}} \right) \cdot \frac{\partial \mathbf{h}_j^{t'}}{\partial W_{ji}}.$$

$$(2.11)$$

We write the term in parentheses into a second term indexed by t:

$$\frac{dE}{dW_{ji}} = \sum_{t'} \sum_{t > t'} L_j^t \frac{\partial z_j^t}{\partial \mathbf{h}_j^t} \frac{\partial \mathbf{h}_j^t}{\partial \mathbf{h}_j^{t-1}} \cdots \frac{\partial \mathbf{h}_j^{t+1}}{\partial \mathbf{h}_j^{t'}} \cdot \frac{\partial \mathbf{h}_j^{t'}}{\partial W_{ji}}.$$
 (2.12)

We then exchange the summation indices to pull out the learning signal L_i^t from the inner summation.

Within the inner summation, the terms $\frac{\partial \mathbf{h}_j^{t+1}}{\partial \mathbf{h}_j^t}$ are collected in an eligibility vector ϵ_{ji}^t and multiplied with the learning signal L_j^t at every time step t. This is crucial for understanding why e-prop is an online training method—local gradients are computed based on traces that are directly accessible at the current time step t, and the eligibility vector operates as a recursively updated "memory" to track previous local hidden state derivatives:

$$\epsilon_{ji}^{t} = \frac{\partial \mathbf{h}_{j}^{t}}{\partial \mathbf{h}_{j}^{t-1}} \cdot \epsilon_{ji}^{t-1} + \frac{\partial \mathbf{h}_{j}^{t}}{\partial W_{ji}}.$$
(2.13)

This is why the ρ and α parameters, which define the decay rate in hidden states and the corresponding eligibility vectors, should be set according to the required working memory in the learning task. The eligibility vector and the hidden state have the same dimension: $\left\{\epsilon_{ji}^t, \mathbf{h}_j^t\right\} \subset \mathbb{R}^d$, where d=2 for all neuron types described in this report.

The *eligibility trace* e_{ji}^t is a product of $L_j^t = \frac{dE}{dz_j^t}$ and the eligibility vector, resulting in the gradient that can be immediately applied at

every time step t, or accumulated and integrated locally on a synapse (see Section 2.1.5.2 for details):

$$\frac{dE}{dW_{ji}} = \sum_{t} \frac{dE}{dz_{j}^{t}} \underbrace{\frac{\partial z_{j}^{t}}{\partial \mathbf{h}_{j}^{t}}}_{\underbrace{t \geq t'}} \underbrace{\frac{\partial \mathbf{h}_{j}^{t}}{\partial \mathbf{h}_{j}^{t-1}} \cdots \frac{\partial \mathbf{h}_{j}^{t+1}}{\partial \mathbf{h}_{j}^{t'}} \cdot \frac{\partial \mathbf{h}_{j}^{t'}}{\partial W_{ji}}}_{e_{ji}^{t}}.$$
(2.14)

This is the main e-prop equation.

2.1.5 Learning procedure

The e-prop equation can be applied to any neuron type with any number of hidden states. In this section, the derivation for ALIF neurons will be detailed.

2.1.5.1 Eligibility trace

Recall the hidden state update equations from Section 2.1.2:

$$v_j^{t+1} = \alpha v_j^t + \sum_{i \neq j} W_{ji}^{\text{rec}} z_i^t + \sum_i W_{ji}^{\text{in}} x_i^{t+1} - z_j^t v_{\text{th}}$$
 (2.4 revisited)

and

$$a_j^{t+1} = \rho a_j^t + z_j^t \tag{2.6 revisited}$$

and the update of the observable state

$$z_j^t = H\left(v_j^t - v_{\text{th}} - \beta a_j^t\right). \tag{2.5 revisited}$$

The ALIF neuron model has a two-dimensional hidden state

$$\mathbf{h}_{j}^{t} = \begin{pmatrix} v_{j}^{t} \\ a_{j}^{t} \end{pmatrix} \tag{2.15}$$

associated with a neuron j and a two-dimensional eligibility vector

$$\epsilon_{ji}^{t} \stackrel{\mathsf{def}}{=} \begin{pmatrix} \epsilon_{ji,v}^{t} \\ \epsilon_{ji,a}^{t} \end{pmatrix} \tag{2.16}$$

associated with a synapse from neuron i to neuron j.

The hidden state derivative $\frac{\mathbf{h}_{j}^{t+1}}{\mathbf{h}_{j}^{t}}$ must be computed to derive the eligibility vector. This hidden state derivative is expressed by a 2×2 matrix of partial hidden state derivatives:

$$\frac{\mathbf{h}_{j}^{t+1}}{\mathbf{h}_{j}^{t}} = \begin{pmatrix} \frac{\partial v_{j}^{t+1}}{\partial v_{j}^{t}} & \frac{\partial v_{j}^{t+1}}{\partial a_{j}^{t}} \\ \frac{\partial a_{j}^{t+1}}{\partial v_{j}^{t}} & \frac{\partial a_{j}^{t+1}}{\partial a_{j}^{t}} \end{pmatrix}. \tag{2.17}$$

The presence of z_j^t , and its relation with the Heaviside step function $H(\cdot)$ in the hidden state updates in Equation 2.4 and Equation 2.6 seems problematic for computing these partial derivatives, because the derivative $\frac{\partial z_j^t}{\partial v_j^t}$ is nonexistent. This is overcome by replacing it with a simple nonlinear function called a pseudo-derivative. Outside of the refractory period of a neuron j, this pseudo-derivative has the form

$$\psi_j^t = \gamma \max\left(0, 1 - \left| \frac{v_j^t - v_{\text{th}} - \beta a_j^t}{v_{\text{th}}} \right| \right)$$
 (2.18)

where γ is a dampening constant, which is set to 0 during the neuron's refractory period.

Now, the partial derivatives in the hidden state derivative can be computed:

$$\frac{\partial v_j^{t+1}}{\partial v_j^t} = \alpha \tag{2.19}$$

$$\frac{\partial v_j^{t+1}}{\partial a_j^t} = 0 (2.20)$$

$$\frac{\partial a_j^{t+1}}{\partial v_j^t} = \psi_j^t \tag{2.21}$$

$$\frac{\partial a_j^{t+1}}{\partial a_j^t} = \rho - \psi_j^t \beta. \tag{2.22}$$

These partial derivatives can be used to compute the eligibility vector:

$$\begin{pmatrix} \epsilon_{ji,v}^{t+1} \\ \epsilon_{ji,a}^{t+1} \end{pmatrix} = \begin{pmatrix} \frac{\partial v_j^{t+1}}{\partial v_j^t} & \frac{\partial v_j^{t+1}}{\partial a_j^t} \\ \frac{\partial a_j^{t+1}}{\partial v_j^t} & \frac{\partial a_j^{t+1}}{\partial a_j^t} \end{pmatrix} \cdot \begin{pmatrix} \epsilon_{ji,v}^t \\ \epsilon_{ji,a}^t \end{pmatrix} + \begin{pmatrix} \frac{\partial v_j^{t+1}}{\partial W_{ji}} \\ \frac{\partial a_j^{t+1}}{\partial W_{ji}} \end{pmatrix}$$
(2.23)

$$= \begin{pmatrix} \alpha & 0 \\ \psi_{j}^{t} & \rho - \psi_{j}^{t} \beta \end{pmatrix} \cdot \begin{pmatrix} \epsilon_{ji,v}^{t} \\ \epsilon_{ji,a}^{t} \end{pmatrix} + \begin{pmatrix} z_{i}^{t-1} \\ 0 \end{pmatrix}$$
 (2.24)

$$= \begin{pmatrix} \alpha \cdot \epsilon_{ji,v}^t + z_i^{t-1} \\ \psi_j^t \epsilon_{ji,v}^t + \left(\rho - \psi_j^t \beta\right) \epsilon_{ji,a}^t \end{pmatrix}. \tag{2.25}$$

This eligibility vector can be recursively applied. For eligibility vectors of synapses that are efferent to input neurons, the input value x_i^t is used in place of z_i^{t-1} in Equation 2.24. Note that the current time index t is used for input neurons to satisfy the online learning principle defined in the model definition in Equation 2.1; neurons receive input from the input at time t, and from the spikes of other neurons sent at time t-1. Furthermore, the absence of $\epsilon_{ji,a}^t$ in the computation of $\epsilon_{ji,v}^{t+1}$ facilitates online training in emulations in non–von Neumann machines, because $\epsilon_{ji,a}^{t+1}$ can be computed before $\epsilon_{ji,v}^{t+1}$, relieving the need to store a temporary copy of its value. In later sections, it is demonstrated that this does not necessarily hold for other neuron models, such as the Izhikevich neuron.

Multiplying the eligibility vector with the partial derivative of the observable state with respect to the hidden state results in the eligibility trace:

$$e_{ji}^{t} = \begin{pmatrix} \epsilon_{ji,v}^{t} \\ \epsilon_{ji,a}^{t} \end{pmatrix} \cdot \begin{pmatrix} \frac{\partial z_{j}^{t}}{\partial v_{j}^{t}} \\ \frac{\partial z_{j}^{t}}{\partial a_{i}^{t}} \end{pmatrix}$$

$$(2.26)$$

$$= \begin{pmatrix} \epsilon_{ji,v}^t \\ \epsilon_{ji,a}^t \end{pmatrix} \cdot \begin{pmatrix} \psi_j^t \\ -\beta \psi_i^t \end{pmatrix} \tag{2.27}$$

$$= \psi_j^t \left(\epsilon_{ji,v}^t - \beta \epsilon_{ji,a}^t \right). \tag{2.28}$$

2.1.5.2 Gradients

Gradient descent is used to apply the weight updates, such that weights are updated by a small fraction η in the negative direction of the estimated gradient of the loss function with respect to the model weights:

$$\Delta W = -\eta \frac{\widehat{dE}}{dW_{ji}} \stackrel{\text{def}}{=} -\eta \sum_{t} \frac{\partial E}{\partial z_{j}^{t}} e_{ji}^{t}. \tag{2.29}$$

Note that for clarity, this section describes e-prop using the stochastic gradient descent. In the actual implementation, the Adam optimization algorithm (Kingma and Ba, 2014) is used (see Section 3.4).

ERROR METRIC In the TIMIT frame-wise phone classification task, there are K=61 output neurons y_k^t where $k\in[1\ldots K].$ These are computed according to

$$y_k^t = \kappa y_k^{t-1} + \sum_j W_{kj}^{\text{out}} z_j^t + b_k,$$
 (2.30)

where $\kappa \in [0,1]$ is the decay factor for the output neurons, W_{kj}^{out} is the weight between neuron j and output neuron k, and b_k is the bias value. The decay factor κ acts as a low-pass filter, smoothening the output values over time and implemented based on the observation that output frame classes typically persist for multiple time steps.

The softmax function $\sigma(\cdot)$ computes the predicted probability π_k^t for class k at time t:

$$\pi_k^t = \sigma_k (y_1^t, \dots, y_K^t) = \frac{\exp(y_k^t)}{\sum_{k'} \exp(y_{k'}^t)}.$$
 (2.31)

This predicted probability is compared to the one-hot probability vector corresponding to the target class label $\pi_k^{*,t}$ at time t using the cross entropy loss function

$$E = -\sum_{t,k} \pi_k^{*,t} \log \pi_k^t,$$
 (2.32)

thereby obtaining the accumulated loss E at time step t.

Since the learning signal L_j^t is defined as the partial derivative of the error E with respect to the observable state z_j^t of a neuron j afferent to an output neurons k, we can derive

$$L_j^t = \frac{\partial E}{\partial z_j^t} = \sum_k B_{jk} \sum_{t'>t} \left(\pi_k^{t'} - \pi_k^{*,t'} \right) \kappa^{t'-t}, \tag{2.33}$$

where B_{jk} is a feedback weight from neuron k back to neuron j. There are multiple strategies for choosing feedback weights. Bellec, Scherr, Subramoney, et al. (2020) noted that a constantly uniform weight matrix yields poor performance, which has been empirically verified in this report. However, when the feedback weight matrix is initialized from a zero-centered normal distribution, it can remain constant, $(W^{\text{out}})^{\top}$, or update according to $(\Delta W^{\text{out}})^{\top}$. These variants are referred to in Bellec, Scherr, Subramoney, et al. (2020) as random, symmetric, and adaptive e-prop, respectively. In this paper, symmetric e-prop is used (i.e., $B_{jk} \stackrel{\text{def}}{=} W_{kj}^{\text{out}}$) unless explicitly stated otherwise.

Note that the term $\kappa^{t'-t}$ in Equation 2.33 is a filter that compensates for the decay factor of output neurons. Note also that this equation does not allow online learning, because future time steps t' are accessed. However, if a low-pass filter with factor κ is applied on the eligibility trace, it will cancel out the effects of the future time steps on the learning signal, and the estimated loss gradient can be approximated. This low-pass filter of the eligibility trace can be implemented in an online fashion by including it as a hidden synaptic variable \bar{e}^t_{ji} . Recall that the estimated loss gradient $\frac{dE}{dW_{ji}}$ is approximated by $\sum_t \frac{\partial E}{\partial z^t_j} e^t_{ji}$. Therefore, after inserting Equation 2.33 in Equation 2.29, the weight update is computed by

$$\Delta W_{ji} = -\eta \sum_{t'} \frac{\partial E}{\partial z_j^{t'}} e_{ji}^{t'} \tag{2.34}$$

$$= -\eta \sum_{t'} \sum_{k} B_{jk} \sum_{t'>t} \left(\pi_k^{t'} - \pi_k^{*,t} \right) \kappa^{t'-t} e_{ji}^{t'}$$
 (2.35)

$$= -\eta \sum_{k,t'} B_{jk} \sum_{t'>t} \left(\pi_k^{t'} - \pi_k^{*,t} \right) \kappa^{t'-t} e_{ji}^{t'}$$
 (2.36)

$$= -\eta \sum_{t} \underbrace{\sum_{k} B_{jk} \left(\pi_k^{t'} - \pi_k^{*,t} \right)}_{=L_j^t} \underbrace{\sum_{t' \le t} \kappa^{t'-t} e_{ji}^{t'}}_{\stackrel{\text{def}}{=} \bar{e}_{i:}^t}, \tag{2.37}$$

where W_{ji} is an input or recurrent weight. By implementing \bar{e}_{ji} as a low-pass filter (with factor κ) of the eligibility trace, the weight update in Equation 2.37 is implemented as a local and online learning algorithm.

The training algorithm for the output weights W^{out} and bias b can be directly derived from gradient descent:

$$\Delta W_{kj}^{\text{out}} = -\eta \sum_{t} \left(\pi_k^t - \pi_k^{*,t} \right) \sum_{t' \le t} \kappa^{t'-t} z_j^t \tag{2.38}$$

and

$$\Delta b_k = -\eta \sum_t \left(\pi_k^t - \pi_k^{*,t} \right). \tag{2.39}$$

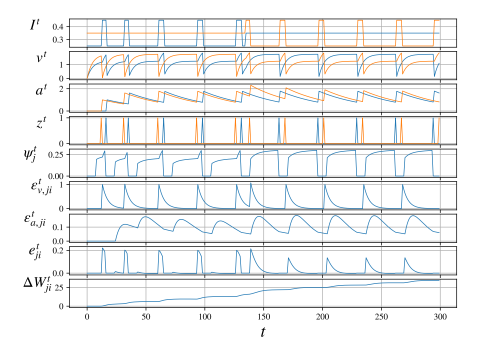


Figure 2.3: A single-synapse demonstration of the evolution of the full hidden state of the ALIF neuron. The blue lines indicate the postsynaptic neuron j, and the orange lines indicate the presynaptic neuron i. Note that the synapse weight increases regardless of the order of spikes, indicating an absence of STDP in the standard e-prop ALIF neuron.

3.1 DATA PREPROCESSING

3.1.1 The TIMIT speech corpus

TIMIT is a speech corpus that contains phonemically transcribed speech (Garofolo et al., 1993), comprising 6300 sentences, 10 spoken by each of the 630 speakers. To include a broad range of dialects all speakers lived in 8 different geographical regions in the United States (as categorized in Labov, Ash, and Boberg (2008)) during their childhood years. Table 3.1 breaks down the precise composition of the dialect distribution.

DIALECT REGION	#MALE	#FEMALE	TOTAL
1 (New England)	31 (63%)	18 (27%)	49 (8%)
2 (Northern)	71 (70%)	31 (30%)	102~(16%)
3 (North Midland)	79 (67%)	23 (23%)	102~(16%)
4 (South Midland)	69 (69%)	31 (31%)	100~(16%)
5 (Southern)	62~(63%)	36 (37%)	98 (16%)
6 (New York City)	30~(65%)	16 (35%)	46 (7%)
7 (Western)	74 (74%)	26 (26%)	100 (16%)
8	22~(67%)	11 (33%)	33 (5%)
All	438 (70%)	192 (30%)	630 (100%)

Table 3.1: Distribution of speakers' dialect regions and sexes. Speakers of the innominate dialect region 8 relocated often during their childhood.

The sentence text can be categorized into 2 dialect sentences, 450 phonetically compact sentences, and 1890 phonetically diverse sentences.

The dialect sentences, which are spoken by all speakers, are designed to expose the dialectical variants of the speakers. The phonetically compact sentences are designed to include many pairs of phones. The phonetically diverse sentences are taken from the Brown Corpus (Kucera, Kučera, and Francis, 1967) and the Playwrights Dialog (Hultzsch et al., 1964) in order to maximize the number of allophones (i. e., different phones used to pronounce the same phoneme). Table 3.2 lists an overview of the distribution of the number of speakers per sentence type.

Each of the sentences is encoded in as a waveform signal in .wav format, and is accompanied by a corresponding text file indicating which phones are pronounced in the waveform, and between which pairs of sample points.

SENTENCE TYPE	#SENTENCES	#SPEAKERS	TOTAL
Dialect	2	630	1260
Compact	450	7	3150
Diverse	1890	1	1890
Total	2342		6300

Table 3.2: Distribution of sentence types.

3.1.2 Data splitting

The TIMIT dataset is split into a training, validation and testing set as in Graves and Schmidhuber (2005) and Bellec, Scherr, Subramoney, et al. (2020). The training set is used to train the network synaptic weights according to the e-prop algorithm. The validation set is used to obtain a well-performing set of hyperparameters. The testing set is used to evaluate the performance of the network after the hyperparameters are obtained.

The TIMIT corpus documentation offers a suggested partitioning of the training and testing data, which is based on the following criteria:

- 1. 70%–80% of the data is used for training, and the remaining 20%–30% for testing.
- 2. No speaker appears in both the training and testing portions.
- 3. Both subsets include at least 1 male and 1 female speaker from every dialect region.
- 4. There is a minimal overlap of text material in the two subsets.
- 5. The test set should contain all phonemes in as many allophonic contexts as possible.

In accordance with these criteria, the TIMIT corpus includes a "core" test set that contains 2 male speakers and 1 female speaker from each dialect, summing up to 24 speakers. Each of these speakers read a different set of 5 phonetically compact sentences, and 3 phonetically diverse sentences that were unique for each speaker. Consequently, the test set comprises 192 sentences $(24 \times (5+3))$ and was selected such that it contains at least one occurrence of each phoneme. In this report, the TIMIT core test set is used, thereby meeting the criteria listed above.

The remaining 4096 sentences are randomly partitioned into 3696 training sentences and 400 validation sentences.

3.1.3 Engineering features

The data preprocessing pipeline is similar to that used in Fayek (2016), which can be summarized by applying a pre-emphasis filter on the

waveforms, then slicing the waveform in short frames, taking their short-term power spectra, computing 26 filterbanks, and finally obtain 12 Mel-Frequency Cepstrum Coefficients (MFCCs). Then, these MFCCs are aligned with the phones found in the TIMIT dataset. An example of a waveform signal is given in Figure 3.1.

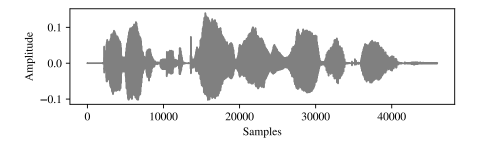


Figure 3.1: A raw waveform signal from the TIMIT dataset.

PRE-EMPHASIS In speech signals, high frequencies generally have smaller magnitudes than lower frequencies. To balance the magnitudes over the range of frequencies in the signal, a pre-emphasis filter y(t) is applied on the waveform signal x(t) defined in Equation 3.1.

$$y(t) = x(t) - 0.97x(t-1) \tag{3.1}$$

This procedure yields the additional benefit of improving the signal-tonoise ratio. An example of a pre-emphasized signal is given in Figure 3.2.

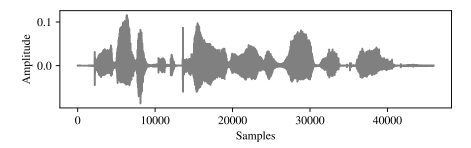


Figure 3.2: A signal after the pre-emphasis filter of Equation 3.1 was applied to it.

FRAMING The waveforms, which are sampled at a rate f_s of 16 kHz, cannot be directly used as input to the model, because they are too long—a typical sentence waveform contains in the order of tens of thousands of samples. Furthermore, the samples are not very informative, because they represent the sound wave of the uttered sound. These sounds are filtered by the shape of the vocal tract, which manifests itself in the envelope of the short time power spectrum of the sound. This power spectrum representation describes the power of the frequency components of the signal over a brief interval. The frequency components are assumed to be

stationary over short intervals—in contrast to the full sentence, which carries its meaning because it is non-stationary. Therefore, the waveform signals are transformed into series of frequency coefficients of short-term power spectra. To obtain these multiple short-term power spectra over the duration of the waveform, it is sliced into brief overlapping frames.

Every 160 samples (equivalent to $10\,\mathrm{ms}$) of a pre-emphasized signal, an interval frame of 400 samples (equivalent to $25\,\mathrm{ms}$) is extracted. This means that the frames overlap by $25\,\mathrm{ms}$. The waveform is zero-padded such that the last frame also has 400 samples. By this process, signal frames $x_i(n)$ are obtained, where n ranges over 1–400, and i ranges over the number of frames in the waveform.

Then, a Hamming window with the form

$$w[n] = a_0 - a_1 \cos\left(\frac{2\pi n}{N-1}\right),\tag{3.2}$$

is applied where N is the window length of 400 samples, $0 \le n < N$, $a_0 = 0.53836$, and $a_1 = 0.46164$. A plot of this window is given in Figure 3.3. This window is applied to reduce the spectral leakage, which

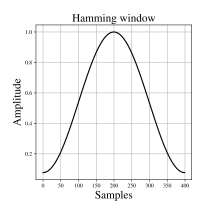


Figure 3.3: The form of the Hamming window applied on signal frames to reduce spectral leakage.

manifests itself though sidelobes in the power spectra. Applying the Hamming window reduces the sidelobes to near-equiripple conditions, minimizing the leakage (Smith, accessed December 2020).

SHORT-TERM POWER SPECTRA The power spectra P_i are obtained for each frame by first taking the absolute K-point discrete Fourier transform (DFT) of the frame samples $x_i(n)$:

$$X_k = \left| \sum_{n=0}^{N-1} x_i(n) \cdot e^{-\frac{i2\pi}{N}kn} \right|, \tag{3.3}$$

where K = 512. This yields the magnitudes of the DCT of the frames (an example is illustrated in Figure 3.4).

The power spectra are obtained using the equation

$$P = \frac{X_k^2}{K},\tag{3.4}$$

an example of which is shown in Figure 3.5.

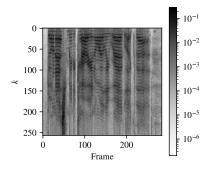


Figure 3.4: The magnitudes of the DFT of a frame.

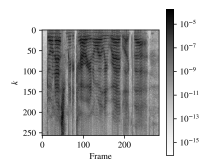


Figure 3.5: The power spectrum of a frame.

MEL FILTERBANK The short-term power spectra are then transformed to Mel-spaced filterbanks. The Mel scale is a scale of pitches that are perceptually equal in distance (Stevens, Volkmann, and Newman, 1937). This is in contrast to the frequency measurement, in which the human cochlea can better distinguish lower frequencies better than higher ones. The aim of converting to the Mel scale is to make every filterbank coefficient feature equally informative, thereby improving the learning performance of the model.

The Mel-spaced filterbank is a set of 40 triangular filters that we apply to each frame in P.

To compute the Mel-spaced filterbank, lower and upper band edges of respectively 0 Hz and $f_s/2=8\,\mathrm{kHz}$ are selected, and convert these to Mels using

$$m(f) = 2595 \log_{10} \left(1 + \frac{f}{700} \right),$$
 (3.5)

where f is the frequency in Hz. This yields a lower band edge of 0 Mels and an upper band edge of approximately 2835 Mels.

The 40 filterbanks are obtained by first spacing 42 points **m** linearly between these bounds (inclusive), thereby obtaining 40 points spaced exclusively between the bounds.

Then, each point m is converted back to Hz using

$$f = 700 \left(10^{m/2595} - 1 \right). \tag{3.6}$$

Resulting Mel-spaced frequencies f are rounded to their nearest Fourier transform bin b using

$$b = |(K+1)\mathbf{f}/fs|. \tag{3.7}$$

The resulting 40 filterbanks with their corresponding Mels and frequencies are listed in Table A.1.

The i^{th} filter in filterbank H_i is a triangular filter that has its lower boundary at b_i Hz, its peak at b_{i+1} Hz, and its upper boundary at b_{i+2} Hz. For other frequencies, they are 0. Therefore, the filterbank can be described by

$$H_{i}(k) = \begin{cases} 0 & k < b_{i} \\ \frac{k-b_{i}}{b_{i+1}-b_{i}} & b_{i} \leq k < b_{i+1} \\ 1 & k = b_{i+1} \\ \frac{b_{i+2}-k}{b_{i+2}-b_{i+1}} & b_{i+1} < k \leq b_{i+2} \\ 0 & b_{i+2} < k \end{cases}$$

$$(3.8)$$

where $0 \le k \le \frac{K}{2}$. These Mel-spaced filters are shown in Figure 3.6.

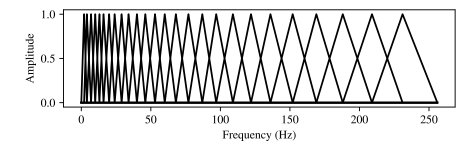


Figure 3.6: The Mel-spaced filterbanks.

After applying the filterbank to the short-term power spectrum, a spectrogram S of the frame (see e.g. Figure 3.7) is obtained.

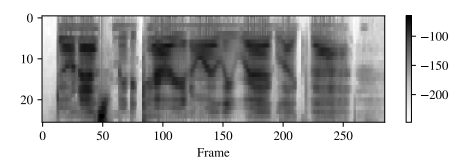


Figure 3.7: An example of a frame spectrogram.

MEL-FREQUENCY CEPSTRAL COEFFICIENTS Coefficients in the spectrograms are strongly correlated, which would negatively impact the learning performance of the model. Therefore, the DCT is applied again to decorrelate the coefficients and obtain the power cepstrum ${\cal C}$ of the speech frame:

$$C(n) = \left| \sum_{n=0}^{N-1} S(n) \cdot e^{-\frac{i2\pi}{N}kn} \right|. \tag{3.9}$$

The first coefficient in C is discarded, because it indicates the average power of the input signal and therefore carries little meaning. Coefficients higher than 13 are also discarded, because they represent only fast changes in the spectrogram and increase the complexity of the input signal while adding increasingly less meaning to it.

Then, the final MFCCs are balanced by centering each frame around the value 0. An example of the final MFCCs is given in Figure 3.8.

TARGET OUTPUT The target output of the model is a frame-wise representation of the phones that are uttered in a sentence. The TIMIT corpus contains text files indicating in what order phones occur in a sentence, and their starting and ending sample points.

These phones are discretized into frames such that they align correctly with the MFCCs. They are represented in one-hot vector encoding. Since the dataset contains 61 different phones, this is also the length of these vectors.

Figure 3.8 illustrates the waveform data and its frame-wise aligned MFCCs and target output. Note that the full dataset of features is standardized per channel; feature channels are first centered around 0, and then divided by their standard deviations.

3.2 ENHANCING E-PROP

This report examines two types of enhancements to applying the e-prop learning algorithm on the TIMIT dataset.

The first type is the effect of the neuron model; particularly, the effect of including STDP behavior is analyzed. The second type is the effect of a multi-layered architecture.

3.2.1 Multi-layer architecture

The multi-layer e-prop architecture can be described in the same formal model as its single-layer counterpart, in which the hidden state is based on temporally (i.e., online) and spatially locally available information at a neuron j:

$$\mathbf{h}_{j}^{t} = M\left(\mathbf{h}_{j}^{t-1}, \mathbf{z}^{t-1}, \mathbf{x}^{t}, \mathbf{W}_{j}\right). \tag{2.1 revisited}$$

For the multi-layer architecture, however, neurons in deeper layers no longer depend on the input, but on the observable states of the previous layer at the same time step, such that at every time step, a full pass through the network is made. We modify the indexing notation accordingly, in order to directly refer to neurons and weights in a particular layer $r \in [1 ... R]$:

$$\mathbf{h}_{rj}^{t} = \begin{cases} M\left(\mathbf{h}_{rj}^{t-1}, \mathbf{z}_{r}^{t-1}, \mathbf{x}^{t}, \mathbf{W}_{rj}\right) & \text{if } r = 1\\ M\left(\mathbf{h}_{rj}^{t-1}, \mathbf{z}_{r}^{t-1}, \mathbf{z}_{r-1}^{t}, \mathbf{W}_{rj}\right) & \text{otherwise,} \end{cases}$$
(3.10)

where \mathbf{h}_{rj}^t (resp. z_{rj}^t) is the hidden state (resp. observable state) of a neuron j in layer r. and $\mathbf{W}_{rj} = \mathbf{W}_{rj}^{\text{in}} \cup \mathbf{W}_{rj}^{\text{rec}}$ is the set of afferent weights to neuron j in layer r.

Similarly, the observable state can be modeled by

$$z_{rj}^{t} = f\left(\mathbf{h}_{rj}^{t}\right) \tag{3.11}$$

and the network output by

$$y_k^t = \kappa y_k^{t-1} + \sum_{j,r} W_{rkj}^{\text{out}} z_{rj}^t + b_k,$$
 (3.12)

where W_{rkj}^{out} is a weight between neuron j in layer r and output neuron k. Note that the summation over r entails that the output layer is connected to all neurons in all layers in the network. This allows trainable broadcast weights in earlier layers, such as those found in symmetric and adaptive e-prop.

MULTI-LAYER ALIF NEURONS An ALIF neuron in a multi-layer architecture is similar to one in a single-layer architecture (see Section 2.1.2). The only difference, apart from the layer indexing, is its activity update. For a multi-layer ALIF neuron, the activity value is given by

$$v_{rj}^{t+1} = \alpha v_{rj}^t + \sum_{i \neq j} W_{rji}^{\text{rec}} z_i^t + \sum_i W_{rji}^{\text{in}} I - z_{rj}^t v_{\text{th}},$$
 (3.13)

where

$$I = \begin{cases} x_i^{t+1} & \text{if } r = 1\\ z_{r-1,i}^{t+1} & \text{otherwise.} \end{cases}$$

$$(3.14)$$

3.2.2 Other neuron types

In this section, the multi-layer STDP-ALIF and Izhikevich neuron models are presented. The advantage of these models over the standard ALIF model is that they naturally elicit STDP. Additionally, the Izhikevich model has an implicit refractory mechanism that is built into its system of equations, making it a more biologically plausible model, as opposed

to the (STDP-)ALIF model that requires an explicit timer variable in a neuron's hidden state (but it is still local and online).

The Izhikevich e-prop neuron model was first presented by Traub et al. (2020) only in a single-synapse demonstration of its STDP properties. In this report, the performance of the e-prop Izhikevich model is experimentally validated in a learning task for the first time. Traub et al. (2020) also described the STDP-LIF, which is a non-adaptive modification of the standard LIF neuron. Here, its adaptive counterpart, the STDP-ALIF model, is derived and validated as well. This allows for a direct comparison between the ALIF and STDP-ALIF models, such that the effects of STDP can be more precisely analyzed.

3.2.2.1 STDP-LIF

The key change between the ALIF and STDP-ALIF neuron is that the latter is reset to zero at a spike event, and when its refractory period of $\delta t_{\rm ref}$ time steps ends. Recall that in contrast, the standard ALIF neuron uses a soft reset by including a term $-z_{rj}^t v_{\rm thr}$ in its activation update equation (Equation 3.13).

The activation update of the STDP-ALIF neuron is therefore:

$$v_{rj}^{t+1} = \alpha v_{rj}^{t} + \sum_{i \neq j} W_{rji}^{\text{rec}} z_{i}^{t} + \sum_{i} W_{rji}^{\text{in}} I - z_{rj}^{t} \alpha v_{rj}^{t} - z_{rj}^{t-\delta t_{\text{ref}}} \alpha v_{rj}^{t}, \quad (3.15)$$

where, again, I is the network input if r = 1, otherwise it is the observable state of the preceding layer (see Equation 3.14).

The hidden state derivative changes accordingly:

$$\frac{\partial v_{rj}^{t+1}}{\partial v_{rj}^{t}} = \alpha - z_{rj}^{t} \alpha - \alpha z_{rj}^{t-\delta t_{\text{ref}}}$$
(3.16)

$$= \alpha \left(1 - z_{rj}^t - z_{rj}^{t-\delta t_{\text{ref}}} \right). \tag{3.17}$$

Note that the absence of a_{rj}^t in the new activation update entails that the other entries of the hidden state Jacobian are equal to those of the ALIF model, i. e., $\frac{\partial v_{rj}^{t+1}}{\partial a_{rj}^t} = 0$, $\frac{\partial a_{rj}^{t+1}}{\partial v_{rj}^t} = \psi_{rj}^t$, and $\frac{\partial a_{rj}^{t+1}}{\partial a_{rj}^t} = \rho - \psi_{rj}^t \beta$.

Using these values, we compute the new eligibility trace:

$$\epsilon_{rji}^{t+1} = \frac{\partial \mathbf{h}_{rj}^{t+1}}{\partial \mathbf{h}_{rj}^{t}} \cdot \epsilon_{rji}^{t} + \frac{\partial \mathbf{h}_{rj}^{t+1}}{\partial W_{rji}}$$
(3.18)

$$= \begin{pmatrix} \alpha \left(1 - z_{rj}^t - z_{rj}^{t-\delta t_{\text{ref}}} \right) \epsilon_{rji,v}^t + z_{ri}^{t-1} \\ \psi_{rj}^t \epsilon_{rji,v}^t + \left(\rho - \psi_{rj}^t \beta \right) \epsilon_{rji,a}^t \end{pmatrix}$$
(3.19)

Note that the observable state of the afferent neuron z_{ri}^{t-1} in Equation 3.19 changes to $z_{r-1,i}^t$ if ϵ_{rji}^{t+1} corresponds to a weight between layer r-1 and r and r > 1. If the weight is instead between the network input and the first layer, then it changes to x_i^t . Note also that this corrects an error in the STDP-LIF model described in Traub et al. (2020), where

the observable state at the current time step t is accessed instead of t-1, thereby diverging from the e-prop model in Equation 3.10.

The eligibility trace remains

$$e_{rji}^{t} = \frac{\partial z_{rj}^{t}}{\partial \mathbf{h}_{rj}^{t}} \cdot \epsilon_{rji}^{t} \tag{3.20}$$

$$= \psi_{rj}^t \left(\epsilon_{rji,v}^t - \beta \epsilon_{rji,a}^t \right). \tag{3.21}$$

By resetting the neuron activation at spike time and after the refractory time, STDP is introduced in the system by clamping the pseudoderivative to a negative value, instead of 0, during the refractory time:

$$\psi_{rj}^{t} = \begin{cases} -\gamma & \text{if } t - t_{z_{rj}} < \delta t_{\text{ref}} \\ \gamma \max\left(0, 1 - \left|\frac{v_{rj}^{t} - v_{\text{th}}}{v_{\text{th}}}\right|\right) & \text{otherwise} \end{cases}$$
(3.22)

The factor of the pseudo-derivative and the eligibility vector can therefore produce both positive or negative eligibility traces and gradients (see Figure 3.10).

3.2.2.2 Izhikevich neuron

The standard system of equations of the Izhikevich neuron is described by

$$v' = 0.04v^2 + 5v + 140 - u + I (3.23)$$

$$a' = 0.004v - 0.02a, (3.24)$$

where v' and a' are the values of the activation value v and recovery variable a at the next time point, and I is the current input to the neuron. Following Traub et al. (2020), the activation reset and refractory period are built into this system of equations by replacing v and a by respectively:

$$\tilde{v}_{rj}^t = v_{rj}^t - \left(v_{rj}^t + 65\right) z_{rj}^t \tag{3.25}$$

$$\tilde{a}_{rj}^t = a_{rj}^t + 2z_{rj}^t, (3.26)$$

such that when a spike event occurs (i. e., $z_{rj}^t = 1$), the activation value is reset to its baseline value of -65, and the recovery variable increases by 2. We describe this "self-resetting" Izhikevich neuron in the context of multi-layer e-prop as follows:

$$v_{rj}^{t+1} = \tilde{v}_{rj}^t + 0.04 \left(\tilde{v}_{rj}^t \right)^2 + 5\tilde{v}_{rj}^t + 140 - \tilde{u}_{rj}^t + I_{rj}^t$$
(3.27)

$$u_{rj}^{t+1} = \tilde{u}_{rj}^t + 0.004\tilde{v}_{rj}^t - 0.02\tilde{u}_{rj}^t. \tag{3.28}$$

The partial derivatives of the hidden state \mathbf{h}_{rj}^t can then be computed:

$$\frac{\partial v_{rj}^{t+1}}{\partial v_{rj}^{t}} = \left(1 - z_{rj}^{t}\right) \left(6 + 0.08v_{rj}^{t}\right) \tag{3.29}$$

$$\frac{\partial a_{rj}^{t+1}}{\partial v_{rj}^t} = -1 \tag{3.30}$$

$$\frac{\partial v_{rj}^{t+1}}{\partial a_{rj}^{t}} = 0.004 \left(1 - z_{rj}^{t} \right) \tag{3.31}$$

$$\frac{\partial a_{rj}^{t+1}}{\partial a_{rj}^t} = 0.98. \tag{3.32}$$

Using these values, we compute the new eligibility trace:

$$\epsilon_{rji}^{t+1} = \frac{\partial \mathbf{h}_{rj}^{t+1}}{\partial \mathbf{h}_{rj}^{t}} \cdot \epsilon_{rji}^{t} + \frac{\partial \mathbf{h}_{rj}^{t+1}}{\partial W_{rji}}$$
(3.33)

$$= \begin{pmatrix} \left(1 - z_{rj}^t\right) \left(6 + 0.08v_{rj}^t\right) \epsilon_{rji,v}^t - \epsilon_{rji,a}^t + z_{ri}^{t-1} \\ 0.004 \left(1 - z_{rj}^t\right) \epsilon_{rji,v}^t + 0.98\epsilon_{rji,a}^t \end{pmatrix}$$
(3.34)

As in Traub et al. (2020), the pseudo-derivative is defined as

$$\psi_{rj}^t = \gamma \exp\left(\frac{\min\left(v_{rj}^t, 30\right) - 30}{30}\right),\tag{3.35}$$

such that

$$\begin{pmatrix} \frac{\partial z_{r_j}^t}{\partial v_{r_j}^t} \\ \frac{\partial z_{r_j}^t}{\partial u_{r_j}^t} \end{pmatrix} = \begin{pmatrix} \psi_{rj}^t \\ 0 \end{pmatrix} \tag{3.36}$$

and therefore only $\epsilon_{rji,v}^t$ is used in computing the eligibility trace:

$$e_{rji}^{t} = \left(\psi_{rj}^{t} \ 0\right) \begin{pmatrix} \epsilon_{rji,v}^{t} \\ \epsilon_{rji,a}^{t} \end{pmatrix} \tag{3.37}$$

$$= \psi_{rj}^t \epsilon_{rji,v}^t. \tag{3.38}$$

However, when inserting these equations in a single-synapse demo, the eligibility vector assumes extremely high or low values (see Figure A.1). This suggests that the Izhikevich neuron does not fit the e-prop framework well. In this report, this is corrected by clipping the value of $\epsilon_{ji,v}^t$ to [-3,3] and $\epsilon_{ji,a}^t$ to [-0.005,0.005]. This correction yields the desired STDP behavior (see Figure 3.11).

3.3 REGULARIZATION

Firing rate regularization and L2 regularization are applied to improve the stability of the learning process and the generalizability of the resulting model. These two regularization methods are motivated by biological plausibility, ease of implementation in the e-prop framework, and improved empirical performance.

3.3.1 Firing rate regularization

A firing rate regularization term is added to individually control the spike frequencies of the neurons. Since spikes in a neuromorphic architecture cost energy, the practical motivation for this regularization term is that it increases the energy efficiency of the model. The biological motivation is that the firing rate of biological neurons is also under homeostatic control (Erickson et al., 2006).

Firing rate is implemented by adding a regularization term E_{reg} to the loss function that penalizes neurons that have a firing rate that is too low or too high:

$$E_{\text{reg}} = \frac{1}{2} \sum_{i} \left(f^{\text{target}} - f_{rj}^{\text{av},t} \right)^{2}, \tag{3.39}$$

where f^{target} is a target firing rate of 10 Hz, and

$$f_{rj}^{\text{av},t} = \frac{1}{t} z_{rj}^{\text{total},t} \tag{3.40}$$

is the running average spike frequency, where $z^{\text{total},t}$ accumulates spikes emitted by neuron j in layer r up to (and including) time step t. Note that $z^{\text{total},0} = 0$, i. e., the accumulation resets at every new training sample. By implementing this sum as a hidden variable, e-prop remains an online training algorithm when firing rate regularization is implemented.

To insert the regularization term into the e-prop framework, we compute the weight update that regularizes the firing rate toward f^{target} through gradient descent, similarly to the main e-prop weight update (Equation 2.29):

$$\frac{\partial E_{\text{reg}}}{\partial z_{rj}^t} = \left(f^{\text{target}} - f_{rj}^{\text{av},t} \right). \tag{3.41}$$

Note that this regularization loss differs from the firing rate regularization in Bellec, Scherr, Subramoney, et al. (2020), in which the firing rate is calculated in an offline fashion, by retroactively computing the average firing rate based on all spikes instead of only accumulated spikes. Note also that in Bellec, Scherr, Subramoney, et al. (2020), $\frac{\partial E_{\text{reg}}}{\partial z_{rj}^t}$ is multiplied with the eligibility trace e_{rji}^t , as in Equation 2.29 to obtain the weight update, whereas in this report, the eligibility trace is omitted, resulting in a number of benefits:

- 1. It allows silent neurons that have infrequently spiking afferent neurons to more easily increase their firing rate, because their low afferent eligibility traces no longer nullifies the regularization gradient, and thereby results in a better empirical learning performance;
- 2. It is more efficient in emulations on von Neumann machines, because the element-wise multiplication of $\frac{\partial E_{\text{reg}}}{\partial z_{rj}^t}$ and the eligibility trace is a relatively large computation on the order $\Theta(n^2)$ that no longer needs to be computed.

3. It is more intuitive, as only the gradient of the firing rate is used to compute the weight update.

We apply the weight update ΔW_{rji} of the regularization gradient using

$$\Delta_{\text{reg}} W_{rji} = -\eta \ c_{\text{reg}} \sum_{t} \left(f^{\text{target}} - f_{rj}^{\text{av},t} \right). \tag{3.42}$$

Note that the regularization gradients can be combined and accumulated over time on the same synaptic variable as the normal gradients, facilitating practical implementation of the learning procedure in both software emulations and neuromorphic embeddings:

$$\Delta W_{rji} = -\eta \sum_{t} \left(c_{\text{reg}} \left(f^{\text{target}} - f_{rj}^{\text{av},t} \right) + L_{rj}^{t} \cdot \bar{e}_{rji}^{t} \right). \tag{3.43}$$

3.3.2 L2 regularization

To regularize weights around 0, a small fraction (parametrized by c_{L2}) of the value of the weight is added to its gradient value at every weight update:

$$\Delta_{L2}W_{rji} = -\eta \ c_{L2} \cdot W_{rji},\tag{3.44}$$

which can be added to the full weight update as an extra term:

$$\Delta W_{rji} = -\eta \left(c_{L2} \cdot W_{rji} + \sum_{t} \left(c_{reg} \left(f^{target} - f_{rj}^{av,t} \right) + L_{rj}^{t} \cdot \bar{e}_{rji}^{t} \right) \right).$$

$$(3.45)$$

The statistical motivation for this extra regularization term is that by softly restricting the weights, the network is less likely to overfit on the training data.

The biological motivation is that biological synapses are regularized by a multiplicative factor to decrease the strength of individual synapses to the same proportion (Turrigiano and Nelson, 2000), likely counteracting the run-away effects that the positive feedback of STDP naturally induces (Siddoway, Hou, and Xia, 2014). Moreover, there are natural bounds of the strength of a biological synapse, measured by the amplitude of the postsynaptic potential, because the number of neurotransmitter vesicles and release sites is physically limited (Del Castillo and Katz, 1954). These natural limits are approximately 0.4 mV to 20 mV (Diaz-Rios and Miller, 2006).

3.4 OPTIMIZER

For simplicity, in this report weight updates are described using stochastic gradient descent:

$$\Delta W_{rji} = -\eta \sum_{t} L_{rj}^{t} \cdot \bar{e}_{rji}^{t}. \tag{3.46}$$

However, the results described in Chapter 4 are obtained using Adam (or Adaptive Moment Estimation) (Kingma and Ba, 2014). This optimization method tracks running averages of the gradient and its second moment (resp. M_{rji} and V_{rji}), and fits the local and online constraints of e-prop, because the running averages are tracked per individual synapse. The Adam weight update in the context of multi-layer e-prop is given by:

$$M_{rji}^{(i+1)} = \beta_1 M_{rji}^{(i)} + (1 - \beta_1) G_{rji}^{(i)}$$
(3.47)

$$V_{rji}^{(i+1)} = \beta_2 V_{rji}^{(i)} + (1 - \beta_2) \left(G_{rji}^{(i)} \right)^2$$
(3.48)

$$\widehat{M}_{rji} = \frac{M_{rji}^{(i+1)}}{1 - \beta_1^{i+1}} \tag{3.49}$$

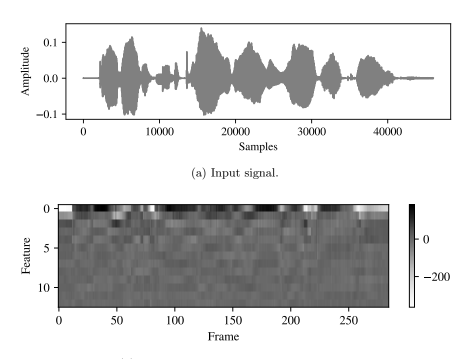
$$\widehat{V}_{rji} = \frac{V_{rji}^{(i+1)}}{1 - \beta_2^{i+1}} \tag{3.50}$$

$$\widehat{V}_{rji} = \frac{V_{rji}^{(i+1)}}{1 - \beta_2^{i+1}}$$

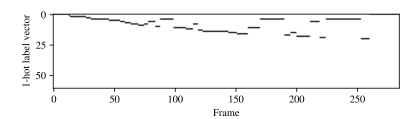
$$\Delta W_{rji}^{(i+1)} = -\eta \frac{\widehat{M}_{rji}}{\sqrt{\widehat{V}_{rji} + 10^{-5}}},$$
(3.50)

where $G^{(i)}$ is the estimated gradient $\sum_t L_{rj}^t \cdot \bar{e}_{rji}^t$ at weight update i, and $\beta_1 = 0.9$ and $\beta_2 = 0.999$ are the forgetting factors for the gradient and its second moment, respectively. The firing rate and L2 regularization terms are omitted here for clarity. Note that the forgetting factors are not indexed, but raised to the power of i + 1, in computing the biascorrected estimates \widehat{M}_{rji} and \widehat{V}_{rji} . Note also that minibatches of size 32 are used to more accurately estimate the gradient and enable a stabler descent in the error landscape. The value of $G_{rji}^{(i)}$ is computed as the mean over the minibatch.

Note that the learning rate is linearly ramped up from 0 to η during the first epoch, such that the initial minibatches are used to aggregate good initial momentum buffers, as the variance is higher when fewer minibatches are processed. This "warming up" of the learning rate is a variance reduction technique that has shown beneficial results in training other models (L. Liu et al., 2019). Empirical observations on the resulting learning curves (see Chapter 4) suggest that this procedure does not obstruct a rapid initial decrease of the loss function.



(b) MFCCs centered around 0 per channel.



(c) Target signal encoded as a one-hot vector changing over time.

Figure 3.8: An alignment of a sample signal with its MFCCs features and target phones.

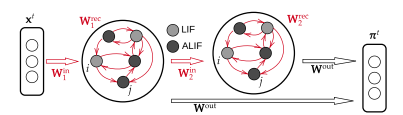


Figure 3.9: An illustration of a multi-layer network architecture. Some details shown in Figure $(2.2\ b)$ are omitted for clarity

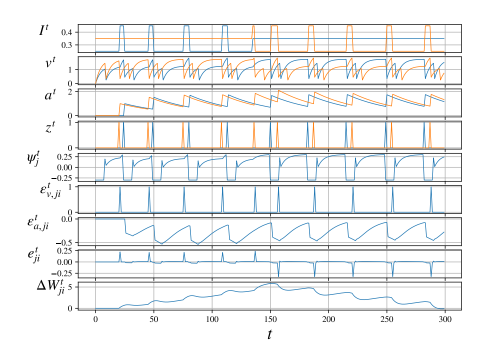


Figure 3.10: A single-synapse demonstration of the STDP-ALIF.

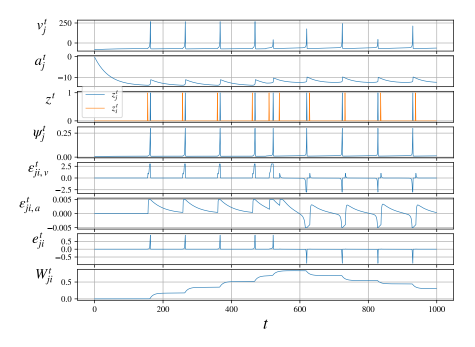


Figure 3.11: A single-synapse demo of the corrected Izhikevich neuron.

In this chapter, the learning performance and regularization behavior of the ALIF, STDP-ALIF, and Izhikevich neurons are compared. Then, the effect of stacking multiple recurrent layers on the learning performance and speed is examined.

4.1 COMPARING NEURON MODELS

ACCURACY The main outcome of the neuron model comparison is that the Izhikevich neuron type performs poorly compared to the ALIF and STDP-ALIF neuron models, which reach a similar classification performance. In Figure 4.1a the Izhikevich neuron reaches a misclassification rate of 94.2% on the test set, which is only slightly better than constantly guessing the most frequent class. The ALIF neuron reaches a test misclassification rate of 50% in relatively few iterations, and the STDP-ALIF stably reaches 50%, and likely lower if training was continued for longer. Note that the test performance was obtained on the model with the best validation accuracy.

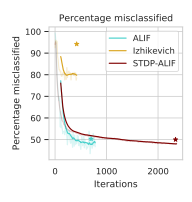
Figure 4.1b illustrates the decrease of the cross-entropy score, which for the ALIF and STDP-ALIF neurons is comparable to that of the misclassification rate. The cross-entropy of the Izhikevich neuron continues to decrease while its classification accuracy stalls, suggesting that it trains its bias toward more frequent phone classes rather than learning a relationship between input MFCCs and classes.

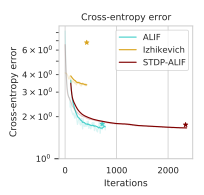
FIRING RATE Figure 4.2a illustrates the effect of the firing regularization term. It can be observed that the ALIF and STDP-ALIF neuron models are able to quickly change their mean spiking frequencies to the desired target frequency of 10 Hz, but the Izhikevich neuron increases past this rate to a spiking frequency of approximately 18 Hz.

Figure 4.2b illustrates the decrease of the regularization error. The ALIF neuron appears to slightly better adapt to the regularization term. The Izhikevich neuron, again, underperforms significantly.

4.2 COMPARING NETWORK DEPTH

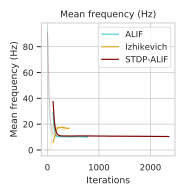
From the comparison between the network depth in Figures 4.3a–4.3c, it can be concluded that single-layer networks train considerably more efficiently and accurately than multi-layer networks. This holds for all tested neuron types. The cross-entropy error, spiking frequency, and regularization error are also better for single-layer networks (see Figure A.2).

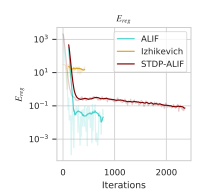




- (a) Percentage of samples wrongly classified.
- (b) Cross-entropy loss (log-scaled).

Figure 4.1: Classification performance on the validation data for each of the three neuron models in a single-layer e-prop model. The opaque lines indicate the running average of the real validation scores indicated by the transparent lines. The star symbols indicate the performances on the test set, with a misclassification rate of 92.4% for the Izhikevich neuron, 50.3% for the ALIF neuron, and 50% for the STDP-ALIF neuron type.





- (a) Mean spiking frequency.
- (b) Regularization error.

Figure 4.2: Effect of firing rate regularization on the validation data for each of the three neuron models.

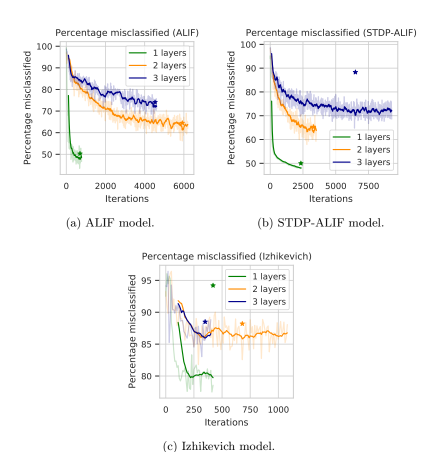


Figure 4.3: Accuracy comparison on the validation data between single- and multi-layer e-prop models.

INTERPRETATION OF RESULTS

POSSIBLE IMPROVEMENTS There are many hypothetical ways of improving the performance or biological plausibility of e-prop that have not yet been considered in this report. For instance, a likely reason that the learning speed of the multi-layer architectures was slower than their single-layer counterparts is that the weights are poorly initialized. Empirical observation of the learning process suggested that spiking activity faded in deeper layers, because the spiking activity from a preceding layer is generally weaker than the input values the first layer receives. Higher weights in-between layers mitigate this fading activity, but require some search to find a good value. In this report, the firing rate regularization term approximated this value, but learning is more efficient with a better initialization, since initial synaptic weights significantly affect the performance of STDP-based SNNs (J. Kim et al., 2020).

Also in this report, certain parameters such as firing rate targets (f^{target}) , activity leak (α) , threshold adaptivity (β) , and feedback signals were constant for all neurons, except the threshold adaptivity, which was 0 for some neurons to emulate non-adaptive LIF neurons. However, future research could examine the effects of sampling some of these parameters from a distribution for each neuron, thereby creating a more diverse population of neurons with different time scales. This sampling requires a careful assessment of the time scales related to the learning task; in particular, the network should be able to process the slowest relevant time scale of the task (Jaeger et al., 2021). In the TIMIT classification task, for instance, this could span the whole time fragment, because initial words give semantic context to all subsequent words. Note also that later words give informative context to earlier words. In Bellec, Scherr, Subramoney, et al. (2020), this context was captured in a bidirectional network, improving the accuracy by nearly 10%. In this report, this was empirically validated as well, but left untreated because a bidirectional network is not biologically plausible, as accessing earlier input values violates the "online" constraint. According to Bellec, Scherr, Subramoney, et al. (2020), e-prop suggests that the experimentally found diverse time constants of the firing activity of populations of neurons in different brain areas (Runyan et al., 2017) are correlated with their capability to handle corresponding ranges of delays in temporal credit assignment for learning. Setting different values for these parameters per layer might also have a beneficial effect; Ahmed et al. (1998) suggested that deeper layers display slower and weaker adaptation rates than shallow layers.

Random dropout of 80% of the recurrent network weights led to better training and validation performances. This suggests that the

effects of the topology within a layer might positively affect the learning process. In the brain, neurons tend to connect to nearby neurons. A simple lattice topology might better approximate the connectivity of the brain, decrease the computational complexity in both emulations in von Neumann machines, and allow easier on-chip implementations in neuromorphic hardware. Hierarchical clustering of neurons might also have a beneficial effect, as this has been demonstrated to improve R-STDP in SNNs (Weidel, Duarte, and Morrison, 2020) and address the scalability issue of SNNs (Carrillo et al., 2012). Neuromorphic computation allows complex network operations (Hasler and Akers, 1990), large-scale conductance-based SNNs (Yang, B. Deng, et al., 2019; Yang, J. Wang, et al., 2019) and asynchronous communication in VLSIs through address event representation (Deiss et al., 1999; Lazzaro et al., 1993).

The connection topology might also change over time through a dynamic pruning and growing of weights and neurons. Here, the biological motivation is that the human brain prunes synaptic connections during early development (Huttenlocher et al., 1979). Elbez et al. (2020) demonstrated that 75% of a SNN can be compressed while preserving its performance, but it is not clear if this can be applied in a biologically plausible way in the e-prop framework. However, integrating stochastic synaptic rewiring (Kappel et al., 2018) into an ALIF network can improve its short-term memory (Bellec, Scherr, Subramoney, et al., 2020).

Finally, synaptic delay might improve the temporal processing power of an e-prop model. In this report, communication between neurons was transmitted as a spike over a synapse with a delay of 1 ms. This delay could be variable (e.g., between 1 and 10 ms), such that potentially informative past inputs are more accurately preserved in synaptic delays, rather than only in eligibility traces and activity loops. This can help deal with tasks that require processing information on multiple time spans (Jaeger et al., 2021). This resembles the variable physical length of myelinated biological synapses and number of nodes of Ranvier along them, affecting the conductance of the action potential from one neuron to another (Bean, 2007).

FUTURE DIRECTIONS As neuromorphic computing matures, neuroscience improves, and DL increasingly hits fundamental limitations, there is an exciting future for biologically plausible SNNs. There is much to gain from cross-fertilization between these fields. The popularity of DL was accelerated by accessible platforms to implement and deploy ANNs. Similar high-level simulation platforms are now in active development, which can integrate the typical behavior of memristive device models into crossbar architectures within DL systems (Lammie et al., 2020).

Recent advances in neuromorphic computing indicate this increasing popularity. Neuromorphic architectures have been used for mapless navigation with 75 times lower power and better performance (Tang, Kumar, and Michmizos, 2020); as low-power solutions for simultaneous localization and mapping of mobile robots (Tang, Shah, and Michmizos,

2019), for planning (Fischl et al., 2017), and control (Blum et al., 2017); and self-repairing SNN for fault detection (Zhu et al., 2017). While crossfertilization between neuromorphic computing and quantum computing is starting to take place (Russek et al., 2016), as quantum superposition and entanglement can be used to process information in parallel and in a high-dimensional state space (Fujii and Nakajima, 2017; Tacchino et al., 2019; Yamamoto et al., 2017), more physics and material science is required to build efficient neuromorphic architectures (Marković et al., 2020). The same holds for the cross-fertilization between neuroscience and learning rules of biologically plausible SNNs. Nanodevices that emulate biological synapses with learning functions can benefit neuromorphic architectures (Ren et al., 2018; Y. Wang et al., 2018; Yao et al., 2017), particularly the two-terminal memristor (Jo et al., 2010; Z. Wang et al., 2017). However, the learning principles of biological NNs are not explored enough to design engineering solutions (Gorban, Makarov, and Tyukin, 2019; Taherkhani et al., 2018). Feedback connections, for which the brain uses neurotransmitters, may become particularly problematic in large-scale neuromorphic systems. Another issue in analog computation is how to match the system's internal temporal processing to that of its inputs. Emulating neural dynamics on a physical substrate is more efficient but requires constraints to match the brain's timescales (Jaeger et al., 2021; Mead, 1990). Future work on e-prop could explore a combination with attention-based models in order to cover multiple timescales (Bellec, Scherr, Subramoney, et al., 2020).

6

CONCLUSION

- CONCLUSION PLACEHOLDERS



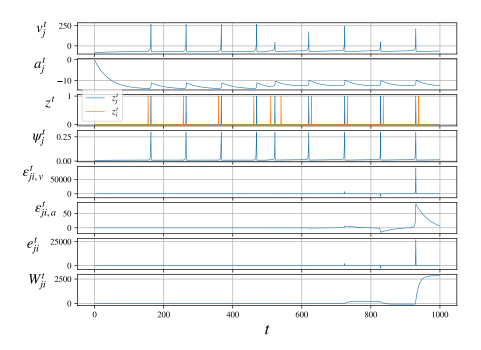


Figure A.1: A single-synapse demo of the uncorrected Izhikevich neuron. Note that $\epsilon^t_{ji,v}$ takes on extreme values and that the eligibility vector flips sign at any pair of spikes.

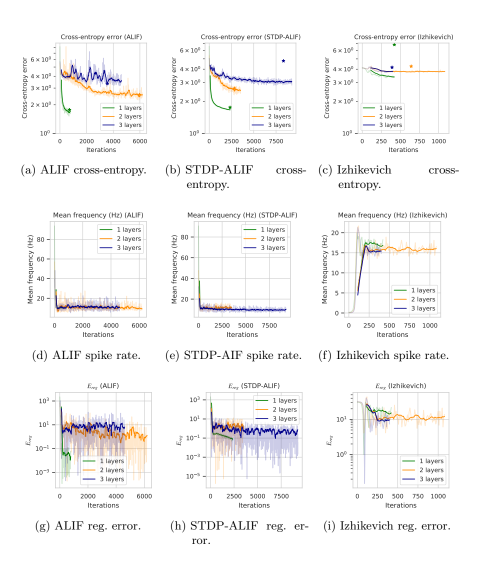


Figure A.2: Cross-entropy rates, mean spiking frequencies, and regularization errors for multi-layer networks.

MELS	ΗZ	FILTERBANK	
0	0	0	
105	68.5	2	
210	143.7	4	
315	226.2	7	
420	316.8	10	
525	416.3	13	
630	525.5	16	
735	645.4	20	
840	777	24	
945	921.5	29	
1050	1080.1	34	
1155	1254.4	40	
1260	1445.4	46	
1365	1655.3	53	
1470	1885.7	60	
1575	2138.6	68	
1680	2416.3	77	
1785	2721.2	87	
1890	3055.9	97	
1995	3423.3	109	
2100	3826.7	122	
2205	4269.5	136	
2310	4755.7	152	
2415	5289.4	169	
2520	5875.3	188	
2625	6518.6	209	
2730	7224.8	231	
2835	8000	256	

Table A.1: Conversion table between linearly spaced Mels and their corresponding frequencies and filterbank boundaries.

SYMBOL	DESCRIPTION	VALUE
α	Activity leak	0.8
β	Adaptivity	0.184
ho	Adaptivity leak	0.975
κ	Output decay	0.8
γ	Pseudoderivative dampening	0.3
$v_{ m th}$	Base threshold	0.95
$\delta t_{ m ref}$	Refractory time	2
η	Learning rate	0.01
β_1	Adam momentum factor 1	0.9
β_2	Adam momentum factor 2	0.999
$c_{ m reg}$	Firing rate regularization	50
$c_{ m L2}$	L2 regularization	10^{-5}
$f^{ m target}$	Target firing rate	0.01
N	Network size	800*

Table A.2: The full list of hyperparameter values. * This is the total number of neurons in multi-layer networks.

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