# A solution to the learning dilemma for recurrent networks of spiking neurons

Guillaume Bellec<sup>1,o</sup>, Franz Scherr<sup>1,o</sup>, Anand Subramoney<sup>1</sup>, Elias Hajek<sup>1</sup>,
 Darjan Salaj<sup>1</sup>, Robert Legenstein<sup>1</sup> & Wolfgang Maass<sup>1,\*</sup>

<sup>5</sup> <sup>1</sup>Institute of Theoretical Computer Science, Graz University of Technology,

<sup>6</sup> Inffeldgasse 16b, Graz, Austria

<sup>7</sup> ° Equal contributions.

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<sup>8</sup> \* To whom correspondence should be addressed; E-mail: maass@igi.tugraz.at.

#### Abstract

Recurrently connected networks of spiking neurons underlie the astounding infor-10 mation processing capabilities of the brain. But in spite of extensive research, it has 11 remained open how they can learn through synaptic plasticity to carry out complex 12 network computations. We argue that two pieces of this puzzle were provided by 13 experimental data from neuroscience. A new mathematical insight tells us how these 14 pieces need to be combined to enable biologically plausible online network learning 15 through gradient descent, in particular deep reinforcement learning. This new learn-16 ing method – called e-prop – approaches the performance of BPTT (backpropagation 17 through time), the best known method for training recurrent neural networks in ma-18 chine learning. In addition, it suggests a method for powerful on-chip learning in 19 novel energy-efficient spike-based hardware for AI. 20

## <sup>21</sup> Introduction

Networks of neurons in the brain differ in at least two essential aspects from deep neural 22 networks in machine learning: They are recurrently connected, forming a giant number 23 of loops, and they communicate via asynchronously emitted stereotypical electrical pulses, 24 called spikes, rather than bits or numbers that are produced in a synchronized manner 25 by each layer of a feedforward deep network. Models that capture primary information 26 processing capabilities of spiking neurons in the brain are well known, and we consider 27 the arguably most prominent one: leaky integrate-and-fire (LIF) neurons, where spikes 28 that arrive from other neurons through synaptic connections are multiplied with the corre-29 sponding synaptic weight, and are linearly integrated by a leaky membrane potential. The 30 neuron fires – i.e., emits a spike – when the membrane potential reaches a firing threshold. 31

But it is an open problem how recurrent networks of spiking neurons (RSNNs) can learn, 32 i.e., how their synaptic weights can be modified by local rules for synaptic plasticity so that 33 the computational performance of the network improves. In deep learning this problem 34 is solved for feedforward networks through gradient descent for a loss function E that 35 measures imperfections of current network performance [1]. Gradients of E are propagated 36 backwards through all layers of the feedforward network to each synapse through a process 37 called backpropagation. Recurrently connected networks can compute more efficiently 38 because each neuron can participate several times in a network computation, and they are 39 able to solve tasks that require integration of information over time or a non-trivial timing 40 of network outputs according to task demands. But since each synaptic weight can affect 41 the network computation at several time points during a recurrent network computation, 42 its impact on the loss function (see Fig. 1a) is more indirect, and learning through gradient 43 descent becomes substantially more difficult. This learning problem is aggravated if there 44 are slowly changing hidden variables in the neuron model, such as neurons with spike-45 frequency adaptation (SFA). Neurons with SFA are quite common in the neocortex [2], 46 and it turns out that their inclusion in the RSNN significantly increases the computational 47 power of the network [3]. In fact, RSNNs trained through gradient descent acquire then 48 similar computing capabilities as networks of LSTM (Long Short-Term Memory) units, 49 the state of the art for recurrent neural networks in machine learning. Because of this 50 functional relation to LSTM networks these RSNN models are referred to as LSNNs [3]. 51

In machine learning one trains recurrent neural networks by unrolling the network into a virtual feedforward network [1], see Fig. 1b, and applying the backpropagation algorithm to that (Fig. 1c). This learning method for recurrent neural networks is called backpropagation through time (*BPTT*) since it requires propagation of gradients backwards in time with regard to the network computation.

With a careful choice of the pseudo-derivative for handling the discontinuous dynamics of spiking neurons one can apply *BPTT* also to RSNNs, and RSNNs were able to learn in this way for the first time to solve really demanding computational tasks (see [3], [4] for preceding results). But the dilemma is that *BPTT* requires storing the intermediate states of all neurons during a network computation, and merging these in a subsequent offline process with gradients that are computed backwards in time (see Fig. 1c, Movie S1 and Movie S2). This makes it very unlikely that *BPTT* is used by the brain [5].

<sup>64</sup> We present a solution to this dilemma in the form of a biologically plausible method for <sup>65</sup> online network learning through gradient descent: *e-prop* (Fig. 1d, see Movie S3). *E-prop* <sup>66</sup> is motivated by two streams of experimental data from neuroscience:

i) Neurons in the brain maintain traces of preceding activity on the molecular level, for
example in the form of calcium ions or activated CaMKII enzymes [6]. In particular,
they maintain a fading memory of events where the presynaptic neuron fired before
the postsynaptic neuron, which is known to induce synaptic plasticity if followed by
a top-down learning signal [7, 8, 9]. Such traces are often referred to as eligibility
traces.

ii) In the brain there exists an abundance of top-down signals such as dopamine, acetyl-

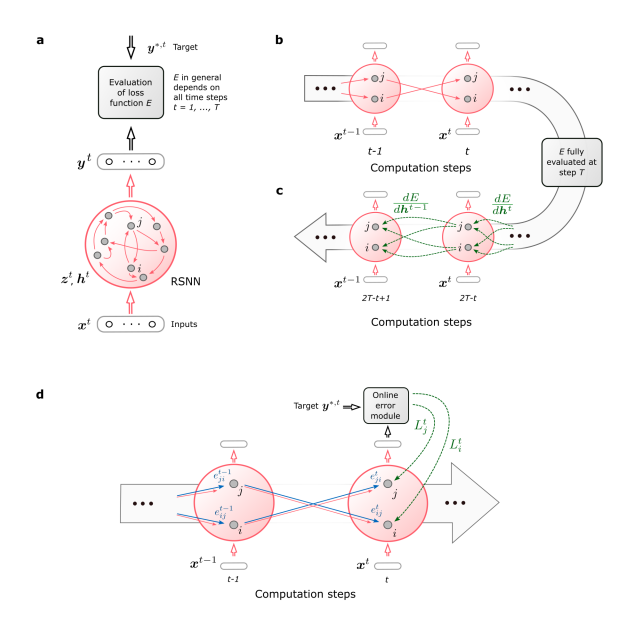


Figure 1: Schemes for RSNNs, *BPTT*, and *e-prop.* a) RSNN with network inputs x, neuron spikes z, and output targets  $y^*$ , for each time step t of the RSNN computation. Output neurons y provide a low-pass filter of a weighted sum of network spikes z. b) *BPTT* computes gradients in the unrolled version of the network. It has a new copy of the neurons of the RSNN for each time step t. A synaptic connection from neuron i to neuron j of the RSNN is replaced by an array of feedforward connections, one for each time step t, that goes from the copy of neuron i in the layer for time step t + 1. All synapses in this array have the same weight: the weight of this synaptic connection in the RSNN. c) Loss gradients of *BPTT* are propagated backwards in time and retrograde across synapses in an offline manner, long after the forward computation has passed a layer. d) Online learning dynamics of *e-prop*. Feedforward computation of eligibility traces is indicated in blue. These are combined with online learning signals according to equation (1).

choline, and neural firing [10] related to the event-related negativity (ERN), that
inform local populations of neurons about behavioral results. Furthermore dopamine
signals [11, 12] have been found to be specific for different target populations of neurons, rather than being global. We refer in our learning model to such top-down
signals as learning signals.

A re-analysis of the mathematical basis of gradient descent in recurrent neural networks 79 tells us how local eligibility traces and top-down learning signals should be optimally com-80 bined to enable network learning through gradient descent – without requiring backpro-81 gation of signals through time. The resulting new learning method, *e-prop*, learns slower 82 than BPTT, but tends to approximate the performance of BPTT, thereby providing a first 83 solution to the learning dilemma for RSNNs. Furthermore *e-prop* also works for RSNNs 84 with more complex neuron models, such as LSNNs. This new learning paradigm for brain-85 like network models elucidates how the brain could learn to recognize phonemes in spoken 86 language (Fig. 2), solve temporal credit assignment problems (Fig. 3), and acquire new 87 behaviors just from rewards (Fig. 4, 5). 88

In such reinforcement learning (RL) tasks the learner needs to explore its environment, 89 and find out which action gets rewarded in what state [13]. There is no "teacher" that 90 tells the learner what action would be optimal; in fact, the learner may never find that 91 out. Nevertheless learning methods such as BPTT are essential for a powerful form of 92 RL that is often referred to as Deep RL [14]. There one trains recurrent artificial neural 93 networks with internally generated teaching signals. We show here that Deep RL can in 94 principle also be carried out by neural networks of the brain, since *e-prop* approximates the 95 performance of *BPTT* also in this RL context. However another new ingredient is needed 96 to prove that. Previous work on Deep RL for solving complex tasks, such as winning 97 Atari games [14], required additional mechanisms to avoid well-known instabilities that 98 arise from using nonlinear function approximators, such as the use of several interacting 99 learners in parallel. Since this parallel learning scheme does not appear to be biologically 100 plausible, we introduce here a new method for avoiding learning instabilities: We show that 101 a suitable schedule for the lengths of learning episodes and learning rates also alleviates 102 learning instabilities in Deep RL. 103

We are not aware of previous work on online gradient descent learning methods for RSNNs, neither for supervised learning nor for RL. There exists however preceding work on online approximations of gradient descent for non-spiking neural networks based on [15], which we review in the Discussion section.

The previous lack of powerful learning methods for RSNNs also affected the develop-108 ment and use of neuromorphic computing hardware, which aims at a drastic reduction 109 in the energy consumption of AI implementations. A substantial fraction of this neuro-110 morphic hardware, such as SpiNNaker [16] or Intel's Loihi chip [17], implements RSNNs 111 and aims at on-chip training of these RSNNs. Although it does not matter here whether 112 the learning algorithm is biologically plausible, the excessive storage and offline processing 113 demands of *BPTT* make this option unappealing for neuromorphic hardware. Hence there 114 also exists a learning dilemma for RSNNs in neuromorphic hardware, which can be solved 115

<sup>116</sup> with *e*-prop.

## 117 **Results**

#### <sup>118</sup> Mathematical basis for e-prop

Spikes are modeled as binary variables  $z_j^t$  that assume value 1 if neuron j fires at time t, 119 otherwise value 0. It is common in models to let t vary over small discrete time steps, e.g. 120 of 1 ms length. The goal of network learning is to find synaptic weights W that minimize 121 a given loss function E. E may depend on all or a subset of the spikes in the network. 122 E measures in the case of regression or classification learning the deviation of the actual 123 output  $y_k^t$  of each output neuron k at time t from its given target value  $y_k^{*,t}$  (Fig. 1a). 124 In reinforcement learning (RL), the goal is to optimize the behavior of an agent in order 125 to maximize obtained rewards. In this case, E measures deficiencies of the current agent 126 policy to collect rewards. 127

The gradient  $\frac{dE}{dW_{ji}}$  for the weight  $W_{ji}$  of the synapse from neuron *i* to neuron *j* tells us how this weight should be changed in order to reduce *E*. The key innovation is that a rigorous proof (see Methods) shows that this gradient can be represented as a sum over the time steps *t* of the RSNN computation, where the second factor is just a local gradient that does not depend on *E*:

$$\frac{dE}{dW_{ji}} = \sum_{t} \frac{dE}{dz_j^t} \cdot \left[\frac{dz_j^t}{dW_{ji}}\right]_{\text{local}} .$$
(1)

This local gradient can be represented as a sum of products of partial derivatives concern-133 ing the hidden state of neuron i up to time t (equation (13)), which can be updated during 134 the forward computation of the RNN by a simple recursion (equation (14)). This term 135  $\left[\frac{dz_j^i}{dW_{ji}}\right]_{\text{local}}$  is not an approximation. Rather, it collects the maximal amount of informa-136 tion about the network gradient  $\frac{dE}{dW_{ii}}$  that can be computed locally in a forward manner. 137 Therefore it is the key-factor of *e-prop*. Since it reduces for simple neuron models – whose 138 internal state is fully captured by its membrane potential – to a variation of terms that 139 are commonly referred to as eligibility traces for synaptic plasticity [9], we also refer to 140

$$e_{ji}^{t} \stackrel{\text{def}}{=} \left[ \frac{dz_{j}^{t}}{dW_{ji}} \right]_{\text{local}} \tag{2}$$

as eligibility trace. But most biological neurons have additional hidden variables that change on a slower time scale, such as for example the firing threshold of a neuron with firing threshold adaptation. Furthermore these slower processes in neurons are essential for attaining with spiking neurons similarly powerful computing capabilities as LSTM networks [3]. Hence the form that this eligibility trace  $e_{ji}^t$  takes for adapting neurons (see equation (25)) is essential for understanding *e-prop*, and it is the main driver behind the resulting qualitative jump in computing capabilities of RSNNs which are attainable through biologically plausible learning. Equations (1) and (2) yield the representation

$$\frac{dE}{dW_{ji}} = \sum_{t} L_j^t \ e_{ji}^t \tag{3}$$

of the loss gradient, where we refer to  $L_j^t \stackrel{\text{def}}{=} \frac{dE}{dz_j^t}$  as the learning signal for neuron j. This equation defines a clear program for approximating the network loss gradient through local rules for synaptic plasticity: Change each weight  $W_{ji}$  at step t proportionally to  $-L_j^t e_{ji}^t$ , or accumulate these "tags" in a hidden variable that is translated occasionally into an actual weight change. Hence *e-prop* is an online learning method in a strict sense (see Fig. 1d and Movie S3). In particular, there is no need to unroll the network as for *BPTT*.

Since the ideal value  $\frac{dE}{dz_j^t}$  of the learning signal  $L_j^t$  also captures influences which the current spike output  $z_j^t$  of neuron j may have on E via future spikes of other neurons, its precise value is in general not available at time t. We replace it by an approximation, such as  $\frac{\partial E}{\partial z_j^t}$ , which ignores these indirect influences. This approximation takes only currently arising losses at the output neurons k of the RSNN into account, and routes them with neuron-specific weights  $B_{jk}$  to the network neurons j (see Fig. 2a):

$$L_j^t = \sum_k B_{jk} \underbrace{(y_k^t - y_k^{*,t})}_{\text{deviation of output } k} .$$
(4)

Although this approximate learning signal  $L_j^t$  only captures errors that arise at the current time step t, it is combined in equation (3) with an eligibility trace  $e_{ji}^t$  that may reach far back into the past of neuron j (see Fig. 3b), thereby alleviating the need to solve the temporal credit assignment problem by propagating signals backwards in time (like in *BPTT*).

There are several strategies for choosing the weights  $B_{jk}$  for this online learning signal. 166 In symmetric e-prop we set it equal to the corresponding weight  $W_{kj}^{\text{out}}$  of the synaptic 167 connection from neuron j to output neuron k, as demanded by  $\frac{\partial E}{\partial z_i^t}$ . Note that this learning 168 signal would actually implement  $\frac{dE}{dz_{i}^{t}}$  exactly in the absence of recurrent connections in the 169 network. Biologically more plausible are two variants of *e-prop* that avoid weight sharing: 170 In random e-prop the values of all weights  $B_{ik}$  – even for neurons j that are not synaptically 171 connected to output neuron k – are randomly chosen and remain fixed, similar to Broadcast 172 Alignment for feedforward networks [18, 19, 20]. In adaptive e-prop we let in addition  $B_{ik}$ 173 for neurons j that are synaptically connected to output neuron k evolve through a simple 174 local plasticity rule that mirrors the plasticity rule applied to  $W_{kj}^{\text{out}}$  (see section S2.3). 175

Resulting synaptic plasticity rules (see Methods) look similar to previously proposed plasticity rules [9] for the special case of LIF neurons without slowly changing hidden variables. In particular they involve postsynaptic depolarization as one of the factors, similarly as the data-based Clopath-rule in [21], see section S6.4 in the supplement for an analysis.

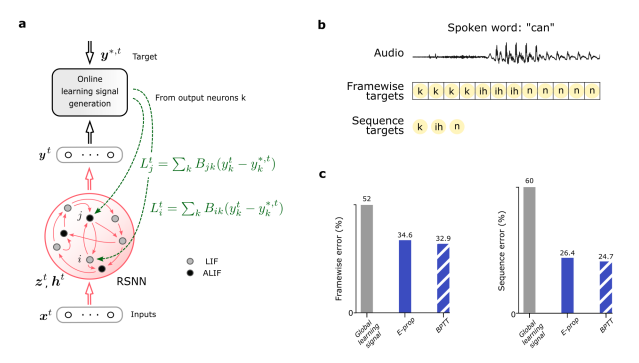


Figure 2: Comparison of the performance of *BPTT* and *e-prop* for learning phoneme recognition (TIMIT data set). a) Network architecture for *e-prop*, illustrated for an LSNN consisting of LIF and ALIF neurons. b) Input and target output for the two versions of TIMIT. c) Performance of *BPTT* and *symmetric e-prop* for LSNNs consisting of 800 neurons for framewise targets and 2400 for sequence targets (*random* and *adaptive e-prop* produced similar results, see Fig. S2). To obtain the "Global learning signal" baselines, the neuron specific feedbacks are replaced with global ones.

## <sup>181</sup> Comparing the performance of e-prop and BPTT for learning <sup>182</sup> spoken phoneme recognition

The phoneme recognition task TIMIT [22] is one of the most commonly used benchmarks 183 for temporal processing capabilities of different types of recurrent neural networks and 184 different learning approaches [23]. It comes in two versions. Both use, as input, acoustic 185 speech signals from sentences that are spoken by 630 speakers from 8 dialect regions of the 186 USA (see the top of Fig. 2b for a sample segment). In the simpler version, used for example 187 in [23], the goal is to recognize which of 61 phonemes is spoken in each 10 ms time frame 188 ("frame-wise classification"). In the more sophisticated version from [24], which achieved 189 an essential step toward human-level performance in speech-to-text transcription, the goal 190 is to recognize the sequence of phonemes in the entire spoken sentence independently 191 of their timing ("sequence transcription"). RSNNs consisting of LIF neurons do not even 192 reach with BPTT good performance for TIMIT [3]. Hence we are considering here LSNNs, 193 where a random subset of the neurons is a variation of the LIF model with firing rate 194 adaptation (ALIF neurons), see Methods. The name LSNN is motivated by the fact that 195

this special case of the RSNN model can achieve through training with *BPTT* similar performance as an LSTM network [3].

E-prop approximates the performance of BPTT on LSNNs for both versions of TIMIT 198 very well, as shown in Fig. 2c. Furthermore LSNNs could solve the frame-wise classification 199 task without any neuron firing more frequently than 12 Hz (spike count taken over 32 200 spoken sentences), demonstrating that they operate in an energy efficient spike-coding – 201 rather than a rate-coding – regime. For the more difficult version of TIMIT we trained as 202 in [24] a complex LSNN consisting of a feedforward sequence of three recurrent networks. 203 Our results show that *e-prop* can also handle learning for such more complex network 204 structures very well. In Fig. S4 we show for comparison also the performance of *e-prop* 205 and *BPTT* for LSTM networks on the same tasks. These data show that for both versions 206 of TIMIT the performance of e-prop for LSNNs comes rather close to that of BPTT for 207 LSTM networks. In addition, they show that *e-prop* provides also for LSTM networks a 208 functionally powerful online learning method. 200

### *E-prop* performance for learning a task where temporal credit assignment is difficult

A hallmark of cognitive computations in the brain is the capability to go beyond a purely 212 reactive mode: to integrate diverse sensory cues over time, and to wait until the right 213 moment arrives for an action. A large number of experiments in neuroscience analyze 214 neural coding after learning such tasks (see e.g. [25, 11]). But it had remained unknown 215 how one can model the learning of such cognitive computations in RSNNs of the brain. In 216 order to test whether *e-prop* can solve this problem, we considered the same task that was 217 studied in the experiments of [25] and [11]. There a rodent moved along a linear track in 218 a virtual environment, where it encountered several visual cues on the left and right, see 219 Fig. 3a and Movie S1. Later, when it arrived at a T-junction, it had to decide whether to 220 turn left or right. It was rewarded when it turned to that side from which it had previously 221 received the majority of visual cues. This task is not easy to learn since the subject needs 222 to find out that it does not matter on which side the last cue was, or in which order the 223 cues were presented. Instead, the subject has to learn to count cues separately for each 224 side and to compare the two resulting numbers. Furthermore the cues need to be processed 225 properly long before a reward is given. We show in Fig. S5 that LSNNs can learn this task 226 via *e-prop* in exactly the same way just from rewards. But since the way how *e-prop* solves 227 the underlying temporal credit assignment problem is easier to explain for the supervised 228 learning version of this task, we discuss here the case where a teacher tells the subject at 229 the end of each trial what would have been the right decision. This still yields a challenging 230 scenario for any online learning method since non-zero learning signals  $L_i^t$  arise only during 231 the last 150 ms of a trial (Fig. 3b). Hence all synaptic plasticity has to take place during 232 these last 150 ms, long after the input cues have been processed. Nevertheless, *e-prop* is 233 able to solve this learning problem, see Fig. 3c and Movie S3. It just needs a bit more time 234 to reach the same performance level as offline learning via BPTT (see Movie S2). Whereas 235

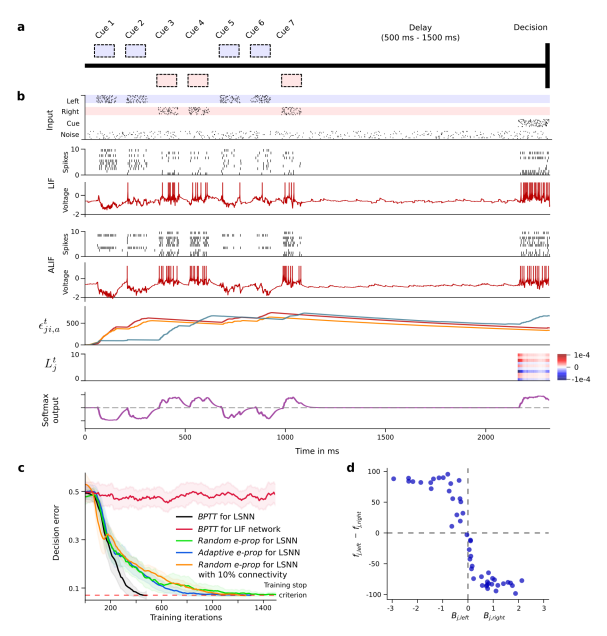


Figure 3: Solving a task with difficult temporal credit assignment. a) Setup of corresponding rodent experiments of [25] and [11], see Movie S1. b) Input spikes, spiking activity of 10 out of 50 sample LIF neurons and 10 out of 50 sample ALIF neurons, membrane potentials (more precisely:  $v_j^t - A_j^t$ ) for two sample neurons j, 3 samples of slow components of eligibility traces, sample learning signals for 10 neurons and softmax network output. c) Learning curves for *BPTT* and two *e-prop* versions applied to LSNNs, and *BPTT* applied to an RSNN without adapting neurons (red curve). Orange curve shows learning performance of *e-prop* for a sparsely connected LSNN consisting of excitatory and inhibitory neurons (Dale's law obeyed). The shaded areas are the 95%-confidence intervals of the mean accuracy computed with 20 runs. d) Correlation between the randomly drawn broadcast weights  $B_{jk}$  for k = left/right for learning signals in *random e-prop* and resulting sensitivity to "left" and "right" input components after learning.  $f_{j,\text{left}}$  ( $f_{j,\text{right}}$ ) was the resulting average firing rate of neuron j during presentation of left (right) cues after learning.

this task can not even be solved by *BPTT* with a regular RSNN that has no adapting neurons (red curve in Fig. 3c), all 3 previously discussed variations of *e-prop* can solve it if the RSNN contains adapting neurons. We explain in section S2.5 how this task can also be solved by sparsely connected LSNNs consisting of excitatory and inhibitory neurons: by integrating stochastic rewiring [26] into *e-prop*.

But how can the neurons in the LSNN learn to record and count the input cues if all the learning signals are identically 0 until the last 150 ms of a 2250 ms long trial (see 2nd to last row of Fig. 3b)? For answering this question one should note that firing of a neuron j at time t can affect the loss function E at a later time point t' > t in two different ways:

i) By affecting future values of slow hidden variables of neuron j (e.g., its firing threshold), which may then affect the firing of neuron j at t', which in turn may directly affect the loss function at time t'.

<sup>248</sup> ii) By affecting the firing of other neurons j' at t', which directly affects the loss function <sup>249</sup> at time t'.

In symmetric and adaptive e-prop one uses the partial derivative  $\frac{\partial E}{\partial z_j^t}$  as learning signal  $L_j^t$  for e-prop – instead of the online not available total derivative  $\frac{dE}{dz_j^t}$ . This blocks the 250 251 flow of gradient information along route ii. But the eligibility trace keeps the flow along 252 route i open. Therefore even symmetric and adaptive e-prop can solve the temporal credit 253 assignment problem of Fig. 3 through online learning: The gradient information that flows 254 along route i enables neurons to learn how to process the sensory cues at time points t 255 during the first 1050 ms, although this can affect the loss only at time points t' > 2100 ms 256 when the loss becomes non-zero. This is illustrated in the 3rd last row of Fig. 3b: The 257 slow component  $\epsilon_{ji,a}^t$  of the eligibility traces  $e_{ji}$  of adapting neurons j decays with the 258 typical long time constant of firing rate adaptation (see equation (24) and Movie S3). 259 Since these traces stretch from the beginning of the trial into its last phase, they enable 260 learning of differential responses to "left" and "right" input cues that arrived over 1050 ms 261 before any learning signals become non-zero, as shown in the 2nd to last row of Fig. 3b. 262 Hence eligibility traces provide "highways into the future" for the propagation of gradient 263 information. These can be seen as biologically realistic replacements for the highways into 264 the past that *BPTT* employs during its backwards pass (see Movie S2). 265

This analysis also tells us when symmetric e-prop is likely to fail to approximate the 266 performance of BPTT: If forward propagation of gradient information cannot reach along 267 route i those later time points t' when the value of the loss function becomes salient. One 268 can artificially induce this in the experiment of Fig. 3 by adding to the LSNN – which has 269 the standard architecture shown in Fig. 2a – hidden layers of a feedforward SNN through 270 which the communication between the LSNN and the readout neurons has to flow. The 271 neurons j' of these hidden layers block route i, while leaving route ii open. Hence the task 272 of Fig. 3 can still be learnt with this modified network architecture by BPTT, but not by 273 symmetric e-prop, see Fig. S8. 274

Identifying tasks where the performance of *random e-prop* stays far behind that of BPTT is more difficult, since error signals are sent there also to neurons that have no direct

connections to readout neurons. For deep feedforward networks it has been shown in [27] 277 that Broadcast Alignment, as defined in [20, 19], cannot reach the performance of Backprop 278 for difficult image classification tasks. Hence we expect that random e-prop will exhibit 279 corresponding deficiencies for difficult classification tasks with deep feedforward SNNs. 280 We are not aware of corresponding demonstrations of failures of Broadcast Alignment for 281 artificial RNNs, although they are likely to exist. Once they are found, they will probably 282 point to tasks where *random e-prop* fails for RSNNs. Currently we are not aware of any. 283 Fig. 3d provides insight into the functional role of the randomly drawn broadcast 284

weights in *random e-prop*: The difference of these weights determines for each neuron *j* whether it learns to respond in the first phase of a trial more to cues from the left or right. This observation suggests that neuron-specific learning signals for RSNNs have the advantage that they can create a diversity of feature detectors for task-relevant network inputs. Hence a suitable weighted sum of these feature detectors is later able to cancel remaining errors at the network output, similarly as in the case of feedforward networks [18].

We would like to point out that the use of the familiar actor-critic method in *reward*based *e-prop*, which we will discuss in the next section, provides an additional channel by which information about future losses can gate synaptic plasticity of the *e-prop* learner at the current time step t: Through the estimate V(t) of the value of the current state, that is simultaneously learnt via internally generated reward-prediction errors.

### 297 Reward-based e-prop

Deep RL has significantly advanced the state of the art in machine learning and AI through 298 clever applications of *BPTT* to RL [14]. We found that one of the arguably most powerful 299 RL methods within the range of deep RL approaches that are not directly biologically 300 implausible, policy gradient in combination with actor-critic, can be implemented with 301 *e-prop.* This yields the biologically plausible and hardware friendly deep RL algorithm 302 reward-based e-prop. The LSNN learns here both an approximation to the value function 303 (the "critic") and a stochastic policy (the "actor"). Neuron-specific learning signals are 304 combined in *reward-based e-prop* with a global signal that transmits reward prediction 305 errors (Fig. 4b). In contrast to the supervised case, where the learning signals  $L_j^t$  depend 306 on the deviation from an external target signal, the learning signals communicate here how 307 a stochastically chosen action deviates from the action mean that is currently proposed by 308 the network. 309

The resulting online synaptic plasticity rule (5) for deep RL is similar to equation (3), except that a fading memory filter  $\mathcal{F}_{\gamma}$  is applied here to the term  $L_j^t \bar{e}_{ji}^t$ , where  $\gamma$  is the given discount factor for future rewards and  $\bar{e}_{ji}^t$  denotes a low-pass filtered copy of the eligibility trace  $e_{ji}^t$  (see Methods). This term is multiplied in the synaptic plasticity rule with the reward prediction error  $\delta^t = r^t + \gamma V^{t+1} - V^t$ , where  $r^t$  is the reward received at time t. This yields an instantaneous weight change of the form:

$$\Delta W_{ji}^t = -\eta \ \delta^t \mathcal{F}_{\gamma} \left( L_j^t \overline{e}_{ji}^t \right) \,. \tag{5}$$

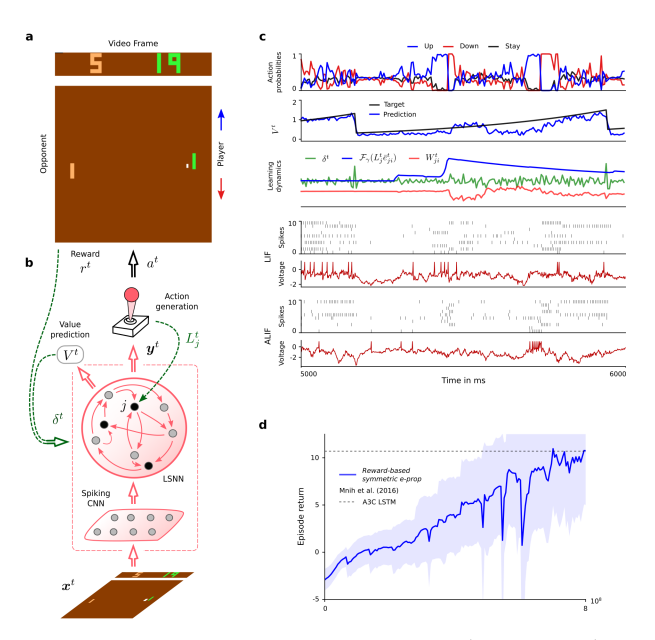


Figure 4: Application of *e-prop* to the Atari game Pong. a) Here the player (green paddle) has to outplay the opponent (light brown). A reward is acquired when the opponent cannot bounce back the ball (a small white square). To achieve this, the agent has to learn to hit the ball also with the edges of his paddle, which causes a less predictable trajectory. b) The agent is realized by an LSNN. The pixels of the current video frame of the game are provided as input. During processing of the stream of video frames by the LSNN, actions are generated by the stochastic policy in an online manner. At the same time, future rewards are predicted. The current error in prediction is fed back both to the LSNN and the spiking CNN that preprocesses the frames. (caption continued on next page)

Figure 4: (continued caption) c) Sample trial of the LSNN after learning with *reward-based e-prop*. From top to bottom: probabilities of stochastic actions, prediction of future rewards, learning dynamics of a random synapse (arbitrary units), spiking activity of 10 out of 240 sample LIF neurons and 10 out of 160 sample ALIF neurons, and membrane potentials (more precisely:  $v_j^t - A_j^t$ ) for the two sample neurons j at the bottom of the spike raster above. d) Learning progress of the LSNN trained with *reward-based e-prop*, reported as the sum of collected rewards during an episode. The learning curve is averaged over 5 different runs and the shaded area represents the standard deviation. More information about the comparison between our results and A3C are given in section S5.3.

Previous 3-factor learning rules for RL were usually of the form  $\Delta W^t = \eta \delta^t \bar{e}_{ii}^t$  [28, 9]. 316 Hence they estimated gradients of the policy just by correlating the output of network 317 neurons with the reward prediction error. The learning power of this approach is known to 318 be quite limited due to high noise in the resulting gradient estimates. In contrast, in the 319 plasticity rule (5) for *reward-based e-prop* the eligibility traces are first combined with a 320 neuron specific feedback  $L_i^t$ , before they are multiplied with the reward prediction error  $\delta^t$ . 321 We show in Methods analytically that this yields estimates of policy- and value gradients 322 similarly as in deep RL with *BPTT*. Furthermore, in contrast to previously proposed 3-323 factor learning rules, this rule (5) is also applicable to LSNNs. 324

We tested *reward-based e-prop* on a classical benchmark task [14] for learning intelligent 325 behavior from rewards: Winning Atari video games provided by the Arcade Learning 326 Environment [29]. To win such game, the agent needs to learn to extract salient information 327 from the pixels of the game screen, and to infer the value of specific actions, even if rewards 328 are obtained in a distant future. In fact, learning to win Atari games is a serious challenge 329 for reinforcement learning even in machine learning [14]. Besides artificial neural networks 330 and BPTT, previous solutions also required experience replay (with a perfect memory of 331 many frames and action sequences that occurred much earlier) or an asynchronous training 332 of numerous parallel agents sharing synaptic weight updates. We show here that also an 333 LSNN can learn via *e-prop* to win Atari games, through online learning of a single agent. 334 This becomes possible with a single agent and without episode replay if the agent uses 335 a schedule of increasing episode lengths –with a learning rate that is inversely related to 336 that length. Using this scheme, an agent can experience diverse and uncorrelated short 337 episodes in the first phase of learning, producing useful skills. Subsequently, the agent can 338 fine-tune its policy using longer episodes. 339

First, we considered the well-known Atari game Pong (Fig. 4a). Here, the agent has to learn to hit a ball in a clever way using up and down movements of his paddle. A reward is obtained if the opponent cannot catch the ball. We trained an agent using *reward-based e-prop* for this task, and show a sample trial in Fig. 4c and Movie S5. In contrast to common deep RL solutions, the agent learns here in a strict online manner, receiving at any time just the current frame of the game screen. In Panel d of Fig. 4 we demonstrate that also this biologically realistic learning approach leads to a competitive score.

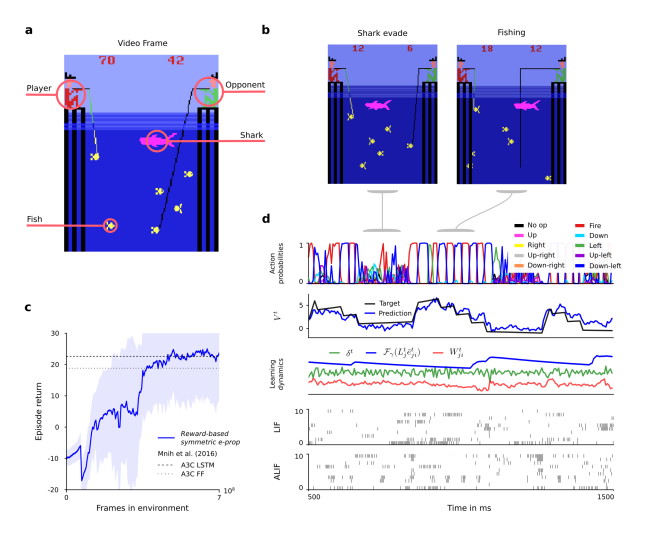


Figure 5: Application of *e-prop* to learning to win the Atari game Fishing Derby. a) Here the player has to compete against an opponent, and try to catch more fish from the sea. b) Once a fish has bit, the agent has to avoid that the fish gets touched by a shark. c) Sample trial of the trained network. From top to bottom: probabilities of stochastic actions, prediction of future rewards, learning dynamics of a random synapse (arbitrary units), spiking activity of 20 out of 180 sample LIF neurons and 20 out of 120 sample ALIF neurons. d) Learning curves of an LSNN trained with *reward-based e-prop* as in Fig. 4d.

If one does not insist on an online setting where the agent receives just the current 347 frame of the video screen but the last 4 frames, winning strategies for about half of the 348 Atari games can already be learnt by feedforward neural networks (see table S3 of [14]). 349 However, for other Atari games, such as Fishing Derby (Fig. 5a), it was even shown in [14] 350 that deep RL applied to LSTM networks achieves a substantially higher score than any deep 351 RL method for feedforward networks which was considered there. Hence, in order to test 352 the power of online *reward-based e-prop* also for those Atari games that require enhanced 353 temporal processing, we tested it on the Fishing Derby game. In this game, the agent has 354 to catch as many fish as possible while avoiding that the shark touches the fish with any 355 part of its body, and that the opponent catches the fish first. We show in Fig. 5c that 356 online reward-based e-prop applied to an LSNN does in fact reach the same performance 357 as reference offline algorithms applied to LSTM networks. We show a random trial after 358 learning in Fig. 5d, where we can identify two different learnt behaviors: 1) evading the 359 shark, 2) collecting fish. The agent has learnt to switch between these two behaviors as 360 required by the situation. 361

In general, we conjecture that variants of *reward-based e-prop* will be able to solve most deep RL tasks that can be solved by online actor-critic methods in machine learning.

## 364 Discussion

We propose that in order to understand the computational function and neural coding of 365 neural networks in the brain, one needs to understand the organization of the plasticity 366 mechanisms that install and maintain these. So far BPTT was the only candidate for 367 that, since no other learning method provided sufficiently powerful computational function 368 to RSNN models. But since BPTT is not viewed to be biologically realistic [5], it does 369 not help us to understand learning in the brain. *E-prop* offers a solution to this dilemma. 370 since it does not require biologically unrealistic mechanisms, but still enables RSNNs to 371 learn difficult computational tasks, in fact almost as well as *BPTT*. Furthermore it enables 372 RSNNs to solve these tasks in an energy efficient sparse firing regime, rather than resorting 373 to rate coding. 374

*E-prop* relies on two types of signals that are abundandly available in the brain, but 375 whose precise role for learning have not yet been understood: eligibility traces and learning 376 signals. Since e-prop is based on a transparent mathematical principle (see equation (3)), 377 it provides a normative model for both types of signals, as well as for synaptic plasticity 378 rules. Interestingly, the resulting learning model suggests that a characteristic aspect of 379 many biological neurons – the presence of slowly changing hidden variables – provides a 380 possible solution to the problem how a RSNN can learn without error signals that propagate 381 backwards in time: Slowly changing hidden variables of neurons cause eligibility traces 382 that propagate forward over longer time spans, and are therefore able to coincide with 383 later arising instantaneous error signals (see Fig. 3b). 384

The theory of *e-prop* makes a concrete experimentally testable prediction: that the time constant of the eligibility trace for a synapse is correlated with the time constant for the history-dependence of the firing activity of the postsynaptic neuron. It also suggests that the experimentally found diverse time constants of the firing activity of populations of neurons in different brain areas [30] are correlated with their capability to handle corresponding ranges of delays in temporal credit assignment for learning.

Finally, *e-prop* theory provides a hypothesis for the functional role of the experimentally 391 found diversity of dopamine signals to different populations of neurons [11]. Whereas 392 previous theories of reward-based learning required that the same learning signal is sent 393 to all neurons, the basic equation (1) for *e-prop* postulates that ideal top-down learning 394 signals to a population of neurons depend on its impact on the network performance (loss 395 function), and should therefore be target specific (see Fig. 2c and section S6.2). In fact, the 396 learning-to-learn result for e-prop in [31] suggests that prior knowledge about the possible 397 range of learning tasks for a brain area could optimize top-down learning signals even 398 further on an evolutionary time scale, thereby enabling for example learning from few or 399 even a single trial. 400

Previous methods for training RSNNs did not aim at approximating BPTT. Instead 401 some of them were relying on control theory to train a chaotic reservoir of spiking neurons 402 [32, 33, 34]. Others used the FORCE algorithm [35, 36] or variants of it [37, 38, 39, 403 35]. However the FORCE algorithm was not argued to be biologically realistic, since the 404 plasticity rule for each synaptic weight requires knowledge of the current values of all other 405 synaptic weights. The generic task considered in [35] was to learn with supervision how 406 to generate patterns. We show in Figs. S1, S7, and Movie S4 that RSNNs can learn such 407 tasks also with a biologically plausible learning method: *e-prop*. 408

Several methods for approximating stochastic gradient descent in *feedforward* networks 409 of spiking neurons have been proposed, see e.g. [40, 41, 42, 43, 44]. These employ – 410 like e-prop – a pseudo-gradient to overcome the non-differentiability of a spiking neuron. 411 as proposed previously in [45, 46]. [40, 42, 43] arrive at a synaptic plasticity rule for 412 feedforward networks that consists - like e-prop - of the product of a learning signal and 413 a derivative (eligibility trace) that describes the dependence of a spike of a neuron j on 414 the weight of an afferent synapse  $W_{ii}$ . But in a recurrent network the spike output of j 415 depends on  $W_{ii}$  also indirectly, via loops in the network that allow that a spike of neuron 416 *i* contributes to the firing of other neurons, which in turn affect firing of the presynaptic 417 neuron *i*. Hence the corresponding eligibility trace can no longer be locally computed if 418 one transfers these methods for feedforward networks to recurrently connected networks. 419 Therefore [40] suggests the need to investigate extensions of their approach to RSNNs. 420

Previous work on the design of online gradient descent learning algorithms for *non*-421 spiking RNNs was based on real-time recurrent learning (RTRL) [15]. RTRL itself has 422 rarely been used since its computational complexity per time-step is  $\mathcal{O}(n^4)$ , if n is the 423 number of neurons. But interesting approximations to RTRL have subsequently been pro-424 posed (see [47] for a review): some stochastic approximations [48] which are  $\mathcal{O}(n^3)$  or only 425 applicable for small networks [49], and also recently two deterministic  $\mathcal{O}(n^2)$  approxima-426 tions [50, 51]. The latter were in fact written at the same time as the first publication 427 of e-prop [31]. A structural difference between this paper and [50] is that their approach 428 requires that learning signals are transmitted between the neurons in the RNN, with sepa-429

rately learnt weights. [51] derived for rate based neurons a learning rule similar to *random e-prop*. But this work did not address other forms of learning than supervised regression,
such as RL, nor learning in networks of spiking neurons, or in more powerful types of RNNs
with slow hidden variables such as LSTM networks or LSNNs.

*E-prop* also has complexity  $\mathcal{O}(n^2)$ , in fact  $\mathcal{O}(S)$  if S is the number of synaptic connec-434 tions. This bound is optimal except for the constant factor- since this is the asymptotic 435 complexity of just simulating the RNN. The key point of e-prop is that the general form (13) 436 of its eligibility trace collects all contributions to the loss gradient that can be locally com-437 puted in a feedforward manner. This general form enables applications to spiking neurons 438 with slowly varying hidden variables, such as neurons with firing rate adaptation, which 439 are essential ingredients of RSNNs to reach the computational power of LSTM networks 440 [3]. We believe that this approach can be extended in future work –with a suitable choice 441 of pseudo-derivatives—to a wide range of biologically more realistic neuron models. It also 442 enables the combination of these rigorously derived eligibility traces with – semantically 443 identical but algorithmically very different – eligibility traces from RL for reward-based 444 *e-prop* (equation (5)), thereby bringing the power of deep RL to RSNNs. As a result, we 445 were able to show in Fig. 2 - 5 that RSNNs can learn with the biologically plausible rules 446 for synaptic plasticity that arise from the *e-prop* theory to solve tasks such as phoneme 447 recognition, integrating evidence over time and waiting for the right moment to act, and 448 winning Atari games. These are tasks that are fundamental for modern learning-based AI, 449 but have so far not been solved with RSNNs. Hence *e-prop* provides a new perspective of 450 the major open question how intelligent behavior can be learnt and controlled by neural 451 networks of the brain. 452

Apart from obvious consequences of e-prop for research in neuroscience and cognitive 453 science, *e-prop* also provides an interesting new tool for approaches in machine learning 454 where *BPTT* is replaced by approximations in order to improve computational efficiency. 455 We have already shown in Fig. S4 that e-prop provides a powerful online learning method 456 for LSTM networks. Furthermore, the combination of eligibility traces from *e-prop* with 457 synthetic gradients from [52] even improves performance of LSTM networks for difficult 458 machine learning problems such as the copy-repeat task and the Penn Treebank word 459 prediction task [31]. Other future extensions of *e-prop* could explore a combination with 460 attention-based models in order to cover multiple timescales. 461

Finally, *e-prop* suggests a promising new approach for realizing powerful on-chip learning of RSNNs on neuromorphic chips. Whereas *BPTT* is not within the reach of current neuromorphic hardware, an implementation of *e-prop* appears to offer no serious hurdle. Our results show that an implementation of *e-prop* will provide a qualitative jump in on-chip learning capabilities of neuromorphic hardware.

## $_{467}$ Methods

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#### 476 Network models

To exhibit the generality of the *e-prop* approach, we define the dynamics of recurrent neural 477 networks using a general formalism that is applicable to many recurrent neural network 478 models, not only to RSNNs and LSNNs. Also non-spiking models such as LSTM networks 479 fit under this formalism (see section S4.3 in the supplement). The network dynamics is 480 summarized by the computational graph in Fig. 6. It uses the function M to define the 481 update of the hidden state of a neuron j:  $\mathbf{h}_{j}^{t} = M(\mathbf{h}_{j}^{t-1}, \mathbf{z}^{t-1}, \mathbf{x}^{t}, \mathbf{W}_{j})$ , where  $\mathbf{W}_{j}$  gathers 482 the weights of synapses arriving at neuron j, and f to define the update of the observable 483 state of a neuron j:  $z_j^t = f(\mathbf{h}_j^t, \mathbf{z}^{t-1}, \mathbf{x}^t, \mathbf{W}_j)$  (f simplifies to  $z_j^t = f(\mathbf{h}_j^t)$  for LIF and ALIF 484 neurons). We chose a discrete time step of  $\delta t = 1$  ms for all our simulations. Control 485 experiments with smaller time steps for the task of Fig. 3, reported in Fig. S6, suggest that 486 the size of the time step has no significant impact on the performance of *e-prop*. 487

**LIF neurons.** Each LIF neuron has a one dimensional internal state – or hidden variable  $h_{489}^{t} - h_j^t$  that consists only of the membrane potential  $v_j^t$ . The observable state  $z_j^t \in \{0, 1\}$  is binary, indicating a spike  $(z_j^t = 1)$  or no spike  $(z_j^t = 0)$  at time t. The dynamics of the LIF model is defined by the equations:

$$v_{j}^{t+1} = \alpha v_{j}^{t} + \sum_{i \neq j} W_{ji}^{\text{rec}} z_{i}^{t} + \sum_{i} W_{ji}^{\text{in}} x_{i}^{t+1} - z_{j}^{t} v_{\text{th}}$$
(6)

$$z_j^t = H\left(v_j^t - v_{\rm th}\right). \tag{7}$$

 $W_{ji}^{\text{rec}}(W_{ji}^{\text{in}})$  is the synaptic weight from network (input) neuron *i* to neuron *j*. The decay factor  $\alpha$  in (6) is given by  $e^{-\delta t/\tau_m}$ , where  $\tau_m$  (typically 20 ms) is the membrane time constant.  $\delta t$  denotes the discrete time step size, which is set to 1 ms in our simulations. *H* denotes the Heaviside step function. Note that we deleted in equation (6) the factor  $1 - \alpha$ that occured in the corresponding equation (4) in the supplement of [3]. This simplifies the notation in our derivations, and has no impact on the model if parameters like the threshold voltage are scaled accordingly. <sup>499</sup> Due to the term  $-z_j^t v_{\text{th}}$  in equation (6), the neurons membrane potential is reduced <sup>500</sup> by a constant value after an output spike, which relates our model to the spike response <sup>501</sup> model [53]. To introduce a simple model of neuronal refractoriness, we further assume <sup>502</sup> that  $z_j^t$  is fixed to 0 after each spike of neuron j for a short refractory period of 2 to 5 ms <sup>503</sup> depending on the simulation.

LSNNs. According to the database of the Allen Institute [2] a fraction of neurons be-504 tween roughly 20 % (in mouse visual cortex) and 40 % (in the human frontal lobe) exhibit 505 spike frequency adaptation (SFA). It had been shown in [3] that the inclusion of neuron 506 models with SFA – via a time-varying firing threshold as slow hidden variable – drastically 507 enhances computing capabilities of RSNN models. Hence we consider here the same simple 508 model for neurons with SFA as in [3], to which we refer as adaptive LIF (ALIF) neuron. 509 This model is basically the same as the  $GLIF_2$  model in the Technical White paper on 510 generalized LIF (GLIF) models from [2]. LSNNs are recurrently connected networks that 511 consist of LIF and ALIF neurons. ALIF neurons j have a second hidden variable  $a_i^t$ , which 512 denotes the variable component of its firing threshold. As a result, their internal state is 513 a 2 dimensional vector  $\mathbf{h}_{j}^{t} \stackrel{\text{def}}{=} [v_{j}^{t}, a_{j}^{t}]$ . Their threshold potential  $A_{j}^{t}$  increases with every 514 output spike and decreases exponentially back to the baseline threshold  $v_{\rm th}$ . This can be 515 described by 516

$$A_i^t = v_{\rm th} + \beta a_i^t , \qquad (8)$$

$$z_j^t = H(v_j^t - A_j^t) , \qquad (9)$$

517 with a threshold adaptation according to

$$a_j^{t+1} = \rho a_j^t + z_j^t , (10)$$

where the decay factor  $\rho$  is given by  $e^{-\delta t/\tau_a}$ , and  $\tau_a$  is the adaptation time constant that is 518 typically chosen to be in the range of the time span of the length of the working memory 519 that is a relevant for a given task. This is a very simple model for a neuron with spike 520 frequency adaptation [3]. We refer to [53, 54, 55] for experimental data and other neuron 521 models. We refer to a recurrent network of spiking neurons (RSNN) as LSNN if some of 522 its neurons are adaptive. We chose a fraction between 25 and 40 % of the neurons to be 523 adapting, like in the data from neocortex [2], with time constants that are roughly on the 524 same time scale as the tasks for which the network is trained. 525

In relation to the more general formalism represented in the computational graph in Fig. 6, equations (6) and (10) define  $M(\mathbf{h}_{j}^{t-1}, \mathbf{z}^{t-1}, \mathbf{x}^{t}, \mathbf{W}_{j})$ , and equations (7) and (9) define  $f(\mathbf{h}_{j}^{t})$ .

Gradient descent for RSNNs. Gradient descent is problematic for spiking neurons because of the step function H in equation (7). We overcome this issue as in [56, 3]: The non-existing derivative  $\frac{\partial z_j^t}{\partial v_j^t}$  is replaced in simulations by a simple nonlinear function of the membrane potential that is called the pseudo-derivative. Outside of the refractory period, we choose a pseudo-derivative of the form  $\psi_j^t = \frac{1}{v_{\text{th}}} \gamma_{\text{pd}} \max\left(0, 1 - \left|\frac{v_j^t - A_j^t}{v_{\text{th}}}\right|\right)$  where  $\gamma_{\text{pd}} = 0.3$ for ALIF neurons, and for LIF neurons  $A_j^t$  is replaced by  $v_{\text{th}}$ . During the refractory period the pseudo derivative is set to 0.

Network output and loss functions. We assume that network outputs  $y_k^t$  are realvalued and produced by leaky output neurons (readouts) k, which are not recurrently connected:

$$y_k^t = \kappa y_k^{t-1} + \sum_j W_{kj}^{\text{out}} z_j^t + b_k^{\text{out}}, \qquad (11)$$

where  $\kappa \in [0, 1]$  defines the leak and  $b_k^{\text{out}}$  denotes the output bias. The leak factor  $\kappa$  is given for spiking neurons by  $e^{-\delta t/\tau_{out}}$ , where  $\tau_{out}$  is the membrane time constant. Note that for non-spiking neural networks (such as for LSTM networks), temporal smoothing of the network observable state is not necessary. In this case, one can use  $\kappa = 0$ .

The loss function  $E(\mathbf{z}^1, \ldots, \mathbf{z}^T)$  quantifies the network performance. We assume that it depends only on the observable states  $\mathbf{z}^1, \ldots, \mathbf{z}^T$  of the network neurons. For instance, for a regression problem we define E as the mean square error  $E = \frac{1}{2} \sum_{t,k} (y_k^t - y_k^{*,t})^2$ between the network outputs  $y_k^t$  and target values  $y_k^{*,t}$ . For classification or RL tasks the loss function E has to be re-defined accordingly.

#### 548 Conventions

Notation for derivatives. We distinguish the total derivative  $\frac{dE}{d\mathbf{z}^t}(\mathbf{z}^1, \dots, \mathbf{z}^T)$ , which takes into account how E depends on  $\mathbf{z}^t$  also indirectly through influence of  $\mathbf{z}^t$  on the other variables  $\mathbf{z}^{t+1}, \dots, \mathbf{z}^T$ , and the partial derivative  $\frac{\partial E}{\partial \mathbf{z}^t}(\mathbf{z}^1, \dots, \mathbf{z}^T)$  which quantifies only the direct dependence of E on  $\mathbf{z}^t$ .

Analogously for the hidden state  $\mathbf{h}_{j}^{t} = M(\mathbf{h}_{j}^{t-1}, \mathbf{z}^{t-1}, \mathbf{x}^{t}, \mathbf{W}_{j})$  the partial derivative  $\frac{\partial M}{\partial \mathbf{h}_{j}^{t-1}}$  denotes the partial derivative of M with respect to  $\mathbf{h}_{j}^{t-1}$ . It only quantifies the direct influence of  $\mathbf{h}_{j}^{t-1}$  on  $\mathbf{h}_{j}^{t}$  and it does not take into account how  $\mathbf{h}_{j}^{t-1}$  indirectly influences  $\mathbf{h}_{j}^{t}$  via the observable states  $\mathbf{z}^{t-1}$ . To improve readability we also use the following abbreviations:  $\frac{\partial \mathbf{h}_{j}^{t}}{\partial \mathbf{h}_{j}^{t-1}} \stackrel{\text{def}}{=} \frac{\partial M}{\partial \mathbf{h}_{j}^{t-1}} (\mathbf{h}_{j}^{t-1}, \mathbf{z}^{t-1}, \mathbf{x}^{t}, \mathbf{W}_{j}), \quad \frac{\partial \mathbf{h}_{j}^{t}}{\partial W_{ji}} \stackrel{\text{def}}{=} \frac{\partial M}{\partial W_{ji}} (\mathbf{h}_{j}^{t-1}, \mathbf{z}^{t-1}, \mathbf{x}^{t}, \mathbf{W}_{j}), \text{ and}$  $\frac{\partial z_{j}^{t}}{\partial \mathbf{h}_{j}^{t}} \stackrel{\text{def}}{=} \frac{\partial f}{\partial h^{t}} (\mathbf{h}_{j}^{t}, \mathbf{z}^{t-1}, \mathbf{x}^{t}, \mathbf{W}_{j}).$ 

Notation for temporal filters. For ease of notation we use the operator  $\mathcal{F}_{\alpha}$  to denote the low-pass filter such that, for any time series  $x_t$ :

$$\mathcal{F}_{\alpha}(x^{t}) = \alpha \mathcal{F}_{\alpha}(x^{t-1}) + x^{t} , \qquad (12)$$

and  $\mathcal{F}_{\alpha}(x^0) = x^0$ . In the specific case of the time series  $z_j^t$  and  $e_{ji}^t$ , we simplify notation further and write  $\overline{z}_j^t$  and  $\overline{e}_{ji}^t$  for  $\mathcal{F}_{\alpha}(z_j)^t$  and  $\mathcal{F}_{\kappa}(e_{ji})^t$ .

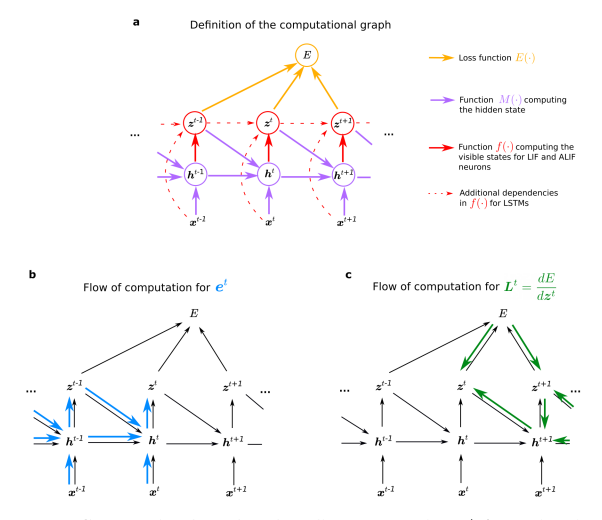


Figure 6: Computational graph and gradient propagations a) Assumed mathematical dependencies between hidden neuron states  $\mathbf{h}_{j}^{t}$ , neuron outputs  $\mathbf{z}^{t}$ , network inputs  $\mathbf{x}^{t}$ , and the loss function E through the mathematical functions  $E(\cdot)$ ,  $M(\cdot)$ ,  $f(\cdot)$  are represented by coloured arrows. **b-c**) The flow of computation for the two components  $\mathbf{e}^{t}$  and  $\mathbf{L}^{t}$  that merge into the loss gradients of equation (3) can be represented in similar graphs. **b**) Following equation (14), the flow of the computation of the eligibility traces  $e_{ji}^{t}$  is going forward in time. **c**) Instead the ideal learning signals  $L_{j}^{t} = \frac{dE}{dz_{j}^{t}}$  require to propagate gradients backward in time. Hence while  $e_{ji}^{t}$  is computed exactly,  $L_{j}^{t}$  is approximated in *e-prop* applications to yield an online learning algorithm.

#### <sup>563</sup> Mathematical basis for *e-prop*

We provide here the proof of the fundamental equation (1) for *e-prop* 

$$\frac{dE}{dW_{ji}} = \sum_{t} \frac{dE}{dz_j^t} \cdot \left[\frac{dz_j^t}{dW_{ji}}\right]_{\text{local}} ,$$

with the new eligibility trace

$$e_{ji}^{t} \stackrel{\text{def}}{=} \left[ \frac{dz_{j}^{t}}{dW_{ji}} \right]_{\text{local}} \stackrel{\text{def}}{=} \frac{\partial z_{j}^{t}}{\partial \mathbf{h}_{j}^{t}} \underbrace{\sum_{t' \leq t} \frac{\partial \mathbf{h}_{j}^{t}}{\partial \mathbf{h}_{j}^{t-1}} \cdots \frac{\partial \mathbf{h}_{j}^{t'+1}}{\partial \mathbf{h}_{j}^{t'}} \cdot \frac{\partial \mathbf{h}_{j}^{t'}}{\partial W_{ji}}}_{\stackrel{\text{def}}{=} \boldsymbol{\epsilon}_{ji}^{t}}$$
(13)

For spiking neurons j we replace the first factor  $\frac{\partial z_j^i}{\partial \mathbf{h}_j^t}$  of  $e_{ji}^t$  by the pseudo-derivative, see [3, 4, 56]. The second factor  $\boldsymbol{\epsilon}_{ji}^t$ , which we call eligibility vector, obviously satisfies the recursive equation

$$\boldsymbol{\epsilon}_{ji}^{t} = \frac{\partial \boldsymbol{h}_{j}^{t}}{\partial \boldsymbol{h}_{j}^{t-1}} \cdot \boldsymbol{\epsilon}_{ji}^{t-1} + \frac{\partial \mathbf{h}_{j}^{t}}{\partial W_{ji}} , \qquad (14)$$

where  $\cdot$  denotes the dot product. This provides the rule for the online computation of  $\boldsymbol{\epsilon}_{ji}^t$ , and hence of  $e_{ji}^t = \frac{\partial z_j^t}{\partial \mathbf{h}_i^t} \cdot \boldsymbol{\epsilon}_{ji}^t$ .

We start from a classical factorization of the loss gradients in recurrent neural networks that arises for instance in equation (12) of [57] to describe *BPTT*. This classical factorization can be justified by unrolling an RNN into a large feedforward network where each layer (*l*) represents one time step. In a feedforward network the loss gradients with respect to the weights  $W_{ji}^{(l)}$  of layer *l* are given by  $\frac{dE}{dW_{ji}^{(l)}} = \frac{dE}{dh_j^{(l)}} \frac{\partial h_j^{(l)}}{\partial W_{ji}^{(l)}}$ . But as the weights are shared across the layers when representing a recurrent network, the summation of these gradients over the layers *l* of the unrolled RNN yields this classical factorization of the loss gradients:

$$\frac{dE}{dW_{ji}} = \sum_{t'} \frac{dE}{d\mathbf{h}_{j}^{t'}} \cdot \frac{\partial \mathbf{h}_{j}^{t'}}{\partial W_{ji}} \,. \tag{15}$$

Note that the first factor  $\frac{dE}{dh_i^{t'}}$  in these products also needs to take into account how the 577 internal state  $\mathbf{h}_{i}$  of neuron j evolves during subsequent time steps, and whether it influences 578 firing of i at later time steps. This is especially relevant for ALIF neurons and other 579 biologically realistic neuron models with slowly changing internal states. Note that this 580 first factor of (15) is replaced in the *e-prop* equation (13) by the derivative  $\frac{dE}{dz_i^t}$  of E with 581 regard to the observable variable  $z_j^{t'}$ . There the evolution of the internal state of neuron j 582 is pushed into the second factor, the eligibility trace  $e_{ji}$ , which collects in *e-prop* all online 583 computable factors of the loss gradient that just involve neurons j and i. 584

Now we show that one can re-factorize the expression (15) and prove that the loss 585 gradients can also be computed using the new factorization (13) that underlies *e-prop*. In 586 the steps of the subsequent proof until equation (19), we decompose the term  $\frac{dE}{d\mathbf{h}_{i}^{t'}}$  into a 587 series of learning signals  $L_j^t = \frac{dE}{dz_j^t}$  and local factors  $\frac{\partial \mathbf{h}_j^{t+1}}{\partial \mathbf{h}_j^t}$  for  $t \ge t'$ . Those local factors 588 will later be used to transform the partial derivative  $\frac{\partial \mathbf{h}_{j}^{t'}}{\partial W_{ji}}$  from equation (15) into the 589 eligibility vector  $\boldsymbol{\epsilon}_{ji}^t$  that integrates the whole history of the synapse up to time t, not just a single time step. To do so, we express  $\frac{dE}{d\mathbf{h}_i^{t'}}$  recursively as a function of the same derivative 590 591 at the next time step  $\frac{dE}{d\mathbf{h}_{i}^{t'+1}}$  by applying the chain rule at the node  $\mathbf{h}_{j}^{t}$  for t = t' of the 592 computational graph shown in Fig. 6c: 593

$$\frac{dE}{d\mathbf{h}_{j}^{t'}} = \frac{dE}{dz_{j}^{t'}} \frac{\partial z_{j}^{t'}}{\partial \mathbf{h}_{j}^{t'}} + \frac{dE}{d\mathbf{h}_{j}^{t'+1}} \frac{\partial \mathbf{h}_{j}^{t'+1}}{\partial \mathbf{h}_{j}^{t'}}$$
(16)

$$= L_j^{t'} \frac{\partial z_j^{t'}}{\partial \mathbf{h}_j^{t'}} + \frac{dE}{d\mathbf{h}_j^{t'+1}} \frac{\partial \boldsymbol{h}_j^{t'+1}}{\partial \boldsymbol{h}_j^{t'}}, \qquad (17)$$

where we defined the learning signal  $L_j^{t'}$  as  $\frac{dE}{dz_j^{t'}}$ . The resulting recursive expansion ends at the last time step T of the computation of the RNN, i.e.,  $\frac{dE}{d\mathbf{h}_j^{T+1}} = 0$ . If one repeatedly substitutes the recursive formula (17) into the classical factorization (15) of the loss gradients, one gets:

$$\frac{dE}{dW_{ji}} = \sum_{t'} \left( L_j^{t'} \frac{\partial z_j^{t'}}{\partial \mathbf{h}_j^{t'}} + \frac{dE}{d\mathbf{h}_j^{t'+1}} \frac{\partial \boldsymbol{h}_j^{t'+1}}{\partial \boldsymbol{h}_j^{t'}} \right) \cdot \frac{\partial \mathbf{h}_j^{t'}}{\partial W_{ji}}$$
(18)

$$= \sum_{t'} \left( L_j^{t'} \frac{\partial z_j^{t'}}{\partial \mathbf{h}_j^{t'}} + \left( L_j^{t'+1} \frac{\partial z_j^{t'+1}}{\partial \mathbf{h}_j^{t'+1}} + (\cdots) \frac{\partial \mathbf{h}_j^{t'+2}}{\partial \mathbf{h}_j^{t'+1}} \right) \frac{\partial \mathbf{h}_j^{t'+1}}{\partial \mathbf{h}_j^{t'}} \right) \cdot \frac{\partial \mathbf{h}_j^{t'}}{\partial W_{ji}} .$$
(19)

The following equation is the main equation for understanding the transformation from 598 *BPTT* into *e-prop*. The key idea is to collect all terms  $\frac{\partial h_j^{t'+1}}{\partial h_i^{t'}}$  which are multiplied with 599 the learning signal  $L_j^t$  at a given time t. These are only terms that concern events in 600 the computation of neuron j up to time t, and they do not depend on other future losses 601 or variable values. To this end, we write the term in parentheses in equation (19) into 602 a second sum indexed by t and exchange the summation indices to pull out the learning 603 signal  $L_j^t$ . This expresses the loss gradient of E as a sum of learning signals  $L_j^t$  multiplied 604 by some factor indexed by ji, which we define as the eligibility trace  $e_{ji}^t \in \mathbb{R}$ . The main factor of it is the eligibility vector  $\boldsymbol{\epsilon}_{ji}^t \in \mathbb{R}^d$ , which has the same dimension as the hidden 605 606

607 state  $\mathbf{h}_{i}^{t}$ :

$$\frac{dE}{dW_{ji}} = \sum_{t'} \sum_{t \ge t'} L_j^t \frac{\partial z_j^t}{\partial \mathbf{h}_j^t} \frac{\partial \mathbf{h}_j^t}{\partial \mathbf{h}_j^{t-1}} \cdots \frac{\partial \mathbf{h}_j^{t'+1}}{\partial \mathbf{h}_j^{t'}} \cdot \frac{\partial \mathbf{h}_j^{t'}}{\partial W_{ji}}$$
(20)

$$= \sum_{t} L_{j}^{t} \frac{\partial z_{j}^{t}}{\partial \mathbf{h}_{j}^{t}} \underbrace{\sum_{t' \leq t} \frac{\partial \boldsymbol{h}_{j}^{t}}{\partial \boldsymbol{h}_{j}^{t-1}} \cdots \frac{\partial \boldsymbol{h}_{j}^{t'+1}}{\partial \boldsymbol{h}_{j}^{t'}} \cdot \frac{\partial \mathbf{h}_{j}^{t'}}{\partial W_{ji}}}_{\stackrel{\text{def}}{=} \boldsymbol{\epsilon}_{ji}^{t}} .$$
(21)

This completes the proof of equations (1), (3), (13).

#### <sup>609</sup> Derivation of eligibility traces for concrete neuron models

The eligibility traces for LSTMs are derived in the supplementary materials. Below we provide the derivation of eligibility traces for spiking neurons.

Eligibility traces for LIF neurons. We compute the eligibility trace of a synapse of a LIF neuron without adaptive threshold (equation (6)). Here the hidden state  $\mathbf{h}_{j}^{t}$  of a neuron consists just of the membrane potential  $v_{j}^{t}$  and we have  $\frac{\partial \mathbf{h}_{j}^{t+1}}{\partial \mathbf{h}_{j}^{t}} = \frac{\partial v_{j}^{t+1}}{\partial v_{j}^{t}} = \alpha$  and  $\frac{\partial v_{j}^{t}}{\partial W_{ji}} = z_{i}^{t-1}$  (for a derivation of the eligibility traces taking the reset into account we refer to section S1.2). Using these derivatives and equation (14), one obtains that the eligibility vector is the low-pass filtered presynaptic spike-train,

$$\boldsymbol{\epsilon}_{ji}^{t+1} = \mathcal{F}_{\alpha}(\boldsymbol{z}_i^t) \stackrel{\text{def}}{=} \boldsymbol{\bar{z}}_i^t , \qquad (22)$$

and following equation (13), the eligibility trace is:

$$e_{ji}^{t+1} = \psi_j^{t+1} \bar{z}_i^t . (23)$$

For all neurons j the derivations in the next sections also hold for synaptic connections from 619 input neurons i, but one needs to replace the network spikes  $z_i^{t-1}$  by the input spikes  $x_i^t$  (the 620 time index switches from t-1 to t because the hidden state  $\mathbf{h}_{j}^{t} = M(\mathbf{h}_{j}^{t-1}, \mathbf{z}^{t-1}, \mathbf{x}^{t}, \mathbf{W}_{j})$ 621 is defined as a function of the input at time t but the preceding recurrent activity). For 622 simplicity we have focused on the case where transmission delays between neurons in the 623 RSNN are just 1 ms. If one uses more realistic length of delays d, this -d appears in 624 equations (23)–(25) instead of -1 as the most relevant time point for presynaptic firing 625 (see section S1.3). This moves resulting synaptic plasticity rules closer to experimentally 626 observed forms of STDP. 627

Eligibility traces for ALIF neurons. The hidden state of an ALIF neuron is a two dimensional vector  $\mathbf{h}_{j}^{t} = [v_{j}^{t}, a_{j}^{t}]$ . Hence a two dimensional eligibility vector  $\boldsymbol{\epsilon}_{ji}^{t} \stackrel{\text{def}}{=} [\boldsymbol{\epsilon}_{ji,v}^{t}, \boldsymbol{\epsilon}_{ji,a}^{t}]$ 

is associated with the synapse from neuron i to neuron j, and the matrix  $\frac{\partial \mathbf{h}_{j}^{i+1}}{\partial \mathbf{h}^{t}}$  is a 2 × 2 630 matrix. The derivatives  $\frac{\partial a_j^{t+1}}{\partial a_j^t}$  and  $\frac{\partial a_j^{t+1}}{\partial v_j^t}$  capture the dynamics of the adaptive threshold. Hence to derive the computation of eligibility traces we substitute the spike  $z_j$  in equation 631 632 (10) by its definition given in equation (9). With this convention one finds that the diagonal 633 of the matrix  $\frac{\partial \mathbf{h}_{j}^{t+1}}{\partial \mathbf{h}_{j}^{t}}$  is formed by the terms  $\frac{\partial v_{j}^{t+1}}{\partial v_{j}^{t}} = \alpha$  and  $\frac{\partial a_{j}^{t+1}}{\partial a_{j}^{t}} = \rho - \psi_{j}^{t}\beta$ . Above and below the diagonal, one finds respectively  $\frac{\partial v_{j}^{t+1}}{\partial a_{j}^{t}} = 0$ ,  $\frac{\partial a_{j}^{t+1}}{\partial v_{j}^{t}} = \psi_{j}^{t}$ . Seeing that  $\frac{\partial \mathbf{h}_{j}^{t}}{\partial W_{ji}} = 0$ . 634 635  $\left[\frac{\partial v_j^t}{\partial W_{ji}}, \frac{\partial a_j^t}{\partial W_{ji}}\right] = \left[z_i^{t-1}, 0\right], \text{ one can finally compute the eligibility traces using equation (13).}$ 636 The component of the eligibility vector associated with the membrane potential remains 637 the same as in the LIF case and only depends on the presynaptic neuron:  $\epsilon_{ji,v}^t = \overline{z}_i^{t-1}$ . 638 For the component associated with the adaptive threshold we find the following recursive 639 update: 640

$$\epsilon_{ji,a}^{t+1} = \psi_j^t \overline{z}_i^{t-1} + (\rho - \psi_j^t \beta) \epsilon_{ji,a}^t , \qquad (24)$$

and, since  $\frac{\partial z_j^t}{\partial h_j^t} = \begin{bmatrix} \frac{\partial z_j^t}{\partial v_j^t}, \frac{\partial z_j^t}{\partial a_j^t} \end{bmatrix} = \begin{bmatrix} \psi_j^t, -\beta \psi_j^t \end{bmatrix}$ , this results in an eligibility trace of the form:

$$e_{ji}^{t} = \psi_{j}^{t} \left( \overline{z}_{i}^{t-1} - \beta \epsilon_{ji,a}^{t} \right).$$

$$(25)$$

Recall that the constant  $\rho = \exp(-\frac{\delta t}{\tau_a})$  arises from the adaptation time constant  $\tau_a$ , which typically lies in the range of hundreds of milliseconds to a few seconds in our experiments, yielding values of  $\rho$  between 0.995 and 0.9995. The constant  $\beta$  is typically of the order of 0.07 in our experiments.

To provide a more interpretable form of eligibility trace that fits into the standard form of local terms considered in 3-factor learning rules [9], one may drop the term  $-\psi_j^t\beta$  in equation (24). This approximation  $\hat{\epsilon}_{ji,a}^t$  of equation (24) becomes an exponential trace of the post-pre pairings accumulated within a time window as large as the adaptation adaptation time constant:

$$\widehat{\epsilon}_{ji,a}^{t+1} = \mathcal{F}_{\rho}\left(\psi_j^t \overline{z}_i^{t-1}\right) . \tag{26}$$

The eligibility traces are computed with equation (24) in most experiments, but the performances obtained with *symmetric e-prop* and this simplification were indistinguishable in the task where temporal credit assignment is difficult of Fig. 3.

#### <sup>654</sup> Synaptic plasticity rules resulting from *e-prop*

An exact computation of the ideal learning signal  $\frac{dE}{dz_j^t}$  in equation (1) requires to backpropagate gradients through time (see Fig. 6c). For online *e-prop* we replace it with the partial derivative  $\frac{\partial E}{\partial z_j^t}$ , which can be computed online. Implementing the weight updates with gradient descent and learning rate  $\eta$ , all the following plasticity rules are derived from the formula

$$\Delta W_{ji}^{\text{rec}} = -\eta \quad \sum_{t} \frac{\partial E}{\partial z_{j}^{t}} e_{ji}^{t} .$$
<sup>(27)</sup>

Note that in the absence of the superscript t,  $\Delta W_{ji}$  denotes the cumulated weight change over one trial or batch of consecutive trials but not the instantaneous weight update. This can be implemented online by accumulating weight updates in a hidden synaptic variable. Note also that the weight updates derived in the following for the recurrent weights  $W_{ji}^{\text{rec}}$ also apply to the inputs weights  $W_{ji}^{\text{in}}$ . For the output weights and biases the derivation does not require the theory of *e-prop*, and the weight updates can be found in the section S3.1.

**Case of regression tasks.** In the case of a regression problem with targets  $y_k^{*,t}$  and outputs  $y_k^t$  defined in equation (11), we define the loss function  $E = \frac{1}{2} \sum_{t,k} (y_k^t - y_k^{*,t})^2$ . This results in a partial derivative of the form  $\frac{\partial E}{\partial z_j^t} = \sum_k W_{kj}^{\text{out}} \sum_{t' \ge t} (y_k^{t'} - y_k^{*,t'}) \kappa^{t'-t}$ . This seemingly provides an obstacle for online learning, because the partial derivative is a weighted sum over future errors. But this problem can be resolved since one can interchange the two summation indices in the expression for the weight updates (see section S3.1). In this way the sum over future events transforms into a low-pass filtering of the eligibility traces  $\overline{e}_{ji}^t = \mathcal{F}_{\kappa}(e_{ji}^t)$ , and the resulting weight update can be written as

$$\Delta W_{ji}^{\text{rec}} = -\eta \sum_{t} \underbrace{\left(\sum_{k} B_{jk}(y_k^t - y_k^{*,t})\right)}_{=L_j^t} \overline{e}_{ji}^t .$$

$$(28)$$

**Case of classification tasks.** We assume that K target categories are provided in the form of a one-hot encoded vector  $\boldsymbol{\pi}^{*,t}$  with K dimensions. We define the probability for class k predicted by the network as  $\pi_k^t = \operatorname{softmax}_k(y_1^t, \ldots, y_K^t) = \exp(y_k^t) / \sum_{k'} \exp(y_{k'}^t)$ , and the loss function for classification tasks as the cross-entropy error  $E = -\sum_{t,k} \pi_k^{*,t} \log \pi_k^t$ . The plasticity rule resulting from *e-prop* reads (see derivation in section S3.1):

$$\Delta W_{ji}^{\text{rec}} = -\eta \sum_{t} \underbrace{\left(\sum_{k} B_{jk}(\pi_k^t - \pi_k^{*,t})\right)}_{=L_j^t} \bar{e}_{ji}^t .$$

$$(29)$$

#### <sup>679</sup> Reward-based e-prop: application of e-prop to deep RL

For reinforcement learning, the network interacts with an external environment. At any time t the environment can provide a positive or negative reward  $r^t$ . Based on the observations  $\boldsymbol{x}^t$  that are perceived, the network has to commit to actions  $a^{t_0}, \dots, a^{t_n}, \dots$  at certain decision times  $t_0, \dots, t_n, \dots$ . Each action  $a^t$  is sampled from a probability distribution  $\pi(\cdot | \boldsymbol{y}^t)$  which is also referred to as the policy of the RL agent. The policy is defined as function of the network outputs  $\boldsymbol{y}^t$ , and is chosen here to be a categorical distribution of K discrete action choices. We assume that the agent chooses action k with probability  $\pi_k^t = \pi(a^t = k | \boldsymbol{y}^t) = \operatorname{softmax}_k(y_1^t, \dots, y_K^t) = \exp(y_k^t) / \sum_{k'} \exp(y_{k'}^t).$ 

The goal of reinforcement learning is to maximize the expected sum of discounted 688 rewards. That is, we want to maximize the expected return at time t = 0,  $\mathbb{E}[R^0]$ , where 689 the return at time t is defined as  $R^t = \sum_{t' \ge t} \gamma^{t'-t} r^{t'}$  with a discount factor  $\gamma \le 1$ . The expectation is taken over the agent actions  $a^t$ , the rewards  $r^t$  and the observations from the 690 691 environment  $\boldsymbol{x}^t$ . We approach this optimization problem by using the actor-critic variant of 692 the policy gradient algorithm, which applies gradient ascent to maximize  $\mathbb{E}[R^0]$ . The basis 693 of the estimated gradient relies on an estimation of the policy gradient, as shown in section 694 13.3 in [13]. There, the resulting weight update is given in equation (13.8), where  $G_t$  refers 695 to the return  $R^t$ . Hence, the gradient  $\frac{d\mathbb{E}[R^0]}{dW_{ji}}$  is proportional to  $\mathbb{E}\left[\sum_{t_n} R^{t_n} \frac{d\log \pi(a^{t_n}|\boldsymbol{y}^{t_n})}{dW_{ji}}\right]$ , 696 which is easier to compute because the expectation can be estimated by an average over 697 one or many trials. Following this strategy, we define the per-trial loss function  $E_{\pi}$  as a 698 function of the sequence of actions  $a^{t_0}, \cdots, a^{t_n}, \cdots$  and rewards  $r^0, \cdots, r^T$  sampled during 690 this trial: 700

$$E_{\pi}(\boldsymbol{z}^{0},\cdots,\boldsymbol{z}^{T},a^{t_{0}},\cdots,a^{t_{n}},\cdots,r^{0},\cdots,r^{T}) \stackrel{\text{def}}{=} -\sum_{n} R^{t_{n}} \log \pi(a^{t_{n}}|\boldsymbol{y}^{t_{n}}) .$$
(30)

701 And thus:

$$\frac{d\mathbb{E}\left[R^{0}\right]}{dW_{ji}} \propto \mathbb{E}\left[\sum_{t_{n}} R^{t_{n}} \frac{d\log \pi(a^{t_{n}} | \boldsymbol{y}^{t_{n}})}{dW_{ji}}\right] = -\mathbb{E}\left[\frac{dE_{\pi}}{dW_{ji}}\right].$$
(31)

Intuitively, given a trial with high rewards, policy gradient changes the network output  $\boldsymbol{y}$ to increase the probability of the actions  $a^{t_n}$  that occurred during this trial. In practice, the gradient  $\frac{dE_{\pi}}{dW_{ji}}$  is known to have high variance and the efficiency of the learning algorithm can be improved using the actor-critic variant of the policy gradient algorithm. It involves the policy  $\pi$  (the actor) and an additional output neuron  $V^t$  which predicts the value function  $\mathbb{E}[R^t]$  (the critic). The actor and the critic are learnt simultaneously by defining the loss function as

$$E = E_{\pi} + c_V E_V , \qquad (32)$$

where  $E_{\pi} = -\sum_{n} R^{t_n} \log \pi (a^{t_n} | \boldsymbol{y}^{t_n})$  measures the performance of the stochastic policy  $\pi$ , and  $E_V = \sum_{t} \frac{1}{2} (R^t - V^t)^2$  measures the accuracy of the value estimate  $V^t$ . Since  $V^t$  is independent of the action  $a^t$  one can show that  $0 = \mathbb{E} \left[ V^{t_n} \frac{d \log \pi (\boldsymbol{a}^{t_n} | \boldsymbol{y}^{t_n})}{d W_{j_i}} \right]$ . We can use that to define an estimator  $\widehat{\frac{dE}{dW_{j_i}}}$  of the loss gradient with reduced variance:

$$-\frac{d\mathbb{E}\left[R^{0}\right]}{dW_{ji}} + c_{V}\mathbb{E}\left[\frac{dE_{V}}{dW_{ji}}\right] \propto \mathbb{E}\left[\frac{dE}{dW_{ji}}\right]$$
(33)

$$= \mathbb{E}\left[\underbrace{-\sum_{t_n} (R^{t_n} - V^{t_n}) \frac{d \log \pi (a^{t_n} | \boldsymbol{y}^{t_n})}{dW_{ji}} + c_V \frac{dE_V}{dW_{ji}}}_{\stackrel{\text{def}}{=} \frac{\widehat{dE}}{dW_{ji}}}\right],$$
(34)

similarly as in equation (13.11) of section 13.4 in [13]. A difference in notation is that  $b(S_t)$ refers to our value estimation  $V^t$ . In addition, equation (34) already includes the gradient  $\frac{dE_V}{dW_{ji}}$  that is responsible for learning the value prediction. Until now this derivation follows the classical definition of the actor-critic variant of policy gradient, and the gradient  $\frac{dE}{dW_{ji}}$ can be computed with *BPTT*. To derive *reward-based e-prop* we follow instead the generic online approximation of *e-prop* as in equation (27) and approximate  $\frac{dE}{dW_{ji}}$  by a sum of terms of the form  $\frac{\partial E}{\partial z_i^t} e_{ji}^t$  with

$$\frac{\widehat{\partial E}}{\partial z_j^t} = -\sum_n (R^{t_n} - V^{t_n}) \frac{\partial \log \pi(a^{t_n} | \boldsymbol{y}^{t_n})}{\partial z_j^t} + c_V \frac{\partial E_V}{\partial z_j^t} .$$
(35)

We choose this estimator  $\widehat{\frac{\partial E}{\partial z_j^t}}$  of the loss derivative because it is unbiased and has a low variance, more details are given in section S5.1. We derive below the resulting synaptic plasticity rule as needed to solve the task of Fig. 4, 5. For the case of a single action as used in Fig. S5 we refer to section S5.1.

When there is a delay between the action and the reward or, even harder, when a 724 sequence of many actions lead together to a delayed reward, the loss function E cannot be 725 computed online because the evaluation of  $R^{t_n}$  requires knowledge of future rewards. To 726 overcome this, we introduce temporal difference errors  $\delta^t = r^t + \gamma V^{t+1} - V^t$  (see Fig. 4), and 727 use the equivalence between the forward and backward view in reinforcement learning [13]. 728 Using the one-hot encoded action  $\mathbb{1}_{a^t=k}$  at time t, which assumes the value 1 if and only if 729  $a^t = k$  (else it has value 0), we arrive at the following synaptic plasticity rules for a general 730 actor-critic algorithm with *e-prop* (see section S5.1): 731

$$\Delta W_{ji}^{\text{rec}} = -\eta \sum_{t} \delta^{t} \mathcal{F}_{\gamma} \left( L_{j}^{t} \bar{e}_{ji}^{t} \right) \quad \text{for}$$
(36)

$$L_{j}^{t} = -c_{V}B_{j}^{V} + \sum_{k} B_{jk}^{\pi}(\pi_{k}^{t} - \mathbb{1}_{a^{t}=k}) , \qquad (37)$$

where we define the term  $\pi_k^t - \mathbb{1}_{a^t=k}$  to have value zero when no action is taken at time t.  $B_j^V$  is here the weight from the output neuron for the value function to neuron j, and the weights  $B_{jk}^{\pi}$  denote the weights from the outputs for the policy. <sup>735</sup> A combination of reward prediction error and neuron-specific learning signal was pre-<sup>736</sup>viously used in a plasticity rule for feedforward networks inspired by neuroscience [58, 59]. <sup>737</sup>Here it arises from the approximation of *BPTT* by *e-prop* in RSNNs solving RL problems. <sup>738</sup>Note that the filtering  $\mathcal{F}_{\gamma}$  requires an additional eligibility trace per synapse. This arises <sup>739</sup>from the temporal difference learning in RL [13]. It depends on the learning signal and <sup>740</sup>does not have the same function as the eligibility trace  $e_{ji}^t$ .

## 741 Code availability

An implementation of *e-prop* solving the tasks of Fig. 2 to 5 is made public together with the publication of this paper https://github.com/IGITUGraz/eligibility\_propagation.

## 744 Data availability

<sup>745</sup> Data for the TIMIT and ATARI benchmark tasks were published in previous works [22, 29].

<sup>746</sup> Data for the temporal credit assignment task are generated by a custom code provided in

<sup>747</sup> the abovementioned code repository.

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Authors contributions GB, FS, AS and WM conceived the work, GB, FS, AS, EH and DS carried out experiments and all authors contributed to the writing of the paper.

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