Training spiking neuronal networks to perform motor control using reinforcement and evolutionary learning

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Artificial neural networks (ANNs) have been successfully trained to perform a wide range of sensory-motor behaviors. In contrast, the performance of spiking neuronal network (SNN) models trained to perform similar behaviors remains relatively suboptimal. In this work, we aimed to push the field of SNNs forward by exploring the potential of different learning mechanisms to achieve optimal performance. We trained SNNs to solve the CartPole reinforcement learning (RL) control problem using two learning mechanisms operating at different timescales: (1) spike-timing-dependent reinforcement learning (STDP-RL) and (2) evolutionary strategy (EVOL). Though the role of STDP-RL in biological systems is well established, several other mechanisms, though not fully understood, work in concert during learning in vivo. Recreating accurate models that capture the interaction of STDP-RL with these diverse learning mechanisms is extremely difficult. EVOL is an alternative method and has been successfully used in many studies to fit model neural responsiveness to electrophysiological recordings and, in some cases, for classification problems. One advantage of EVOL is that it may not need to capture all interacting components of synaptic plasticity and thus provides a better alternative to STDP-RL. Here, we compared the performance of each algorithm after training, which revealed EVOL as a powerful method for training SNNs to perform sensory-motor behaviors. Our modeling opens up new capabilities for SNNs in RL and could serve as a testbed for neurobiologists aiming to understand multi-timescale learning mechanisms and dynamics in neuronal circuits.

KEYWORDS
reinforcement learning, evolutionary strategy, spiking neuronal networks, computer simulation, sensory-motor behavior, video games, spike-timing dependent plasticity (STDP)
Introduction

Reinforcement learning (RL) problems offer an ideal framework for comparing the learning strategies of an interactive, goal-seeking agent (Sutton and Barto, 2018). Most often, the best learning strategy is evaluated based on time efficiency and algorithm complexity. While there are many deep reinforcement learning algorithms for solving dynamical control problems (Mnih et al., 2015), biologically realistic network architectures and training strategies are not yet as efficient. In this work, using the CartPole RL problem, we compare biologically inspired learning algorithms based on the training efficiency and the resulting network dynamics.

As spiking neural networks (SNNs) are shown to be Turing-complete (Maass, 1996b) and computationally more powerful than artificial neural networks (ANNs) (Maass, 1997, 1996a), efficient learning strategies are still actively investigated (Tavanaei et al., 2019). SNNs have been effective for pattern recognition problems (Gupta and Long, 2007; Escobar et al., 2009; Kasabov et al., 2014; Tavanaei and Maida, 2017; Mozafari et al., 2018) but are rarely used for solving reinforcement learning control problems. As spiking neurons operate in the time domain, we show that RL problems are suitable for evaluating training strategies and providing insight into neural circuit dynamics.

Traditionally, when SNN models are trained to perform a behavior using biologically inspired learning mechanisms, algorithms used are variations on either spike timing dependent plasticity (STDP) (Tavanaei et al., 2019) or evolutionary strategies (EVOL) (Espinal et al., 2014). For learning behaviors from the reinforcement learning domain, STDP can be extended to use reward modulated plasticity (Chadderdon et al., 2012; Neymotin et al., 2013; Hazan et al., 2018; Patel et al., 2019; Anwar et al., 2022a), an algorithm denoted spike-timing dependent reinforcement learning (STDP-Rl). In this work, we introduce a new EVOL variation adapted from non-spiking neural networks (Salimans et al., 2017). Alternatively, there are many algorithms that learn behaviors through back-propagation (Bengio et al., 2017; Liu et al., 2017; Mostafa et al., 2017; Huh and Sejnowski, 2018; Bohre et al., 2022). Since back-propagation is challenging to properly implement in SNNs with arbitrary architectures (Patel et al., 2019; Anwani and Rajendran, 2020), we have not used it in this work.

Spike-timing dependent reinforcement learning trains SNNs by establishing associations between the neurons encoding the sensory environment and neurons producing an action or sequence of actions, such that appropriate actions are produced for specific sensory cues. The sensory-motor associations are established from reward-modulated synaptic weight changes that occur at each time step of the simulation. Hence, STDP-Rl trains at the individual level, as we consider each separate initialization of an SNN network a separate “individual”.

In our SNNs, we simulate individual neurons as event-based dynamical units that mimic functions of their biological counterparts, like adaptation, bursting, and depolarization blockade (Neymotin et al., 2011). For training SNNs based on population-level fitness metrics, we adapt EVOL (Salimans et al., 2017), which have been shown to be efficient in training Artificial Neural Networks on similar problems (Salimans et al., 2017; Chrabaszcz et al., 2018). The EVOL algorithm treats the synaptic weight as an individual’s genome, then perturbs the genome using the mutation genetic operator to produce an offspring population. Each offspring is an individual SNN that interacts with the environment by receiving sensory stimuli and producing motor actions based on its internal firing patterns. Based on the offspring fitness, we combine the population genomes to yield the next generation of genomes. In the EVOL algorithm, SNN weights are not adapting throughout the interaction with the environment.

To summarize, in this work, we investigate two algorithms for training SNNs on the CartPole RL problem: individual-level training using STDP-Rl and population-level EVOL. Our contributions are as follows: (1) we analyze the efficiency of biologically inspired training strategies for solving the CartPole RL problem and (2) we analyze the effect between individual-level and population-level learning on SNN’s sensorimotor mappings and neuronal dynamics.

Materials and methods

CartPole game

To test different learning strategies, we chose the classic CartPole problem of balancing a vertical pole on a cart (Barto et al., 1983; Geva and Sitte, 1993). All the simulations were run using the CartPole-v1 environment (Figure 1 left) available in the OpenAI Gym platform (Brockman et al., 2016). To keep the pole balanced (Figure 1 left), a force to the left (-1) or the right (+1) must be applied at each time step of the simulation. Once the force is applied, a new game-state is generated, resulting from the interaction between the previous game-state and the applied action. The environment is fully described by four observations: cart position, cart horizontal velocity, pole angle with respect to a vertical axis, and angular velocity of the pole. For simplicity, we will now reference those four observations as position, velocity, angle, and angular velocity (see Figure 1 left). The position is a relative distance from the center of the screen with positive (negative) values to the right (left). Similarly, the pole’s angle represents positive values for angles to the right (clockwise). The velocity and angular velocity represents the rate of change of the position and the angle, respectively. The

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1 https://www.gymlib.com/dev/
game is played in episodes, where each episode consists of multiple steps (left or right) taken to keep the pole balanced. The pole loses its balance when either the angle is greater than 15 degrees from vertical in either direction or the position is more than 2.4 units from the center in either direction. Each episode is allowed a maximum duration of 500 steps. An episode can be instantiated to different initial positions, which deterministically affects the trajectory through observational space.

Simulations

We used the NEURON simulation environment (Carnevale and Hines, 2006) with the NetPyNE package (Dura-Bernal et al., 2019) for all modeling. NEURON simulates the firing of individual neurons based on the integration of input activation. Neurons are assembled into populations and into a connected network using the NetPyNE package that further coordinates the network simulation environment. The integration of the CartPole environment and the NetPyNE network was implemented in Python. The CartPole environment and the network simulation (the agent) are synchronized every time step T (50 ms) in the following way:

- at the beginning of the time step, the environment is translated into neural activity in the input population (ES);
- for the duration of the time T, neurons are spiking based on induced activity; and
- at the end of the time step (after 50 ms), higher relative activation in the motor populations (EM-L, EM-R) determines the agent’s action.

In the following sections, we will present the setup of the pre-plasticity simulation: the excitatory/inhibitory neurons, the network inputs, the movement generation, and the weights initialization.

Constructing a spiking neuronal network model to play CartPole

Our SNN model was adapted from one of our recent models (Anwar et al., 2022a). To allow the SNN model to capture the game-state space reliably, we included 80 neurons in the sensory area (ES) with four subpopulations (20 neurons each), each to independently encode position, velocity, angle, and angular velocity (Figure 1 right box, “ES”). Each neuron was assigned to encode a different receptive field. Since the game’s goal was to balance the pole, that would require more precision in encoding sensory information near balanced states, around the absolute value of 0. To capture higher sensory precision utilizing smaller receptive fields around the balanced state and less precision utilizing larger receptive fields at peripheries, 80 neurons were assigned receptive fields to each neuron based on percentiles of a Gaussian distribution with a peak value of 0 centered around the 11th neuron. As such, lower indices neurons (neurons 1–10) encode negative values of the state variables in decreasing order. Similarly, higher indices neurons (neurons 11–20) encode positive values of the state variables in increasing order. All four ES populations were assigned receptive fields using Gaussian distributions with each input state’s expected mean and variance.

At each game step, 4 ES neurons, one from each subpopulation, were activated, informing the SNN model about the game state. To allow association between individual state-variable representing a game-state, we included 40 neurons in the association area “EA” (Figure 1 right box, “EA”), which received inputs from the ES neuronal population. Each neuron in EA was connected to motor areas EM-L and EM-R, generating Right- and Left-actions (Figure 1 “action” arrow) by comparing the number of spikes in those populations (winner-takes-all). If both subpopulations have the same number of spikes, then a random move is performed.

To prevent hyperexcitability and depolarization-block (Chadderdon et al., 2012; Neymotin et al., 2013; Anwar et al., 2022a), we included inhibitory neuronal populations (10 IA, 10 IAL, 10 IM, and 10 IML) and synapse weight normalization steps (detailed below).

Integrate-and-fire neuron

Individual neurons were modeled as event-driven, rule-based dynamical units with many of the key features found in real neurons, including adaptation, bursting, depolarization blockade, and voltage-sensitive NMDA conductance (Lytton and Stewart, 2006; Lytton and Omurtag, 2007; Lytton et al., 2008; Neymotin et al., 2011). Event-driven processing provides a faster alternative to network integration: a presynaptic spike is an event that arrives after a delay at a postsynaptic neuron; this arrival is then a subsequent event that triggers further processing in the postsynaptic neurons. Neurons were parameterized (Table 1) as excitatory (E), fast-spiking inhibitory (I), and low threshold activated inhibitory (II). Each neuron had a membrane voltage state variable ($V_m$), with a baseline value determined by a resting membrane potential parameter ($V_{rest}$). After synaptic input events, if $V_m$ crossed the spiking threshold ($V_{thresh}$), the cell would fire an action potential and enter an absolute refractory period, lasting $t_{AR}$ ms. After an action potential, an after-hyperpolarization voltage state variable ($V_{AHP}$) was increased by a fixed amount $\Delta V_{AHP}$, and then $V_{AHP}$ was subtracted from $V_m$. Then $V_{AHP}$ decayed exponentially (with the time constant $t_{AHP}$) to 0. Simulating the depolarization blockade, a neuron cannot fire if $V_m$ surpasses the blockade voltage ($V_{block}$). The relative refractory period was simulated after an action potential by increasing the firing threshold $V_{thresh}$ by $W_{RR}(V_{block}-V_{thresh})$, where $W_{RR}$ was a unitless weight parameter. $V_{thresh}$ then decayed exponentially to its baseline value with a time constant $t_{RR}$.
FIGURE 1
The CartPole game environment (left) interfacing with the SNN model (right). Left box: At each step, a new game state is produced, which is described by the variables (as labeled): Position, Velocity, Angle, and Angular Velocity. Arrow “state”: The values of these variables are used to activate a unique quadruple of neurons in the “ES” neuronal population (1 for each state variable). Right box: The neuronal network is represented as a diagram, as each dot represents one neuron, and arrows (light gray and light red) represent connectivity between populations of neurons. The light gray and red arrows represent excitatory and inhibitory synapses between neuronal populations, respectively. There are four excitatory neuronal populations: “ES” with 80 neurons (black-bordered box); “EA” with 40 neurons (green-bordered box); “EM-L” with 20 neurons (brown-bordered box); and “EM-R” with 20 neurons (cyan-bordered box). There are four inhibitory neuronal populations: “IA” with 10 neurons; “IAL” with 10 neurons; “IM” with 10 neurons; and “IML” with 10 neurons (each within a red-bordered box). Excitatory neuronal populations have outgoing excitatory connections, while inhibitory neuronal populations have outgoing inhibitory connections. The connections marked with a ‘±’ sign represent the connections that undergo synaptic plasticity: the connections between populations: “ES” to “EA”, “EA” to “EM-L”, and “EA” to “EM-R”. Arrow “action”: The activity within the “EM-L” and “EM-R” neuronal populations determines the action performed by the agent. Higher activity within either neuronal population will determine the agent to make a move to the left or to the right, as exemplified by the large red arrows in the left box. Green arrow “reward”: for the Reinforcement Learning training strategy, the state of the environment is used to dictate the reward (or punishment) administered to the network for a correct (or incorrect) move.

TABLE 1 Parameters of the neuron model for each type.

| Cell type                  | $V_{\text{rest}}$ (mV) | $V_{\text{thresh}}$ (mV) | $V_{\text{block}}$ (mV) | $\tau_{AR}$ (ms) | $W_{RR}$ | $\tau_{RR}$ (ms) | $\Delta V_{\text{AHP}}$ (mV) | $\tau_{AHP}$ (ms) |
|---------------------------|------------------------|--------------------------|-------------------------|------------------|----------|------------------|-----------------------------|------------------|
| Excitatory (E)            | -65                    | -40                      | -25                     | 5                | 0.75     | 8                | 1                           | 400              |
| Inhibitory (I)            | -63                    | -40                      | -10                     | 2.5              | 0.25     | 1.5              | 0.5                         | 50               |
| Low-threshold Inhibitory (IL) | -65                 | -47                      | -10                     | 2.5              | 0.25     | 1.5              | 0.5                         | 50               |

$V_{\text{rest}}$, resting membrane potential; $V_{\text{thresh}}$, spiking threshold; $V_{\text{block}}$, depolarization blockade voltage; $\tau_{AR}$, absolute refractory time constant; $W_{RR}$, relative refractory weight; $\tau_{RR}$, relative refractory time constant; $\Delta V_{\text{AHP}}$, after-hyperpolarization increment in voltage; $\tau_{AHP}$, after-hyperpolarization time constant.

Synaptic mechanisms

In addition to the intrinsic membrane voltage state variable, each cell had four additional voltage state variables $V_{\text{syn}}$, corresponding to the synaptic inputs. These represent AMPA (AM2), NMDA (NM2), and somatic and dendritic GABAA (GA and GA2) synapses. At the time of input events, synaptic weights were updated by step-wise changes in $V_{\text{syn}}$, which were then added to the cell’s overall membrane voltage $V_{\text{m}}$.

To allow for dependence on $V_{\text{m}}$, synaptic inputs changed $V_{\text{syn}}$ by $dV = W_{\text{syn}}(1 - V_{\text{m}}/E_{\text{syn}})$, where $W_{\text{syn}}$ is the synaptic weight, and $E_{\text{syn}}$ is the reversal potential relative to $V_{\text{rest}}$. The following values were used for the reversal potential $E_{\text{syn}}$: AMPA, 0 mV; NMDA, 0 mV; and GABAA, −80 mV. After synaptic input events, the synapse voltages $V_{\text{syn}}$ decayed exponentially toward 0 with time constants $\tau_{\text{syn}}$. The following values were used for $\tau_{\text{syn}}$: AMPA, 20 ms; NMDA, 300 ms; somatic GABAA, 10 ms; and dendritic GABAA, 20 ms.
The delays between inputs to dendritic synapses (dendritic GABA) and their effects on somatic voltage were selected from a uniform distribution ranging between 3–12 ms, while the delays for somatic synapses (AMPA, NMDA, somatic GABA) were selected from a uniform distribution ranging from 1.8 to 2.2 ms. Synaptic weights were fixed between a given set of populations except for those involved in learning (see the “±” sign in Figure 1 Right box and plasticity “on” in Table 2).

The neuronal weights

The neurons are organized into three overall layers: Sensory, Association, and Motor (Figure 1 right box). The sensory layer consists of the excitatory sensory neuronal population (ES), which contains a total of 80 excitatory neurons. The association layer consists of 40 excitatory neurons in the excitatory association neuronal population (EA), 10 fast-spiking inhibitory neurons in the inhibitory association neuronal population (IA), and 10 low threshold activated inhibitory neurons in the inhibitory association neuronal population (IAL). Similarly, the motor layer consists of 40 excitatory neurons in the excitatory motor neuronal population (EM), 10 fast-spiking inhibitory neurons in the inhibitory motor neuronal population (IM), and 10 low threshold activated inhibitory neurons in the “low” inhibitory association neuronal population (IML). Furthermore, the EM neuronal population is split into 20 neurons associated with left movements (EM-L) and 20 neurons associated with right movements (EM-R).

The weights between populations were adjusted to allow reliable transmission of spiking activity across different layers/areas of the SNN model. Each row in Table 2 describes the synaptic connectivity between two different neuronal populations (no self connectivity) and follows the same diagram of neuronal connectivity described in Figure 1. Neurons belonging to the presynaptic population have axons that project to neurons in the postsynaptic population. Each neuron projects to a fixed number of postsynaptic neurons, a

| Presynaptic population | Postsynaptic population | Connection convergence | Synapse type | Synaptic weight: $W_{syn}$ | Plasticity (empty for Off) |
|------------------------|-------------------------|------------------------|--------------|-----------------------------|---------------------------|
| ES                     | EA                      | 25                     | AM2          | 10.0                         | On                        |
|                        |                         |                        | NM2          | 0.196                       |                           |
| EA                     | IA                      | 15                     | AM2          | 5.85                         |                           |
|                        |                         |                        | NM2          | 0.0585                       |                           |
| EA                     | IAL                     | 15                     | AM2          | 5.94                         |                           |
|                        |                         |                        | NM2          | 0.294                        |                           |
| EA                     | EM                      | 20                     | AM2          | 6.5                          | On                        |
|                        |                         |                        | NM2          | 0.1                          |                           |
| IA                     | EA                      | 4                      | GA           | 18.0                         |                           |
| IA                     | IA                      | 1                      | GA           | 4.5                          |                           |
| IA                     | IAL                     | 2                      | GA           | 4.5                          |                           |
| IAL                    | EA                      | 4                      | GA2          | 5.0                          |                           |
| IAL                    | IA                      | 2                      | GA2          | 2.25                         |                           |
| IAL                    | IAL                     | 1                      | GA2          | 5.5                          |                           |
| EM (EM-L + EM-R)       | IAL                     | 16                     | AM2          | 5.85                         |                           |
|                        |                         |                        | NM2          | 0.0585                       |                           |
| EM (EM-L + EM-R)       | IML                     | 16                     | AM2          | 2.94                         |                           |
|                        |                         |                        | NM2          | 0.294                        |                           |
| IM (EM-L + EM-R)       | EM                      | 4                      | GA           | 18.0                         |                           |
| IM (EM-L + EM-R)       | IML                     | 16                     | AM2          | 2.94                         |                           |
| IM                     | EM                      | 4                      | GA           | 18.0                         |                           |
| IM                     | IML                     | 2                      | GA           | 4.5                          |                           |
| IM                     | EM                      | 2                      | GA           | 4.5                          |                           |
| IM                     | IML                     | 1                      | GA           | 4.5                          |                           |
constant defined as the connection convergence (Table 2). The individual neuronal connections are picked randomly at the initialization process, and different random seed values generate different connections, hence different networks. The excitatory neuron have both AMPA (AM2) and NMDA (NM2) synaptic connections, while the inhibitory neurons either have somatic GABA synapses (GA) for fast-spiking inhibitory neurons or have the dendritic GABA synapses (GA2) for low threshold activated inhibitory neurons. The excitatory neuron connection was picked based on previous studies and fine-tuned on this network to start with a biologically reasonable spiking pattern (2–20 Hz).

Some of the synapse weights can be changed throughout the course of the training simulation as they undergo synaptic plasticity. For this work, we limited synaptic plasticity to AMPA synapses between excitatory populations (Table 2: Plasticity column). We found that this limitation is not hindering the network’s ability to learn the dynamical behavior of the CartPole problem, and it rather simplifies the network analysis. To have a consistent comparison of different training strategies, we used the same initialization methods and plastic synapses as defined above and in Table 2.

Training strategies

Spike-timing dependent reinforcement learning

To train the neuronal networks, we used an existing STDP-RL mechanism, developed based on the distal reward learning paradigm proposed by Izhikevich (2007), with variations used in spiking neuronal network models (Chadderdon et al., 2012; Neymotin et al., 2013; Chadderdon and Sporns, 2006; Duran-Bernal et al., 2016; Anwar et al., 2022a). Our version of STDP-RL (Figure 2A) uses a spike-time-dependent plasticity mechanism together with a reward or punishment signal for

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**FIGURE 2**

Training SNN using STDP-RL and EVOL strategies. (A) In STDP-RL, when a postsynaptic neuron produced a spike within a short-time interval of a presynaptic spike, the synapse between the pair of neurons was tagged with a decaying eligibility trace. The tagged synapse was strengthened or weakened proportional to the value of eligibility trace for a reward or punishment, respectively. (B) Schematic showing the steps of evolutionary strategies training algorithm (EVOL).
the potentiation or depression of the targeted synapses. When a postsynaptic spike occurred within a few milliseconds of the presynaptic spike, the synaptic connection between this pair of neurons became eligible for STDP-RL and was tagged with an exponentially decaying eligibility trace. An exponentially decaying eligibility trace was included to assign temporally distal credits to the relevant synaptic connections. Later, when a reward or a punishment was delivered before the eligibility trace decayed to zero, the weight of the tagged synaptic connection was increased or decreased, depending on the ‘critic’ value and sign, i.e., increase for reward or decrease for punishment. Since we used an eligibility trace with an exponent of 1 in this work, the change in synaptic strength was constant at the time of the critic’s delivery.

Traditionally, when using STDP-RL for learning behavior, all plastic synaptic connections in the neuronal network model are treated equally (non-targeted STDP-RL). This strategy considers that the underlying causality between pre and postsynaptic neurons and the associated reinforcement automatically changes only relevant synaptic connections. On top of the traditional STDP-RL approach, we used two recently developed versions of targeted reinforcement by selectively delivering reward and punishment to different subpopulations of the Motor population (EM) (Anwar et al., 2022a). In the first variation (targeted RL main), we delivered reward or punishment only to the neuronal subpopulation that generated the action (EM-LEFT or EM-RIGHT). In the second variation (targeted RL both), we additionally delivered opposite and attenuated reinforcement to the opposite-action neuronal subpopulation. Both targeted methods ensured that the learning was specific to the part of the circuit which generated the action. Moreover, we explored delivering (attenuated) “critic” values to the neuronal populations one synapse away from those directly generating motor actions (EA population). We used and evaluated all six STDP-RL mechanisms (three targeted RL versions X two non-motor RL versions) for learning performance during hyperparameter search (see below for details). In all cases, although there is evidence of plasticity involving inhibitory interneurons in vivo (Vogels et al., 2011; Anwar et al., 2017), for the sake of simplicity, STDP-RL was only applied between excitatory neurons.

Critic

For STDP-RL, the model relies on a critic to provide essential feedback on the model’s actions (Figure 2A). For CartPole, we picked a critic that responds positively to movements that bring the vertical pole closer to a balanced position. We computed a loss for each position, determined by the absolute values of the angle and the angular velocity input states. The critic’s returned value will be the difference between the loss of the previous state and the loss of the current state. If the loss between the following states increases due to the agent’s move, the critic will return a negative value, corresponding to a punishment. Similarly, a decrease in loss will return a positive critic value, corresponding to a reward. If the agent could not decide on a motor move due to identical spiking activity in both subpopulations, then a constant negative punishment is returned. Additionally, as the critic is dominated by punishment at the beginning of training, to avoid the weights decreasing to zero, the model needs an associated boost in positive rewards ($\eta_{\text{positivity}}$). The critic is finally capped at a minimum and maximum value to keep rewards within the interval: $[-\max_{\text{reward}}, \max_{\text{reward}}]$

$$\text{loss}(t) = \sqrt{\text{ang}(t)^2 + \eta_{\text{angvel}} \cdot \text{angvel}(t)^2}$$  \hspace{1cm} (1)

$$\text{reward}(t) = \begin{cases} 0 & \text{if } \text{loss}(t - 1) < 10^{-2} \\ -\max_{\text{reward}} / \eta_{\text{positivity}} & \text{if } \text{no}_\text{move}(t) \\ \max_{\text{reward}} / \eta_{\text{positivity}} & \text{if } \text{loss}(t) < 10^{-2} \\ \text{loss}(t - 1) - \text{loss}(t) & \text{otherwise} \end{cases}$$  \hspace{1cm} (2)

$$f(r) = \begin{cases} r \cdot \eta_{\text{positivity}} & \text{if } r > 0 \\ r & \text{otherwise} \end{cases}$$  \hspace{1cm} (3)

$$\text{critic}(t) = \max(-\max_{\text{reward}}, \min(f(\text{reward}(t)) \cdot \text{gain}, \max_{\text{reward}}))$$  \hspace{1cm} (4)

Where:

- $\text{ang}(t)$ and $\text{angvel}(t)$ represent the input states at time step $t$;
- $\eta_{\text{angvel}}$ represents the angular velocity bias, used to balance the dominance of each input state;
- $\eta_{\text{positivity}}$ represents the positivity bias to reinforce rewarding behavior;
- $\max_{\text{reward}}$ is the maximum reward used, fixed at 1.0 throughout the whole experiment;
- $\text{gain}$ represents a final multiplier that increases the absolute reward value.

The critic was implemented as a crude reinforcer of synaptic plasticity, working in conjunction with STDP events. As we found in the HyperParameter search (described below), most of the hyperparameters of the critic have little influence over the final performance of the model, and we believe that many different critic functions would have been suitable for our analysis. More importantly, for synaptic weight normalization, the critic values are further modulated by output gain and homeostatic gain control, as described below.

Hyperparameter search

We first trained our SNN model using the STDP-RL parameters’ values as used in earlier studies (Dura-Bernal...
and found that the model did not perform very well since it could not learn to balance the CartPole for 50 steps per episode (averaged over 100 episodes). The low performance indicated that these parameters might not be optimal for training with STDP-RL. To find an optimal training strategy, we perturbed the parameters of our STDP-RL training setup. We identified nine hyperparameters to optimize (Supplementary Table 1), related specifically to the STDP-RL mechanism, that are independent of each other. Out of the nine: three parameters influence the timing and weight update of the STDP-modulated AMPA synapse (AMPA-RLwindhebb, AMPA-RLhebbwt); four parameters determine the area of effect of STDP (Targeted_RL_Type, Non-Motor_RL, Targeted_RL_Opp_EM, Targeted_RL_Non_Motor); and two parameters influence the reinforcement derived from the critic (Critic Positivity Bias $\eta^{\text{positivity}}$, Critic Angry Bias $\eta^{\text{angry}}$). To identify the best combination of the hyperparameters for training the network, we ran multiple random hyperparameter searches on those nine hyperparameters (Supplementary Table 1).

For the first hyperparameter search, we evaluated the nine parameters above by training networks with random choices of those hyperparameters. From the 7,200 possible combinations, we sampled 50 combinations and trained randomly-initialized models using STDP-RL for 500 s in simulation time. Finally, we evaluated the performance of those models based on the average steps per episode over 100 episodes during training (Supplementary Figure 1). Only the Targeted_RL_Type hyperparameter performance distributions showed a significant deviation from the mean performance (ANOVA $p$-value $< 10^{-5}$), but it failed the homogeneity of variance assumption test.

For the second step of the hyperparameter search, we continued training from the four best models from the previous step and evaluated the nine hyperparameters. For this step, we had 6,912 combinations from which we sampled 54 combinations that we trained for 2000 seconds in simulation time. Similarly, we evaluated the performance of those models based on the average steps per episode over 100 episodes during training (Supplementary Figure 2). The initialization model choice displayed a significant contribution to the final model performance (ANOVA $p$-value = 0.00012). Moreover, the hyperparameter defining the maximum time between pre- and postsynaptic spike (AMPA-RLwindhebb), also showed a minimal deviation from the mean performance (ANOVA $p$-value = 0.027).

For the third step of the hyperparameter search, we continued training from the best four models from the previous step and evaluated the nine hyperparameters. For this step, we had 1,728 combinations from which we sampled 56 combinations that we trained for 2,000 s in simulation time. Similarly, we evaluated the performance of those models based on the average steps per episode over 100 episodes during training (Supplementary Figure 3). Again, the initialization model choice displayed a significant contribution to the final model performance (ANOVA $p$-value = 0.00023). Moreover, the hyperparameter defining the choice of activating plasticity for the synapses within the ES $\rightarrow$ EA pathway (Non-Motor_RL) showed a minimal deviation from the mean performance (ANOVA $p$-value = 0.005). This latter finding was further tested in detail by training multiple models with and without ES $\rightarrow$ EA plasticity (see section “Results”).

Most of the hyperparameters had a minimal effect on the final training performance for the STDP-RL model as there is no significant difference between the performance of models with different hyperparameter values. Interestingly, the hyperparameter search revealed better preference when using the “targeted RL both” paradigm. These findings suggest that targeted plasticity of specific motor areas could enhance the learning ability of the model, consistent with earlier findings (Chadderdon et al., 2012; Hazan et al., 2018; Patel et al., 2019; Anwar et al., 2022a).

**Training protocol for the STDP-RL model**

Since we couldn’t establish the best choice for each of the hyperparameters, we used the training protocol of the best model resulting from the third hyperparameter step. Hence, we trained our STDP-RL models with a sequence of the hyperparameter values for different durations (500s, 2000s, 2000s). Then, continuing training after the 4500s mark, we used the hyperparameters of the third configuration. The exact hyperparameter values used are highlighted in bold in Supplementary Table 1.

**Evolutionary strategies**

The EVOL algorithm has been shown to be an effective gradient-free method for training ANNs in reinforcement learning control problems (Salimans et al., 2017) (note that the term evolutionary here is only used in a loose sense, and the algorithmic implementation is not intended to accurately follow evolutionary processes observed in biology). Here we adapt this learning technique to SNNs to solve the CartPole problem by procedurally adjusting the plastic weights of the SNN. Our method doesn’t change the timing delays, contrary to some EVOL used for SNNs (Altamirano et al., 2015). It should be noted that in the ANN implementation of EVOL weights are allowed to be unrestricted in value, so additive weights were used. As SNNs don’t have negative weights, we instead use a multiplicative noise, i.e., we increase or decrease the weights by a randomly selected percentage. In this way, we are able to restrict
the SNN weights to valid positive values while still effectively searching all possible parameterizations.

Formally our EVOL algorithm (Figure 2B) consists of the following steps to change the weight for each synapse, performed in parallel for the whole model: (1) at iteration \( i \), keep track of the current best synapse weight: \( w_{ij}^* \); (2) sample a population \( (P) \) of weight perturbations \( \epsilon_{ij} \) from the normal distribution; (3) for each weight perturbation \( (\epsilon_j) \), evaluate the whole network weights \( (w_i \ast (1 + \sigma \ast \epsilon_j)) \) for a fixed number of episodes \( (X) \); (4) measure the fitness \( (F_j) \) as the average count of steps achieved during the \( X \) episodes; (5) Normalize population fitness values \( (N_j) \) by subtracting the population mean fitness and dividing by the population mean standard deviation \( (\sigma) \); (6) modulate the weight perturbations based on the normalized fitness and derive a new best synapse weight:

\[
w_{ij+1} = w_{ij} \ast (1 + \alpha \ast \sigma \ast \epsilon \ast N / P)
\]  

(5)

Where \( \alpha \) is the learning rate, \( \sigma \) is the noise variance, and \( \epsilon \) and \( N \) are the vector representations of the weight perturbations for each synapse and the normalized fitness, respectively. We only update the weights that undergo synaptic plasticity (Table 2). The weights are initialized \( (w_{ij}) \) as the same initial weights we used for the STDP-RL model. In this case, STDP was fully deactivated and the EVOL training procedure updated the synaptic weights every iteration.

We trained using the EVOL algorithm on multiple models for 1,500 iterations and a population of \( P = 10 \) with synaptic weight perturbations of \( \sigma = 0.1 \) variance. We used a learning rate of \( \alpha = 1.0 \). We used 5 and 10 episodes (value \( X \) above) during fitness evaluation. The model weights were initialized based on the best and worst models trained with STDP-RL.

Synaptic weight normalization

Training the model with STDP-RL, the synaptic weights tend to increase without bound, leading to epileptic activity (Rowan et al., 2014). To avoid this behavior, we incorporated biologically-realistic normalization methods (Rowan and Neymotin, 2013; Sanda et al., 2017). Apart from the inhibition mechanisms described earlier, we used the following techniques: balancing synaptic input, output balancing, and homeostatic gain control.

To balance the combined synaptic input to each neuron, we normalize the total reception weight (defined as the sum of synaptic weights on each postsynaptic neuron from multiple presynaptic neurons) to its initial values every 25 time steps. This procedure keeps the neuronal inputs constant by decreasing the weights of specific unused synapses or promoting beneficial synapses, creating synaptic competition.

On the neuronal output side, we want to prevent synapses from overwhelming a primary synapse with constant rewards. To avoid such ever-increasing weights, we normalize each synapse's reinforcement by scaling it based on the neuron's total transmission weight (defined as the sum of synaptic weights from a presynaptic neuron onto multiple postsynaptic neurons). Hence, a synapse with a high transmission weight compared to initialization will decrease the reward strength and increase the punishment strength during such STDP-RL events. Conversely, a synapse with a low transmission weight compared to the initialization will increase the reward strength while decreasing the punishment strength. For this normalization step, we cap the maximum scaling factor for promoting STDP-RL events to 0.1 for scaling down and 2.0 for scaling up.

As a final normalization method, we want to keep neuronal populations within a target spiking rate (measured over 500 time steps). The homeostatic gain control method updates the target population synaptic inputs every 75 time steps after checking if the firing rate differs from the target firing rate (5.5 Hz for EA and 6.0 Hz for EM). The homeostatic gain control normalization method doesn’t update the synaptic weights directly but rather scales the expected total reception weight (by 0.01%) that is later used for normalization during the balancing of synaptic inputs (see above). This procedure reduces high neuronal activity and promotes baseline activity (Rowan and Neymotin, 2013; Sanda et al., 2017). As this update is infrequent (every 75 time steps) and has a minor indirect update (scaling by 0.01%), over a large amount of training, this method has an effect similar to network noise.

Validation, testing, and all-inputs datasets

For testing the trained models, we used a validation dataset for selecting the best model time-point and a testing dataset for reporting the final model performance. For those two datasets, we set a seed value for the OpenAI gym environment to fix the episodes and consistently get the same episode initialization. This environment seed \((S_{r1}, S_{r2}, \ldots, S_{r9})\) should not be confused with the model seed \((\text{Seed}-6, \text{Seed}-3, \ldots)\) that we use to randomly initialize the neuronal connections. The episodes are fixed for our datasets when they have the same initial starting conditions for all four environment parameters (position, velocity, angle, and angular velocity). While training the models, the episodes are not fixed and are randomly selected for each new training instantiation. To choose the best time-point of a trained model, we evaluate the model with weights at different time-points throughout training using a validation dataset with 100 fixed episodes. Further, we evaluate the picked model and report the performance on the testing dataset, which contains another 100 fixed episodes.

As each input parameter space is discretized into the activation of twenty individual neurons, the total possible combinations of inputs is 20^5. We tested the models on all the possible combinations, the All-Inputs dataset, by activating each possible input combination at a time. Sensory activations were interleaved with blank periods (no stimulation) to ensure that the activation does not have spikes into the next sensory activation.
Software

All modeling and analysis source code is available on github at the following repository location https://github.com/NathanKlineInstitute/netpyne-STDP.

Results

Like in any multilayer artificial neural network, the biggest challenge of training SNN models to perform sensory-motor behaviors is to come up with a network design and choose the right set of learning parameters resulting in optimal performance. This leaves model developers with many possibilities. Instead of trying many network configurations, here we constrained our network to a minimal configuration (see Methods) based on our previous study (Anwar et al., 2022a), where the SNN models learned to play a racket-ball game showing robust performance using several STDP-RL setups. Our model included 3 layers of excitatory neuronal populations: ES representing a sensory area, EA representing an association area, and EM representing a motor area with probabilistic inter-area connections and fixed initial synaptic weights (see section “The Neuronal Weights”). Inhibitory neurons in each area were included to prevent network hyperexcitability after increased synaptic weights from learning. The activity in the network propagated in a single direction: ES → EA → EM. To compare different learning strategies, we further constrained our network to two sets of plasticity-based configurations: 1. only EA → EM synapses are plastic, and 2. Both ES → EA and EA → EM synapses are plastic. Using these two plasticity constraints, we trained our SNNs with STDP-RL and EVOL methods, evaluated their performances, and dissected the circuits post-training to compare the underlying dynamics resulting in diverse behavioral performances.

Training multilayered SNN models using STDP-RL improved performance

We first trained our SNN model using STDP-RL, allowing only EA → EM synaptic connections to adapt. Since the number of neurons was small in each layer and inter-layer connections were probabilistic, there was a possibility that each initialization would result in a slightly different connectivity pattern, which might affect the learning capabilities of the model. To assess the effect of network initializations on the training, we trained the model using 10 different initializations (random number generator seeds). Each training session included a large number of CartPole episodes, where each episode was randomly initialized. Such random initializations of the game exposed the model to different sensory states representing the game environment and required the model to learn different action strategies during each episode. Learning a sensory-action strategy for an episode resulting in a higher performance does not necessarily mean that the model will perform equally well during the next episode, as evident by large fluctuations in the performance during the training (blue dots in Figure 3A). Such variability in the performance makes it difficult to accurately quantify the learning performance of the model. Therefore, we averaged the performance over 100 episodes to quantify the learning quality of the model (Figure 3A). We continued training each model until the performance converged or started decreasing, which typically required training over simulation durations of 25000 seconds. We observed a strong effect of the model initialization on its performance, as the maximum performances of the models during training ranged from 75 to 157, with a mean of 118 steps per episode (Figures 3B,E).

To further investigate whether allowing synaptic connections between the sensory and the association layer (ES → EA) to learn via STDP-RL in addition to EA → EM synaptic connections will enhance the capability of our model to perform better, we repeated training simulations using 10 different network initializations (same random number generating seeds as in Figure 3B). The models showed large variance in the performance from episode to episode during training as well as among themselves, similar to the models with only EA → EM plasticity (Figures 3C vs. A). Although for many seeds, the peak performance was more consistent (Figure 3E right bar) regardless of varying performance trajectories (Figure 3D). Surprisingly, the performance did not improve significantly (Figures 3B,D,E). Note that we used the same learning parameters in these simulations as in the models where only EA → EM synaptic connections were plastic (parameter values optimized using hyperparameter search, see section “Materials and Methods”). Since these models did not perform better than the models with only EA → EM plasticity, we only considered models with plasticity in the EA → EM pathway for the remainder of the manuscript when comparing to EVOL-based models and analyses.

As shown in Figure 3, we observed large variability in performance from episode to episode due to the dynamically evolving sensory-state of the environment across episodes. Therefore, it remains unclear whether the performance will sustain after learning is stopped and weights are frozen. To test post-training performance, we further evaluated only two models with unique weights based on their performance during training; the best performing model (seed-6 in Figure 3B) and the worst performing model (seed-3 model, Figure 3B). We evaluated the selected models using 100 episodes with the same game initializations for both models and compared them to the model before training and to a random choice...
null model (Figure 4). During training, both models improved their performance throughout STDP-RL training as we see an increase from 19 to 130 median steps per episode for the Seed-6(best) model and from 19 to 53 median steps per episode for the Seed-3(worst) model. These performances were better than before training compared to the random choice null models (Figure 4). Although the performance remained suboptimal (i.e., 500 steps per episode), both models showed enhanced performance after training (Figure 4). One-way ANOVA on the logarithm of the model performances before and after training resulted in $p < 10^{-10}$.

Multilayered SNN models trained using EVOL achieve optimal performance

Variability in performance due to different network configurations resulting from different network initializations suggests that the performance of these models is sensitive to the initial synaptic connections and corresponding weights. Although synaptic weights between two populations of neurons in the neocortex may vary greatly, in modeling SNNs, the initial weights are often drawn from a normal distribution around arbitrarily chosen and tested values, which may bias the ability of the network to learn a behavior. Evolutionary algorithms can allow parsing the weights of synaptic connections over a large range of values. The subsequent performance evaluation of each set of synaptic weights can guide the model to evolve along complex and highly non-linear trajectories, eventually producing models with better performance. To that end, we next evaluated the potential of our models to learn sensory-motor behavior utilizing the EVOL algorithm.

Similar to training using STDP-RL, we first trained our SNN model using the EVOL algorithm with plasticity
limited to the $EA \rightarrow EM$ pathway. While the STDP-RL training performance plateaued before reaching an average of 160 steps per episode, some of the EVOL models performed better than 200 steps per episode (Figures 5B,E). In addition, we observed a relatively smaller variance in performance from one iteration to the next (Figure 5A). One reason for the low variance in performance across iterations is due to the averaging over 50 episodes (5 episodes per perturbation) per iteration compared to showing each individual episode in STDP-RL learning models (Figure 3A). The performance of our EVOL-trained models increased further when using SNN models with both $ES \rightarrow EA$ and $EA \rightarrow EM$ plastic synaptic connections. In fact, the average performance increased to the optimal performance independent of the model seed but also on the game’s initial state. Altogether the EVOL model showed optimal performance independent of the model and game initializations, thereby also demonstrating robustness.

Variability in the performance of models trained using STDP-RL is partially related to the differences in the sensory environment (game initializations)

To demonstrate that the model learned the behavior and did not forget it, we earlier compared two different STDP-RL models (with different seeds) with the random-action generating and untrained models (Figure 4) and found variable yet sustained learning in the STDP-RL models. We also observed negligible variance in the performance for models trained using EVOL. As demonstrated earlier (Figures 3–5 and Table 3), this performance variability could be related to the initialization of synaptic weights or the presence/absence of synaptic connections between specific pairs of neurons or on the initial game-state (note that we reset the game-states at the end of each episode). To test the latter possibility, we handpicked nine unique initial game-states based on the performance of each model on the initial evaluation. We repeatedly played the initial game-states on each trained model for 25 episodes (Figures 3–5). The performance of our EVOL-trained models increased further when using SNN models with both $ES \rightarrow EA$ and $EA \rightarrow EM$ plastic synaptic connections. In fact, the average performance increased to the optimal performance independent of the model seed but also on the game’s initial state. Altogether the EVOL model showed optimal performance independent of the model and game initializations, thereby also demonstrating robustness.
FIGURE 5
Performance of the SNN model when training with EVOL. (A,B) Training SNN model with only \( EA \rightarrow EM \) plastic. (A) Training performance (defined as the length of time the model could keep the pole balanced) of a model initialized using Seed-3. During each iteration, 10 models with unique weights were evaluated on 5 episodes. Each red dot shows an average performance of (10 models x 5 episodes/model = 50 episodes), whereas each green dot and blue dot show minimum and maximum performance of 50 episodes. (B) Training performance of 10 randomly initialized models. All models were trained for the same number of iterations as opposed to the same amount of simulated time. (C,D) Same as A and B, respectively, with both \( ES \rightarrow EA \) and \( EA \rightarrow EM \) synaptic connections plastic. (E) Comparing the peak average performance of 10 randomly initialized models with only \( EA \rightarrow EM \) plastic synapses vs. both \( ES \rightarrow EA \) and \( EA \rightarrow EM \) plastic synapses. The box plot represents the distribution of maximum averages over 100 steps of the models. Each dot of the scatter plot represents the maximum average over 100 steps of one of the randomly initialized models.

TABLE 3 Performance comparison of training strategies on the testing dataset.

| Model                      | Average on test dataset | Median on test dataset | Trained episodes | Iterations | Population |
|----------------------------|-------------------------|------------------------|------------------|------------|------------|
| Seed-6 after initialization| 23.38                   | 19.5                   | 0                | –          | –          |
| Seed-6 after STDP-RL training| 144.67                 | 130.5                  | 4226             | –          | –          |
| Seed-6 after EVOL training | 499.42                  | 500.0                  | 80,000           | 1,600      | 10         |
| Seed-3 after initialization| 23.04                   | 19.0                   | 0                | –          | –          |
| Seed-3 after STDP-RL training| 64.14                  | 53.0                   | 8,577            | –          | –          |
| Seed-3 after EVOL training | 499.09                  | 500.0                  | 80,000           | 1,600      | 10         |

Individual neurons develop a preference for a ‘specific’ action generation after training

Comparing the activity patterns of the models while playing CartPole may not reveal developed features of the full neuronal network, possibly because of a different set of game-states evoked during each episode of CartPole. Therefore, to uncover the response properties of all network model elements, we simulated the trained models by sequentially activating all possible sensory inputs, i.e., \( 20^4 \) unique quadruplets (20 neurons to encode each of the 4 sensory parameters), and recorded the responses of all neurons as well as the associated action (i.e., if firing rate of EM-L > firing rate of EM-R, action will be Move-Left, and if firing rate of EM-R > firing rate of EM-L, action will be Move-Right) in the SNN model. During these activations, we kept the model isolated from the game to allow full control of the model’s sensory state.
Activation of each sensory neuron led to the activation of many neurons at multiple postsynaptic levels, pushing the model toward making a decision (Move-Left or -Right). Since the decisions were always based on the population level activity of motor areas, it was non-trivial to establish causality. Instead, we compared the average action selected when each sensory neuron is activated and the cascading activations throughout the rest of the network (Figure 7). For each activated sensory neuron (y-axis in Figure 7), we first counted the number of times each neuron of the network (x-axis in Figure 7) triggered during a "Move-Left" or "Move-Right" action. Further, we assigned it a Move-Left or Move-Right preference (heatmap color in Figure 7) if it was more active during Left or Right actions, respectively. The colormaps (Figure 7) show the normalized count of preferred action generation contribution of each neuron in the network, referred to as "Action selectivity" later in the text. Note that we used the negative sign (-) with "Action selectivity" to indicate "Move-Right" actions and the positive (+) sign to indicate "Move-Left" actions. Before training (Figure 7A), all the neurons in the circuit were minimally biased toward each action as "Action selectivity" values shown in the colormap are 0 (stay), marginally greater than 0.5 (for "Move-Left" action), or smaller than -0.5 (for "Move-Right" action). After training (Figures 7B–D), the preference of some neurons increased toward a single action as the range of colors in the map expanded toward +1 (for left action) and -1 for right action.

In the STDP-RL model with the worst performance (Worst Seed in Figure 7B), most of the neurons in EM-L (EM-R) participated dominantly in generating "Move-Left" ("Move-Right") actions without any clear preference for particular input features, similar to the INITIAL network (Figure 7A). For sparser input values of "Position" and "Angular Velocity," some neurons in EM-L flipped their preference slightly to "Move Right" (blue colored pixels for EM-L neurons in Figure 7B), and some neurons in EM-R flipped their preference slightly to 'Move Left' (red colored pixels for EM-R neurons in Figure 7B). Any effect on behavior due to sensory input parameters which were not used as 'critic' (e.g., "Position" and "Velocity") for training using STDP-RL were surprising. In contrast, in the STDP-RL model with the best performance (Best Seed in Figure 7C), most of the excitatory motor neurons in the circuit developed a non-overlapping association between positive angular velocity (ES neuron IDs 70–79) and "Move-Right" action, and negative angular velocity (ES neuron IDs 60–69) and "Move-Left" action. This means almost all of the neurons in EM-L were robustly activated only when the game-state had a negative angular velocity, and almost all of the neurons in EM-R were robustly activated only when the game-state had a positive angular velocity. We observed a similar developed preferences for neurons in the EVOL model (Figure 7D), except that for positive or negative angular velocity, only a subset of EM-L and EM-R were activated, i.e., not all EM-L or EM-R neurons were showing the same level of preference for the actions. In addition to angular velocity, we observed a similar effect for angle as neurons in EM-L were activated to generate 'Move-Left' actions only for positive angles as indicated by Neurons 40–50.
FIGURE 7
Action selectivity maps show differential participation of neurons in the network in action generation. In response to a particular sensory input, each neuron was considered to contribute to Move-left or Move-right actions and then labeled with the frequency of the number of times it was activated relative to the frequency of those actions (with positive values indicating activations relative to the number of moves left, and negative values indicating activations relative to moves right). (A) INITIAL, (B) STDP-RL (Worst seed), (C) STDP-RL (Best seed), and (D) EVOL. Note that propensity toward an action is defined as the number of times a neuron is active during a particular action choice. For example, if out of 8,000 sensory inputs, 6,000 times the action was Move left and 2,000 times the action was Move Right, for each of those sets of conditions, we compute how many times each neuron was active. If the neuron was active 3,000 out of the 6,000 moves left, then the participation value is 0.5. If the same neuron gets activated 2,000 times to move right, then its participation value is -1 (negative sign is for move right, positive for move left).
whereas neurons in EM-R were activated to generate “Move-Right” actions only for negative angles. The emerging preference of EM-L and EM-R neurons for positive and negative values of angles and angular velocities in the EVOL model (Figure 7D) might explain its superior performance over models trained with STDP-RL, where we observed such preference only for angular velocity (Figure 7C). Moreover, despite being not used for training, an emerging preference for ‘Angle’ and ‘Angular velocity’ in the SNN trained using EVOL showed that these two parameters were sufficient to capture sensory-motor behavior tested in this study.

For the EVOL model, we also observed a complete deactivation of some neurons (vertical white lines in Figure 7D) that might have been counterproductive in performing a correct movement. The EVOL model showed greater variability in the final total reception weight for postsynaptic neurons, including higher weight decreases that effectively deactivated neurons (Supplementary Figure 4). This neuron pruning strategy was not employed by the STDP-RL learning algorithm and might be a potential reason for the stunted game performance and the variability based on initial connections.

Training using EVOL enabled SNNs to learn broader sensory-motor associations

Plotting the activity levels of neurons during specific actions in response to each pair of sensory-inputs could diffuse the learned sensory-motor associations, especially since the model learned not about individual sensory parameters but a full game-state consisting of four sensory parameters at each game step. Therefore, we next computed sensory-motor maps for pairs of sensory input parameters (Figure 8). Before training, for only a few pairs of sensory input parameters, the model had a weak preference to move left, right, or stay (Figure 8A). After training using STDP-RL, the model with the worst performance (marked with Worst Seed in Figure 8B) developed a modest action preference for a few values of position (see the horizontal yellow line at Position: “18” for “Move Right” in Figure 8B), velocity (see horizontal yellow lines at Velocity: “14” and “16” for “Move Right”, “13” and “15” for “Move Left” heatmaps in Figure 8B) and angles (see horizontal yellow lines at “4”, “8” and “14-17” for “Move Right” and “7”, “9”, “13” and “18” for “Move Left” heatmaps in Figure 8B) but somewhat stronger action preference for positive and negative values of Angular velocity around “5 and 6” for “Move Right” and “14” for “Move Left” heatmaps (angular velocity; Figure 8B). On the other hand, the model with the best performance (marked with Best Seed in Figure 8C) developed a stronger action preference for a broader range of positive and negative values of Angular velocity around “10” for “Move Right” and “Move Left” heatmaps (see broader yellow bands on “0-9” for “Move Left” and “10-19” for “Move Right” in Figure 8C) in addition to some input values of other sensory parameters. These results are in line with Figures 7B,C, confirming that the STDP-RL trained models only learned robust action preference based on Angular velocity and could not learn from any other parameter robustly. In contrast, in models using EVOL, we found quite a few input-output associations (weak association for Position: “0”, “2”, “6-7” and “19” for “Move Left” and “1-5”, “8-18” for “Move Right” heatmaps; stronger association for Angle: “0-9” for “Move Left” and “10-19” for “Move Right” heatmaps; stronger association for Angular velocity: “0-9” for “Move Left” and “10-19” for “Move Right” heatmaps in Figure 8D), indicating distributed learned associations between actions and some values of all sensory input parameters. These results clearly indicate that better performance of models trained using EVOL could be attributed to the use of more sensory information in decision making because the model learned to take action based on two sensory signals, i.e., Angle and Angular velocity, in contrast with STDP-RL trained models, which learned to take action only based on Angular velocity. However, some of these differences could result from the STDP-RL critic explicitly using only two sensory signals: angle and angular velocity (see section “Materials and Methods”). Using more advanced critic methods for STDP-RL could produce different outcomes.

Discussion

In this work, we developed and trained an SNN using biologically inspired STDP-RL and EVOL algorithms. One of our goals was to investigate learning algorithms that operate at different timescales, and determine their strengths and weaknesses, to offer insights into how SNNs learn sensorimotor control tasks. Comparing the performance of our SNN model trained using STDP-RL and EVOL (Figures 4 vs. 5 and Table 3), we demonstrated that both strategies could train SNNs to play CartPole. There were also noticeable differences in resulting neuronal dynamics and behavior (Figures 7, 8). In addition, the EVOL showed excellent performance in training our SNNs to play CartPole (Figure 5).

As mentioned, the design of the two algorithms is substantially different in several regards. First, STDP-RL relies on an explicit hand-crafted critic, which in our case used two variables (angle and angular velocity) to inform the model of the quality of its decisions. Second, the STDP-RL critic used short intervals to evaluate the model, close to the timescale of individual moves. In contrast, EVOL evaluated the quality of a model using only the final game score, evaluated over substantially longer episodic timescales. Some of these differences are not intrinsic to the individual algorithms. However, EVOL allows for a more flexible design, and the longer timescales can offer a more accurate evaluation of a model’s quality.
It is interesting to see how training our models using these different strategies produced similar sensory-motor associations without explicitly training the models to execute specific motor actions for a given game-state. Regardless of the training algorithm, all models learned to associate actions dominantly with specific values of Angular velocity (Figures 7, 8). Analyzing the single input-action mappings (Figure 7), we did not find strong or broad associations between the Position, Angle, and Velocity with actions, which appeared in analyzing the pairwise input-action mappings (Figure 8). It is possible that the associations between sensory inputs and motor actions for other parameters were not fully revealed because the SNN was trained using all four parameter values, and some weak associations between parameters other than Angular velocity may be broadly present, but did not appear in analysis which was limited to pairwise inputs. As noted above, the STDP-RL critic also depended upon only two parameter values, potentially explaining these findings. Another reason for limited tuning for some parameters could be that during training, the receptive fields for those parameters were not experienced by the model, and therefore, the model could not learn any specific associations for those inputs. This limited tuning is indicated in the receptive fields parsed during the episodes of the game played after training (Figure 7).

Our modeling underlines the benefits of training from a set of multiple initial network configurations, achieved through varying synaptic connection weights or network architectures (Stanley and Miikkulainen, 2002; Neymotin et al., 2013), and testing the populations’ performance. The best-performing model can then be used for longer-term training.
An alternative is to use multiple models from the population and ideally promote model diversity. We highlight these issues by employing different training strategies in this work. In our STDP-RL model, we first selected a set of hyperparameters suitable for optimizing learning and then tuned for initial synaptic weights using the fixed optimal hyperparameter values. Thus, the initial weights and the learning parameter values differed from those at the beginning of the hyperparameter search. Furthermore, when we trained our SNN model using STDP-RL and a different set of initial weights (by changing random numbers), we observed variable performance, showing a large influence of the initial parameters on learning behavior. In our EVOL strategy, without using any explicit synaptic learning mechanism, we evaluated the model’s performance with different synaptic weights and found synaptic weight distributions showing good performance. Randomizing initial weights did not impact the performance as strongly as in STDP-RL trained models. Comparing the performance of models using these two training strategies (Figures 3 vs. 5), we clearly showed that EVOL produced models with robust and optimal performance. Although models trained using EVOL generated different distributions of synaptic weights, the robust peak performance was demonstrated, which is consistent with the observed variability in circuit elements of biological neuronal circuits (Bucher et al., 2005; Marder and Goaillard, 2006; Goaillard et al., 2009; Calabrese et al., 2011, 2016; Marder, 2011; Marder and Taylor, 2011; Roffman et al., 2012; Golowasch, 2014; Hamood and Marder, 2014; Anwar et al., 2022a,b).

In the past, the use of evolutionary algorithms in neurobiological models has mainly been limited to optimizing individual neurons (Van Geit et al., 2007, 2008; Rumbell et al., 2016; Neymotin et al., 2017), or neuronal networks through hyperparameter tuning (Dura-Bernal et al., 2017). Although more recent work makes changes to network architectures (Stanley et al., 2019), modifications of synaptic weight matrices in spiking or biophysical neuronal networks have rarely been performed, partly due to the large computational costs associated with searching through the high dimensional space. Here we have demonstrated that evolutionary algorithms operating on synaptic weight matrices are an effective strategy to train SNNs to perform behaviors in a dynamic environment.

We previously used the STDP-RL learning rules to train a visual/motor cortex model to play Pong (Anwar et al., 2022a). However, that model required additional complexity for encoding the visual scene (object location, motion direction). This complexity made it more challenging to decipher the role of different components/parameters of the learning algorithms and how to optimize them. In the present SNN model, the lack of a visual cortex was a simplification that allowed us to perform a more extensive hyperparameter search to increase the chances for STDP-RL to succeed. After hyperparameter optimization, the STDP-RL algorithm effectively produced good performance in CartPole. Furthermore, using CartPole and the simpler sensory/motor cortex model also allowed us to test long optimizations using EVOL in parallel on supercomputers. In the future, with the knowledge gained here, we will try our new algorithms using more complex models, tasks, and environments.

Despite the differences between ANNs and SNNs, learning in both is primarily realized by adjusting the weights of connections or synaptic strengths among interconnected neurons. In ANNs, this usually occurs through the following sequence, repeated many times: (1) inputs are sampled, (2) corresponding outputs are evaluated, and (3) weights are adjusted to minimize the output error via back-propagation through hidden network layers (Schmidhuber, 2015). In SNNs, the weights are often adjusted using hebbian or spike-timing dependent plasticity (STDP) rules (Dan and Poo, 2004, 2006; Farries and Fairhall, 2007; Izhikevich, 2007; Caporale and Dan, 2008), although more intricate learning rules could be used depending on the level of biophysical detail in the spiking neuron model (Bitnner et al., 2017). These strategies are useful when there is a temporally proximate relationship between inputs and outputs and there is no feedback involved.

Sensory-motor RL is a more difficult problem to solve because behaviors require evaluation of many sub-actions and are associated with different environmental cues integrated over time. Moreover, the reward/punishment feedback is delivered later, which makes it challenging to attribute reward/punishment only to relevant actions and neuron groups producing those actions (Izhikevich, 2007). Recent ANNs have taken advantage of a replay and update strategy to re-sample previous experiences and shape the action policy for given sensory cues maximizing the cumulative reward (Hayes et al., 2021). In SNNs using STDP-RL, eligibility traces can be used to associate reward/punishment to corresponding actions and neuronal assemblies backward in time, as we have implemented in our work here: when a postsynaptic neuron fires within a few milliseconds of presynaptic neurons firing, a synaptic eligibility trace is activated, allowing the synapse to undergo potentiation or depression during the following 1-5 seconds (Chadderdon et al., 2012; Hazan et al., 2018; Patel et al., 2019; Anwar et al., 2022a). STDP-RL trains SNNs by establishing associations between the neurons encoding the sensory environment and neurons producing actions or sequences of actions, such that appropriate actions are produced for specific sensory cues. The sensory-motor associations are established from reward-modulated synaptic weight changes at each timestep. STDP-RL trains at the individual model level, as we consider each separate initialization of an SNN network a separate “individual”.

In contrast to individual-based algorithms such as STDP-RL, evolutionary algorithms operate at vastly different timescales and typically use populations of models (Feldman et al., 1996; Parisi et al., 2019). Evolution is successful when individuals who are fit enough to produce offspring pass...
their genes to the next generation (Garrett, 2012). While individual learning is restricted to an animal's lifespan, it still confers powerful competitive advantages. Individual learning feeds into the evolutionary process: animals that learn the idiosyncrasies of their environment, including its threats and rewards, are more likely to survive and propagate. The obvious genomic storage limitations prevent the encoding of all important environmental information within an animal's genome (Zador, 2019; Koulakov et al., 2021), underlining how individual learning must complement evolution. As noted by James Mark Baldwin, learning on an individual level drives the evolutionary process (Baldwin, 1896). To simulate this process, we envision a strategy that implicitly chooses the next generation's weights and connectivity patterns based on learning success, thereby accelerating population-level fitness improvements. Such a process is consistent with the accelerated learning of models when deploying the Baldwin effect, as has been recently demonstrated in embodied intelligence tasks (Gupta et al., 2021). The extent to which these strategies translate to more complex tasks and circuit architectures could offer further insights into multi-scale neurobiological learning.

**Author's note**

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**Data availability statement**

The data presented in this study were generated using code published in online repositories. The names of the repository/repositories and accession number(s) can be found in the article/Methods.

**Author contributions**

HA, HH, and SN supervised the research. All authors performed the research, analyzed data, contributed concepts/ideas, contributed software, and wrote/edited the manuscript.

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**Conflict of interest**

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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**Supplementary material**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fncom.2022.1017284/full#supplementary-material
Lyton, W. W., Omurtag, A., Neymotin, S. A., and Hines, M. L. (2008). Just-in-time connectivity for large spiking networks. *Neural Comput.* 20, 2745–2756. doi: 10.1162/neco.2008.10-07-622

Lyton, W. W., and Stewart, M. (2006). Rule-based firing for network simulations. *Neurocomputing* 69, 1160–1164. doi: 10.1016/j.neucom.2005.12.066

Maass, W. (1996a). Noisy Spiking Neurons with Temporal Coding have more Computational Power than Sigmoidal Neurons. *Adv. Neural Inf. Proc. Syst.* 9, 211–217. doi: 10.1162/nips.1997.9.2.279

Maass, W. (1996b). Lower bounds for the computational power of networks of spiking neurons. *Neural Comput.* 8, 1–40. doi: 10.1162/neco.1996.8.1.1

Maass, W. (1997). Networks of spiking neurons: The third generation of neural network models. *Neural Netw.* 10, 1659–1671. doi: 10.1016/S0893-6080(97)00011-7

Marder, E. (2011). Variability, compensation, and modulation in neurons and circuits. *Proc. Natl. Acad. Sci. U.S.A.* 108, 15542–15548. doi: 10.1073/pnas.1010674108

Marder, E., and Goaillard, J.-M. (2006). Variability, compensation and homeostasis in neuron and network function. *Nat. Rev. Neurosci.* 7, 563–574. doi: 10.1038/nrn1949

Marder, E., and Taylor, A. L. (2011). Multiple models to capture the variability in biological neurons and networks. *Nat. Neurosci.* 14, 133–138. doi: 10.1038/nn.2735

Mnih, V., Kavukcuoglu, K., Silver, D., Rusu, A. A., Veness, J., Bellmere, M. G., et al. (2015). Human-level control through deep reinforcement learning. *Nature* 518, 529–533. doi: 10.1038/nature14236

Mostafa, H., Pedroni, S., and Cauwenberghs, G. (2017). Hardware-efficient on-line learning through pipelined truncated-error backpropagation in binary-state networks. *Front. Neurosci.* 11:496. doi: 10.3389/fnins.2017.00496

Mozafari, M., Kheradpisheh, S. R., Masquelier, T., Nowzari-Dalini, A., and Ganjtabesh, M. (2018). First-Spike-Based Visual Categorization Using Reward-Modulated STDP. *IEEE Trans. Neural Netw. Learn. Syst.* 29, 6178–6190. doi: 10.1109/TNNLS.2018.2826721

Neymotin, S. A., Chadderdon, G. L., Kerr, C. C., Francis, J. T., and Lyton, W. W. (2013). Reinforcement learning of two-joint virtual arm reaching in a computer model of sensorimotor cortex. *Neural Comput.* 25, 3263–3293. doi: 10.1162/NECO_a_00521

Neymotin, S. A., Lee, H., Park, E., Fenton, A. N., and Lyton, W. W. (2011). Emergence of physiological oscillation frequencies in a computer model of neocortex. *Front. Comput. Neurosci.* 5:19. doi: 10.3389/fncom.2011.00019

Neymotin, S. A., Suter, B. A., Dura-Bernal, S., Shepherd, G. M. G., Migliore, M., and Lytton, W. W. (2017). Optimizing computer models of corticospinal neurons to replicate in vitro dynamics. *J. Neurophysiol.* 117, 488–492. doi: 10.1152/jn.00570.2016

Parisi, G. I., Kemker, R., Part, J. L., Kanan, C., and Wermter, S. (2019). Continued lifelong learning with neural networks: A review. *Neural Netw.* 113, 54–71. doi: 10.1016/j.neunet.2019.01.012

Paton, D., Hasan, H., Saunders, D. J., Siegelmann, H., and Kozma, R. (2019). Improved robustness of reinforcement learning policies upon conversion to spiking neuronal network platforms applied to ATARI games. In arXiv [cs.LG]. arXiv. http://arxiv.org/abs/1903.11012. doi: 10.1016/j.neunet.2019.08.009

Roffman, R. C., Nories, B. J., and Calabrese, R. L. (2012). Animal-to-animal variability of connection strength in the leech heartbeat central pattern generator. *J. Neurophysiol.* 107, 1681–1693. doi: 10.1152/jn.00903.2011

Rowan, M., and Neymotin, S. (2013). "Synaptic scaling balances learning in a spiking model of neocortex," in *Adaptive and natural computing algorithms. ICANNGA 2013*. Lecture notes in computer science, Vol. 7824, eds M. Tomassini, A. Antoniou, F. Daolio and P. Buesser (Berlin: Springer). doi: 10.1007/978-3-642-37213-1_3

Rowan, M. S., Neymotin, S. A., and Lytton, W. W. (2014). Electrostimulation to reduce synaptic scaling driven progression of Alzheimer’s disease. *Front. Comput. Neurosci.* 8.39. doi: 10.3389/fncom.2014.00039

Rumbell, T. H., Dragojlik, D., Yadav, A., Hof, P. R., Luebke, J. L., and Weaver, C. M. (2016). Automated evolutionary optimization of ion channel conductances and kinetics in models of young and aged rhesus monkey pyramidal neurons. *J. Comput. Neurosci.* 41, 65–90. doi: 10.1007/s10827-016-0605-9

Salimans, T., Ho, J., Chen, X., Sidor, S., and Sutskever, I. (2017). Evolution strategies as a scalable alternative to reinforcement learning. *In arXiv [stat.ML]. arXiv. Available online at: http://arxiv.org/abs/1703.03864* (accessed September 7, 2017).

Sand, P., Skorheim, S., and Bazzhenov, M. (2017). Multi-layer network utilizing rewarded spike time dependent plasticity to learn a foraging task. *PloS Comput. Biol.* 13:e1005705. doi: 10.1371/journal.pcbi.1005705

Schmidhuber, J. (2015). Deep learning in neural networks: An overview. *Neural Netw.* 61, 85–117. doi: 10.1016/j.neunet.2014.09.003

Stanley, K. O., Clune, J., Lehman, J., and Miikkulainen, R. (2019). Designing neural networks through neuroevolution. *Nat. Mach. Intell.* 1, 24–35. doi: 10.1038/s41922-018-0006-z

Stanley, K. O., and Miikkulainen, R. (2002). Evolving neural networks through augmenting topologies. *Evol. Comput.* 10, 99–127. doi: 10.1162/1063656027621169811

Sutton, R. S., and Barto, A. G. (2018). *Reinforcement Learning*, second Edn. Cambridge, MA: MIT Press.

Tavanaei, A., Ghodrati, M., Kheradpisheh, S. R., Masquelier, T., and Maida, A. (2019). Deep learning in spiking neural networks. *Neural Netw.* 111, 47–63. doi: 10.1016/j.neunet.2018.12.002

Tavanaei, A., and Maida, A. (2017). "Bio-inspired multi-layer spiking neural network extracts discriminative features from speech signals," in *Neural information processing. ICONIP 2017*. Lecture notes in computer science, Vol. 10639, eds D. Liu, S. Xie, Y. Li, D. Zhao and E. S. El-Alfy (Cham: Springer). doi: 10.1007/978-3-319-70136-3_95

Van Geit, W., Achard, P., and De Schutter, E. (2008). Automated neuron tuning package for a wide range of electrophysiological neuron models. *Front. Neuroinf.* 1, 1. doi: 10.3389/neuro.11.001.2007

Van Geit, W., De Schutter, E., and Achard, P. (2008). Automated neuron model optimization techniques: A review. *Biol. Cybern.* 99, 241–251. doi: 10.1007/s00422-008-0257-6

Vogels, T. P., Sprekeler, H., Zenke, F., Clopath, C., and Gerstner, W. (2011). Inhibitory plasticity balances excitation and inhibition in sensory pathways and memory networks. *Science* 334, 1569–1573. doi: 10.1126/science.1211995

Zador, A. M. (2019). A critique of pure learning and what artificial neural networks can learn from animal brains. *Nat. Commun.* 10:3770. doi: 10.1038/s42003-019-11786-6