Learning the synaptic and intrinsic membrane dynamics underlying working memory in spiking neural network models

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Summary

Recurrent neural network (RNN) model trained to perform cognitive tasks is a useful computational tool for understanding how cortical circuits execute complex computations. However, these models are often composed of units that interact with one another using continuous signals and overlook parameters intrinsic to spiking neurons. Here, we developed a method to directly train not only synaptic-related variables but also membrane-related parameters of a spiking RNN model. Training our model on a wide range of cognitive tasks resulted in diverse yet task-specific synaptic and membrane parameters. We also show that fast membrane time constants and slow synaptic decay dynamics naturally emerge from our model when it is trained on tasks associated with working memory (WM). Further dissecting the optimized parameters revealed that fast membrane properties and slow synaptic dynamics are important for encoding stimuli and WM maintenance, respectively. This approach offers a unique window into how connectivity patterns and intrinsic neuronal properties contribute to complex dynamics in neural populations.

1 Introduction

2 Neurons in the cortex form recurrent connections that give rise to the complex dynamic processes 3 underlying computational functions [1–4]. Previous studies have used models based on recurrent 4 neural networks (RNNs) of continuous-rate units to characterize network dynamics behind neural 5 computations and to validate experimental findings [5–10]. However, these models do not explain 6 how intrinsic membrane properties could also contribute to the emerging dynamics.

7 Rate-based encoding of information has been reliably observed in experimental settings [8]. 8 However, recent studies demonstrated that membrane potential dynamics along with spike-based 9 coding are also capable of reliably transmitting information [11–13]. In addition, the intrinsic 10membrane properties of inhibitory neurons, including the membrane time constant and rheobase (minimum current required to evoke a single action potential), were different in two higher-order 11 cortical areas [14]. These findings strongly indicate that neuronal intrinsic properties, often ignored 12in previous computational studies employing rate-based RNNs, are crucial for better understanding 13how distinct subtypes of neurons contribute to information processing. 14

15Rate-based RNNs can be easily trained by stochastic gradient-descent to perform specified cognitive tasks [15]. However, similar supervised learning methods cannot be used to train spiking 1617RNNs due to the non-differentiable behavior of action potentials [16]. Thus, several methods introduced differentiable approximations of the non-differentiable spiking dynamics [17–20]. These stud-1819ies directly applied backpropagation to tune synaptic connections for task-specific computations. 20Other methods that do not rely on gradient computations have been also utilized to train spiking networks. One such method is based on the first-order reduced and controlled error (FORCE) 2122algorithm previously developed for rate RNNs [6]. The FORCE-based methods are capable of 23training spiking networks, but training all the parameters including recurrent connections could 24become computationally inefficient [21–23]. Lastly, recent studies successfully converted rate-based networks trained with a gradient-descent method to spiking networks for both convolutional and 25recurrent neural networks [24, 25]. Since these models are built on rate-coding networks, the result-26ing spiking models do not take advantage of the rich spiking dynamics. Moreover, these previous 27models assume that all the units in a trained network are equivalent, even though experimental 2829evidence shows that neurons in biological neural networks are highly heterogeneous. Such diversity

30 has a vital role in efficient neural coding [26].

31 Here, we present a new approach that can directly train not only recurrent synapses but also membrane-related parameters of a spiking RNN model. Our method utilizes mollifier functions [27] 32to alter the spiking dynamics to be differentiable, and a gradient-descent method is applied to tune 33 34the model parameters. These parameters are composed of synaptic parameters including recurrent connections and several important spiking-related parameters such as membrane time constant and 35 action potential threshold. Neurons with diverse and heterogeneous intrinsic parameters emerged 36 from training our spiking model on a wide range of cognitive tasks. Furthermore, we observed that 37both synaptic and spiking parameters worked in a synergistic manner to perform complex tasks 38 39 that required information integration and working memory.

40 **Results**

41 Here, we provide an overview of the method that we developed to directly train spiking recurrent 42 neural network (RNN) models (for more details see Methods). Throughout the study, we considered 43 recurrent network models composed of leaky integrate-and-fire (LIF) units whose membrane voltage 44 dynamics were governed by:

$$\tau_{m,i}\frac{dv_i}{dt} = -\left(v_i(t) - v_{\text{rest}_i}\right) + R_i I_i(t) \tag{1}$$

45 where $\tau_{m,i}$ is the membrane time constant of unit i, $v_i(t)$ is the membrane voltage of unit i at time 46 t, $v_{\text{rest},i}$ is the resting potential of unit i, and R_i is the input resistance of unit i. $I_i(t)$ represents 47 the current input to unit i at time t, which is given by:

$$I_{i}(t) = \sum_{j=1}^{N} s_{ij}(t) + I_{\text{ext}_{i}}(t)$$
(2)

48 where N is the total number of units in the network, $s_{ij}(t)$ is the synaptic input from unit j to 49 unit i at time t, and $I_{\text{ext},i}(t)$ is the external current source into unit i at time t. We used a single 50 exponential synaptic filter to model the synaptic input (s):

$$\tau_{ij} \frac{ds_{ij}}{dt} = -s_{ij}(t) + \sum_{\substack{t_j^{(k)} < t}} w_{ij} \delta(t - t_j^{(k)})$$
(3)

51 where τ_{ij} is the decay time constant of the synaptic current from unit j to unit i, w_{ij} is the synaptic 52 strength from unit j to unit i, $t_j^{(k)}$ denotes the time of the k-th action potential of unit j, and $\delta(x)$ 53 is the Dirac delta function. Once the membrane voltage of the unit i crosses its action potential 54 threshold (ϑ_i) , its membrane voltage is brought back down to its reset voltage $(v_{\text{reset},i})$.

Each LIF unit is characterized by five distinct parameters: membrane time constant $(\tau_{m,i})$, resting potential $(v_{\text{rest},i})$, input resistance (R_i) , action potential threshold (ϑ_i) , and reset potential $(v_{\text{reset},i})$. In addition, there are two trainable synaptic parameters: synaptic strength (w_{ij}) and synaptic decay time constant (τ_{ij}) from unit j to unit i.

59In order to tune all the parameters described above to produce functional spiking RNNs capable 60 of performing cognitive tasks, we employed the commonly used gradient-descent method known as backpropagation through time (BPTT; [28]) with a few important modifications. We utilized mol-61lifter gradient approximations to avoid the non-differentiability problem associated with training 62 63 spiking networks with backpropagation [27]. Furthermore, we optimized each of the model param-64eters (except for the synaptic connectivity weights) in a biologically plausible range (see Methods). 65 We also employed the weight parametrization method proposed by Song et al. to impose Dale's principle [29] (see Methods). All the spiking RNN models trained in the study used the parameter 66 67 value ranges listed in Supplementary Table 1 unless otherwise noted.

68 Units with diverse parameter values emerge after training. We applied our method to train 69 spiking networks to perform the context-dependent input integration task previously employed by Mante et al. [8]. Briefly, Mante et al. trained rhesus monkeys to flexibly integrate sensory inputs 70(color and motion of randomly moving dots presented on a screen). A contextual cue was given to 71instruct the monkeys which sensory modality (color or motion) they should attend to. The monkeys 72were required to employ flexible computations as the same modality could be either relevant or 73irrelevant depending on the contextual cue. Several previous modeling studies have successfully 7475implemented a simplified version of the task and reproduced the neural dynamics present in the 76experimental data with both continuous-rate RNNs and spiking RNNs converted from rate RNNs [25, 29, 30]. With our method, we were able to directly train the first, to our knowledge, spiking 77 RNNs with heterogeneous units whose parameters were within biologically plausible limits. 78

79 In order to train spiking RNNs to perform the input integration task, we employed a task

80 paradigm similar to the one used by previous computational studies [8, 25, 29, 30]. A recurrently 81 connected network received two streams of noisy input signals along with a constant-valued signal 82 that encoded the contextual cue (Fig. 1A). The input signals were sampled from a standard Gaus-83 sian distribution (i.e., with zero mean and unit variance) and then shifted by a positive or negative "offset" value to simulate the evidence presented in the input modalities. The network was trained 84 to produce an output signal approaching either +1 or -1 depending on the cue and the evidence 85 present in the input signal: if the cued input had a positive mean, the output signal approached 86 +1, and vice versa (Fig. 1B top). The input signal, 150 ms in duration, was given after a fixation 87 88 period (300 ms), and the network was trained to produce an output signal immediately after the 89 offset of the input signal.

We trained 20 spiking RNNs to perform the context-based input integration task. All the trainable parameters were initialized with random numbers drawn from a standard Gaussian distribution and re-scaled to the biologically plausible ranges (see Methods and Supplementary Table 1). Each network was trained until the training termination criteria were satisfied (see Methods). On average, 508.21 ± 45.96 training trials were needed for a network to meet the training termination conditions. After training, a wide distribution of the parameters emerged for both excitatory and inhibitory populations (Fig. 1C, top).

97Consistent with the previous experimental recordings from cortical neurons, the inhibitory units 98in our trained RNNs fired at a higher rate compared to the excitatory units [31]. The higher average 99 firing rates of the inhibitory units were largely due to the intrinsic properties that resulted from 100 training. Compared to the excitatory population, the inhibitory units in the trained RNNs had 101 significantly larger input resistance, smaller membrane time constants, and more depolarized resting potential (Fig. 1C; P < 0.0001, two-sided Wilcoxon rank-sum test). The action potential thresholds 102103and the reset potentials were significantly more depolarized for the inhibitory group. Furthermore, 104the time constants of the inhibitory synaptic current variable were significantly larger than the 105excitatory synaptic decay time constants (Fig. 1C).

106 Working memory requires distinct parameter distributions. The context-dependent input 107 integration task considered in the previous section did not require complex cognitive skills such as 108 working memory (WM) computations. In order to explore what parameter values are essential for

109WM tasks, we modified the paradigm to incorporate a WM component by adding a delay period 110after the delivery of the input signals. The RNN model was trained to integrate the noisy input 111 signals, sustain the integrated information throughout the 300 ms delay period, and produce an 112output signal (Fig. 1B bottom). We again trained 20 models for the modified integration task with 113the same training termination criteria (see Methods). This task required more training trials (on 114 average 1618.10 ± 345.54), but all the models were successfully trained within 2000 training trials. 115Overall, the distributions of the trained parameters were similar to those observed from the 116RNNs trained on the non-WM version of the task (Fig. 1D). The parameters that were significantly different between the two RNN models were the membrane time constant and the synaptic decay 117118time constant. The inhibitory units from the WM model displayed much faster membrane dynamics

119 and slower synaptic decay compared to the inhibitory population of the non-WM model (P < 120 0.0001, two-sided Wilcoxon rank-sum test).

121To ensure that the patterns of the trained parameters and the distinct distributions of the two 122parameters (τ_m and τ) observed from the delayed integration model were indeed associated with 123WM computations, we trained RNNs on two additional WM-related tasks: delayed matched-tosample (DMS) and delayed discrimination (DIS) tasks. For each task, we again trained 20 RNNs. 124125Both task paradigms included two sequential stimuli separated by a brief delay period. For the 126DMS task, the two input stimuli were either +1 or -1; if the two sequential had the same sign (i.e., +1/+1 or -1/-1), the network was trained to have an output signal approaching +1, 127while if the two stimuli had different signs (i.e., +1/-1 or -1/+1), the output signal approached 128-1 (Fig. 2A; see Methods). The two input stimuli for the DIS task were sinusoidal waves with 129different frequencies, modeled after the task used by Romo et al. [32] where monkeys were trained 130to discriminate two vibratory stimuli. If the first stimulus had a higher (lower) frequency, our RNN 131model was trained to produce a positive (negative) output signal (Fig. 2B; see Methods). 132

133 It took longer to train our model on these two tasks compared to the delayed integration task 134 $(7103.95 \pm 3738.65 \text{ trials for the DMS task and } 6985.47 \pm 2112.34 \text{ trials for the DIS task})$. The 135 distributions of the tuned parameters from the two WM tasks were similar to the distributions 136 obtained from the delayed integration task (Fig. 2C and D). More importantly, we again observed 137 significantly faster membrane voltage dynamics and slower synaptic decay from the inhibitory

138 units in the DMS and DIS models compared to the inhibitory units from the non-WM task. These 139 findings strongly suggest that the two parameters (τ_m and τ) of the inhibitory group contribute to 140 important dynamics associated with WM.

141Shared intrinsic properties across different working memory tasks. Prefrontal cortex and 142other higher-order cortical areas have been shown to integrate information in a flexible manner and switch between tasks seamlessly [8]. Along this line of thought, we hypothesized that the intrinsic 143144properties optimized for one WM task should be generalizable to other tasks that also require 145WM. In order to test this hypothesis, we re-trained all the RNNs that were trained in the previous 146sections to perform the DMS task without tuning the intrinsic parameters. For example, given a network trained on the non-WM integration task, we froze its intrinsic $(R, \tau_m, v_{\text{rest}}, v_{\text{reset}} \vartheta)$ along 147with the synaptic decay time constant (τ) and optimized the recurrent connections (W) only using 148149BPTT (see Methods). Therefore, each of the 20 RNNs trained for each of the four tasks (non-WM 150integration, delayed integration, DMS, and DIS tasks) was re-trained to perform the DMS task. 151As expected, the average number of trials required to successfully retrain the RNNs previously 152trained for the DMS task was low at 4408.95 \pm 3596.27 (Fig. 3A). The number of trials required to re-train the RNNs from the DIS task was also low at 4180.30 \pm 2692.81. The RNNs trained 153154for the delayed integration task took longer to re-train at 5391.85 ± 2197.99 . The non-WM RNNs required the most number of training trials to perform the DMS task (9647.55 \pm 2933.17). These 155findings indicate that the intrinsic properties from one WM model are transferable to other WM 156models. 157

158Based on these previous results, the membrane time constant (τ_m) and the synaptic decay (τ) 159variables appeared to be the two most important parameters for the transferability of WM. To test this, we repeated the re-training procedure with both τ_m and τ either fixed ("frozen") or optimized 160("tuned") for the non-WM RNNs (see Methods). For the "frozen" condition (i.e., τ_m and τ frozen 161162while the other parameters optimized), the number of trials required to re-train the non-WM RNNs 163to perform the DMS task was high and not significantly different from the number of trials it took with the intrinsic parameters fixed (Fig. 3B). On the other hand, re-tuning only τ_m and τ with the 164165other parameters fixed (i.e., "tuned" condition) resulted in a significant reduction in training time 166(Fig. 3B), suggesting that these two parameters are indeed critical for performing WM. Optimizing

167 both τ_m and τ resulted in a significant decrease in τ_m for both excitatory and inhibitory populations 168 (Fig. 3C). The synaptic decay values decreased for the excitatory units after re-tuning (Fig. 3D 169 left). For the inhibitory population, τ was significantly increased (Fig. 3D right).

170 Membrane and synaptic decay time constants critical for WM maintenance. Pyramidal 171 excitatory neurons and parvalbumin (PV) interneurons make up the majority of the neuronal cell 172 population in the cortex, and they have been shown to specialize in fast and reliable encoding of 173 information with high temporal precision [33]. To further investigate if the fast membrane and 174 slow synaptic dynamics of the units from our WM RNNs are aligned with previous experimental 175 findings and to probe how they contribute to WM maintenance, we manipulated τ_m and τ during 176 different epochs of the DMS task paradigm.

177For each of the RNNs trained from the DMS task, we first divided the population into two 178subgroups based on their τ_m values (see Methods). The short τ_m group contained units whose τ_m 179was smaller than the lower quartile value, while the long τ_m group contained units whose τ_m was 180greater than the upper quartile. During each of the four epochs (fixation, first stimulus, delay, and 181 second stimulus), we then inhibited the two τ_m subgroups separately by hyperpolarizing them and 182assessed the task performance (see Methods). As shown in Fig. 4, inhibiting the short τ_m subgroup 183during the two stimulus windows significantly impaired task performance (Fig. 4B and D), while 184disrupting the long τ_m group did not result in significant changes in task performance in all four 185task epochs.

We repeated the above analysis with two subgroups derived from a quartile split of the synaptic decay time constant (τ ; see Methods). Suppressing the synaptic connections in the long τ subgroup during the first stimulus window and the delay period significantly impaired task performance (Fig. 4B and C). Inhibiting the short τ group at any of the four epochs did not affect the task performance.

191 Therefore, the units with the fast membrane voltage dynamics (τ_m) were important for encoding 192 of stimuli, while the slow synaptic dynamics (τ) were critical for maintaining the first stimulus 193 information throughout the period spanning from the first stimulus window to the end of the delay 194 window.

195 Discussion

In this study, we presented a new method for directly training spiking RNNs with a gradient-based supervised training algorithm. Our approach allows optimizing not only the synaptic variables but also parameters intrinsic to spiking dynamics. By optimizing a wide range of parameters, we first demonstrated that units with diverse features emerged when the model was trained on a cognitive task (Figs. 1 and 2). We also showed that fast membrane dynamics combined with a slow synaptic property are critical for performing WM tasks (Figs. 3 and 4). Diversity is a basic biological principle that emerged here as a basic computational principle in spiking neural models.

203Previous modeling studies have trained RNNs to perform cognitive tasks [8, 34, 35]. Although 204some of these studies were able to train spiking RNN models, the intrinsic parameters of spiking neurons were not included as trainable variables. By using the mollifier approximation [27], we 205206developed a comprehensive framework that can tune both connectivity and spiking parameters 207using a gradient-descent method. Training spiking RNNs on multiple tasks using our method 208revealed functional specialization of excitatory and inhibitory neurons. More importantly, our 209approach allowed us to identify fast membrane voltage dynamics as an essential property required 210to encode incoming stimuli robustly for WM tasks.

211 Previous computational studies employing RNNs assumed that all the units in a network shared the same intrinsic parameters and optimized only synaptic connectivity patterns during training. 212213Recent studies developed models that give rise to units with heterogeneous intrinsic properties. 214For example, a new activation function that is tunable for each neuron in a network was recently 215proposed [36]. In addition, we recently trained synaptic decay time constants in a rate RNN model 216[25]. Although these methods produce heterogeneous units, they do not incorporate parameters 217inherent to spiking mechanisms. Our method not only allows direct training of synaptic weights 218of spiking RNNs that abide by Dale's principle, but also enables training of synaptic and intrinsic 219membrane parameters for each neuron.

Although our method was successful at training spiking RNNs with biological constraints, the gradient-based method employed in the present study is not biologically plausible. In cortical neural networks, local learning rules, such as spike-timing-dependent plasticity (STDP), were observed, but the gradient-descent algorithm used in our method is neither local to synapses nor local in time [16]. However, this non-locality allowed our method to train intrinsic membrane and connectivity parameters, creating biologically plausible neural architectures that solve specified problems. The learning algorithm for spiking neurons makes it possible to uncover neural dynamics hidden in experimental data [8, 29, 37], thus emphasizing that a biologically realistic model can be constructed by non-biological means.

229Another limitation of our framework arises from our spiking neuron model. Although we were 230able to train models with heterogeneous neurons the leaky integrate-and-fire model used in the 231present study can only capture the dynamics of fast-firing neurons due to the lack of adaptation 232[38]. In particular, several other types of neurons, such as regular-firing and bursting neurons, 233are also common in cortical networks [39]. Applying our method to spiking neuron models with 234adaptation currents, such as those in Hodgkin-Huxley models model [40] and adaptive exponential integrate-and-fire model [41], will be an interesting next step to further investigate the role of 235236neurons from various firing classes in information processing.

In summary, we provide a novel approach for directly training both connectivity and membrane parameters in spiking RNNs. Training connectivity and intrinsic membrane parameters revealed distinct populations only identifiable by their parameter values, thus enabling investigation of the roles played by specific populations in the computation processes. This lays the foundation for uncovering how neural circuits process information with discrete spikes and building more powerefficient spiking networks.

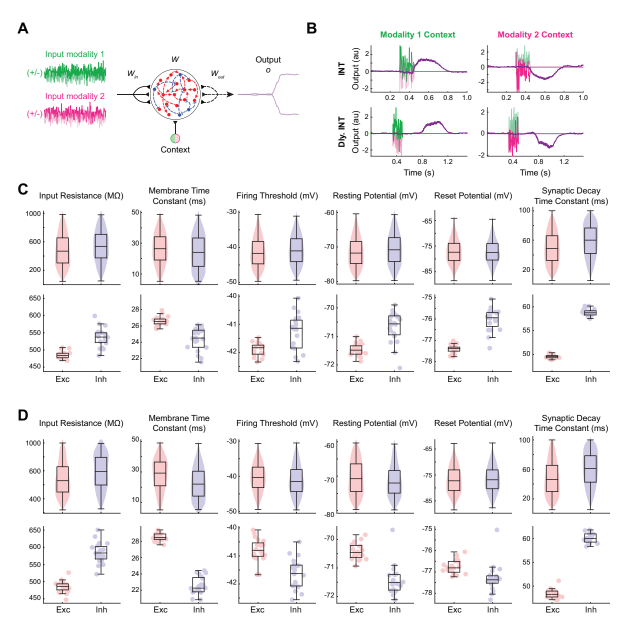


Fig. 1 | Biologically realistic spiking network performing a context-dependent input integration task. (A) Schematic diagram of the RNN model trained for the context-dependent integration task. Two streams of noisy input signals (green and magenta lines) along with a context signal were delivered to the LIF RNN. The network was trained to integrate and determine if the mean of the cued input signal (i.e., cued offset value) was positive ("+" choice) or negative ("-" choice) without or with a delay period at the end of the noisy input signals. (B) Example input and output signals from example RNNs trained to perform the task without (top row; INT) or with a delay period (bottom row; Dly. INT). (C) Distributions of the optimized parameters for the excitatory (red) and inhibitory (blue) units across all 20 models trained for the INT task. Top, distributions pooled from all the units from 20 models. Bottom, each dot represents the average value from one network. (D) Distributions of the optimized parameters for the excitatory (red) and inhibitory (blue) units across all 20 models trained for the Dly. INT task. Top, distributions pooled from all the units from 20 models. Bottom, each dot represents the average value from one network. Bottom, each dot represents the average value from 20 models. Bottom, each dot represents the average value from one network.

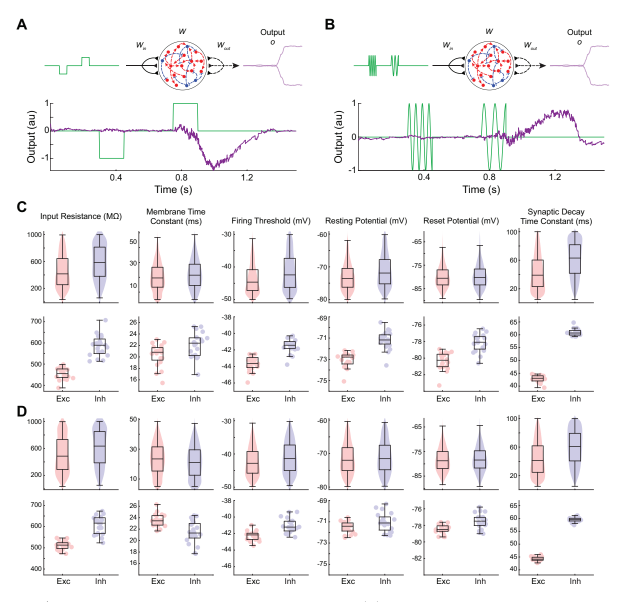


Fig. 2 | RNNs trained for two additional WM tasks. (A) Schematic illustrating the task paradigm for the delayed match-to-sample (DMS) task (top) and input and output signals from an example trained RNN (bottom). (B) Schematic illustrating the task paradigm for the delayed discrimination (DIS) task (top) and input and output signals from an example trained RNN (bottom). (C) Distributions of the optimized parameters for the excitatory (red) and inhibitory (blue) units across all 20 models trained for the DMS task. Top, distributions pooled from all the units from 20 models. Bottom, each dot represents the average value from one network. (D) Distributions of the optimized parameters for the excitatory (red) and inhibitory (blue) units across all 20 models trained for the DIS task. Top, distributions pooled from all the units from 20 models. Bottom, each dot represents the average value from 20 models. Bottom, each dot represents the average value from all the units from 20 models. Bottom, each dot represents the units from 20 models. Bottom, each dot represents the units from 20 models. Bottom, each dot represents the units from 20 models. Bottom, each dot represents the units from 20 models. Bottom, each dot represents the units from 20 models. Bottom, each dot represents the average value from one network.

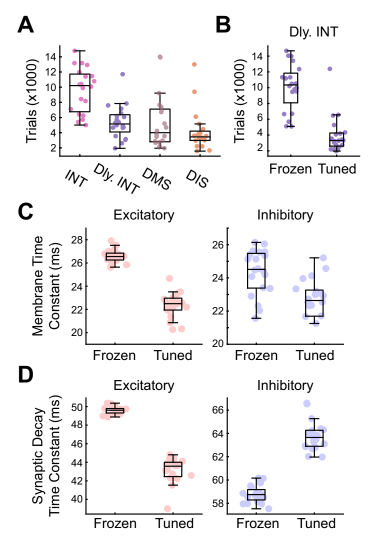


Fig. 3 | Retraining RNN models to perform the DMS task. (A) Number of training trials required to retrain the models previously trained for the INT, Dly. INT, DMS, or DIS tasks to perform the DMS task. (B) Number of training trials required to retrain the Dly. INT RNNs to perform the DMS task with the membrane time constant (τ_m) and synaptic decay time constant (τ) frozen or tuned. (C) Distribution of the membrane time constant values for the excitatory (red) and inhibitory (blue) units for the two conditions (frozen and tuned). Each dot represents the average value from one network. (D) Distribution of the synaptic decay time constant values for the excitatory (red) and inhibitory (blue) units for the two conditions (frozen and tuned). Each dot represents the average value from one network.

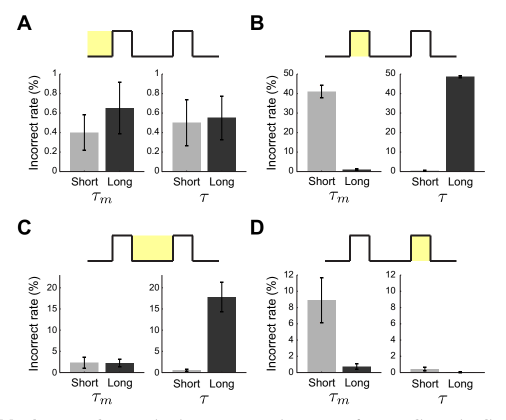


Fig. 4 | Membrane and synaptic time constants important for encoding stimuli and WM maintenance. (A–D) DMS task performance when short τ_m , long τ_m , short τ , or long τ units were inhibited during the fixation (A), first stimulus window (B), delay period (C), or second stimulus window (D).

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339 Author contributions

Y.L., R.K., and T.J.S. designed the study and wrote the manuscript. Y.L. performed the analysesand simulations.

342 Declaration of interests

343 The authors declare no competing interests.

344 Methods

345 Spiking network structure and discretization. Our spiking RNN model consisted of N integrate346 and-fire (LIF) units is governed by

$$\tau_{m,i}\frac{dv_i}{dt} = -\left(v_i(t) - v_{\text{rest}_i}\right) + R_i I_i(t) + \xi \tag{4}$$

347 where $\tau_{m,i}$ is the membrane time constant of unit i, $v_i(t)$ is the membrane voltage of unit i at 348 time t, $v_{\text{rest},i}$ is the resting potential of unit i, and R_i is the input resistance of unit i, and ξ is the 349 membrane voltage spontaneous fluctuation. $I_i(t)$ represents the current input to unit i at time t, 350 which is given by:

$$I_{i}(t) = \sum_{j=1}^{N} s_{ij}(t) + I_{\text{ext}_{i}}(t)$$
(5)

351 where N is the total number of units in the network, $s_{ij}(t)$ is the filtered spike train of unit j to 352 unit i at time t, and $I_{\text{ext}_i}(t)$ is the external current source into unit i at time t. For this study, 353 N = 400 for all tasks and networks trained.

354 The external current $I_{ext}(t)$ encodes the task-specific input at time t:

$$\boldsymbol{I}_{\text{ext}}(t) = W_{\text{in}}\boldsymbol{u}(t) \tag{6}$$

where the time-varying stimulus signals $\boldsymbol{u}(t) \in \mathbb{R}^{N_{\text{in}} \times 1}$ are fed into the network via $W_{\text{in}} \in \mathbb{R}^{N \times N_{\text{in}}}$, which can be viewed as presynaptic connections to the network that convert analog input into firing rates. N_{in} corresponds to the number of channels in the input signal.

358 We used a single exponential synaptic filter:

$$\tau_{ij}\frac{ds_{ij}}{dt} = -s_{ij}(t) + \sum_{\substack{t_j^{(k)} < t}} w_{ij}\delta(t - t_j^{(k)})$$
(7)

359 where τ_{ij} is the synaptic decay time constant from unit j to unit i, w_{ij} is the synaptic strength 360 from unit j to unit i, $t_j^{(k)}$ denotes the time of the k-th action potential of unit j, and $\delta(x)$ is the 361 Dirac delta function. Once the membrane voltage of the unit i crosses its action potential threshold 362 (ϑ_i) , its membrane voltage is brought back down to its reset voltage $(v_{\text{reset},i})$.

363 The output of our spiking model at time t is given by

$$o(t) = W_{\text{out}}\boldsymbol{r}(t) \tag{8}$$

364 where $W_{\text{out}} \in \mathbb{R}^{1 \times N}$ are the readout weights, and $\boldsymbol{r}(t) \in \mathbb{R}^{N \times 1}$, which can be interpreted as the 365 firing rate of units, are given by

$$\tau_{r,i} \frac{dr_i}{dt} = -r_i(t) + \sum_{\substack{t_i^{(k)} < t}} \delta(t - t_i^{(k)})$$
(9)

366 where $\tau_{r,i}$ is the synaptic decay time constant of firing rate estimate for unit *i*.

367 We converted the continuous-time differential equations to discrete-time iterative equations 368 and used numerical integration (Euler's method) to solve the equations. The membrane voltage 369 $\boldsymbol{v} \in \mathbb{R}^{1 \times N}$ at step n + 1 is given by

$$\boldsymbol{v}^{(n+1)} = \tilde{\boldsymbol{v}}^{(n)} + \frac{\Delta t}{\boldsymbol{\tau}_m} \left(-\left(\tilde{\boldsymbol{v}}^{(n)} - \boldsymbol{v}_{\text{rest}} \right) + \boldsymbol{I}^{(n+1)} \odot \boldsymbol{R} \right) + c \mathcal{N}(0, \Delta t)$$
(10)

370 where Δt is the sampling rate (or step size), which was set $\Delta t = 1$ ms for this study, $\boldsymbol{\tau}_m \in \mathbb{R}^{1 \times N}$ is 371 the membrane time constant, $\boldsymbol{v}_{\text{rest}} \in \mathbb{R}^{1 \times N}$ is the resting potential, \odot refers to Hadamard operation 372 (element-wise multiplication), $\dot{-}$ refers to the element-wise division, and $\boldsymbol{R} \in \mathbb{R}^{1 \times N}$ is the input 373 resistance. The term $c\mathcal{N}(0, \Delta t)$ injects spontaneous membrane fluctuations, where $\mathcal{N}(0, \Delta t) \in$ 374 $\mathbb{R}^{1 \times N}$ is a Gaussian random vector consisting of N independent Gaussian random variables with 375 mean 0 and variance Δt , and c is the scaling constant for the amplitude of fluctuations, set as c = 5376 throughout the study.

There are two time-varying terms in Eq. 10, the membrane voltage after reset $(\tilde{\boldsymbol{v}}^{(n)})$ and input current $(\boldsymbol{I}^{(n+1)})$. The voltage reset in the LIF model after action potentials at step n is formulated as

$$\tilde{\boldsymbol{v}}^{(n+1)} = \boldsymbol{v}^{(n+1)} + \left(\boldsymbol{v}_{\text{reset}} - \boldsymbol{v}^{(n+1)}\right) \odot H\left(\boldsymbol{v}^{(n+1)} - \boldsymbol{\vartheta}\right)$$
(11)

380 where $\boldsymbol{v}_{\text{reset}} \in \mathbb{R}^{1 \times N}$ is the reset potential, $\boldsymbol{\vartheta} \in \mathbb{R}^{1 \times N}$ is the action potential thresholds, and H(x) is 381 the element-wise Heaviside step function. The term $H\left(\boldsymbol{v}^{(n+1)} - \boldsymbol{\vartheta}\right)$ represents the spiking output

382 activities at step n + 1. The input current at step n + 1 is given by

$$\boldsymbol{I}^{(n+1)} = S^{(n)} \cdot \boldsymbol{1} + W_{\text{in}} \boldsymbol{u}^{(n+1)}$$
(12)

383 where $\mathbf{1} \in \mathbb{R}^{1 \times N}$ is the column vector with all ones and $S^{(n)}$ is the filtered spike train matrix at 384 step n, which follows the iteration

$$S^{(n)} = S^{(n-1)} + \frac{\Delta t}{T} \left(-S^{(n-1)} + W \odot H \left(\boldsymbol{v}^{(n)} - \boldsymbol{\vartheta} \right) \right)$$
(13)

385 where $T \in \mathbb{R}^{N \times N}$ is the matrix of synaptic decay time constants and $W \in \mathbb{R}^{N \times N}$ is the matrix of 386 synaptic strengths. Here, $W \in \mathbb{R}^{N \times N}$ is a matrix and $H(\boldsymbol{v}^{(n)} - \boldsymbol{\vartheta}) \in \mathbb{R}^{1 \times N}$ is a row vector. The 387 notation $A \odot \boldsymbol{v}$ refers to element-wise multiplication of matrix A row by row with the row vector \boldsymbol{v} . 388 The output at step n + 1 is computed by

$$o^{(n+1)} = W_{\text{out}} \boldsymbol{r}^{(n+1)} \tag{14}$$

389 in which

$$\boldsymbol{r}^{(n+1)} = \boldsymbol{r}^{(n)} + \frac{\Delta t}{\boldsymbol{\tau}_r} \left(-\boldsymbol{r}^{(n)} + H\left(\boldsymbol{v}^{(n+1)} - \boldsymbol{\vartheta}\right) \right)$$
(15)

390 where $\boldsymbol{\tau}_r \in \mathbb{R}^{1 \times N}$ is the synaptic decay time constants of firing rate estimate.

391 **Training details.** In this study, we only used the supervised backpropagation of errors learning 392 algorithm. The loss function (\mathcal{L}) is defined in terms of the root mean square error (RMSE) with 393 respect to a task-specific target signal (z) and the network output signal (o):

$$\mathcal{L} := \sqrt{\left(\sum_{n=1}^{M} \left(z^{(n)} - o^{(n)}\right)^2\right)} \tag{16}$$

394 where M is the total time steps.

We used Adaptive Moment Estimation (ADAM) stochastic gradient descent algorithm [42] with mini-batch training. The mollifier gradient approximations were employed to address nondifferentiability problem associated with the spiking process (see Mollifier gradient approximations). The learning rate was set to 0.01, the batch size was set to 10, and the first and

second moment decay rates were 0.9 and 0.999, respectively. The trainable parameters include in-400 put weights (W_{in}) , synaptic strengths (W), readout weights (W_{out}) , synaptic decay time constants 401 (T), membrane time constants $(\boldsymbol{\tau}_m)$, input resistances (\boldsymbol{R}) , resting potentials (\boldsymbol{v}_{rest}) , reset voltages 402 (\boldsymbol{v}_{reset}) , action potential thresholds $(\boldsymbol{\vartheta})$, and synaptic decay time constants for firing rate estimates 403 $(\boldsymbol{\tau}_r)$.

A nonlinear projected gradient method was used to constrain parameters within the biologically realistic ranges described in Supplementary Table 1. A linear projection map forces some solutions to be projected on the boundary. That is, there are always some units whose parameters take the min and max values of the constraint. On the other hand, a nonlinear projection guarantees that no values are on the boundary almost surely, a more realistic situation to consider. Specifically, to bound a parameter p at iteration i + 1 into the range $[p_{min}, p_{max}]$, we have

$$\tilde{p}_{i+1} = \sigma(p_{i+1}) \cdot (p_{\max} - p_{\min}) + p_{\min} \tag{17}$$

410 where \tilde{p}_{i+1} is the projected solution of parameter p at iteration i + 1, p_{i+1} is the unconstrained 411 solution given by the gradient descent algorithm at iteration i + 1, p_{max} and p_{min} are the maximum 412 and minimum values of parameter p, and $\sigma(x)$ is the sigmoid function, defined as

$$\sigma(x) := \frac{1}{1 + \exp(-x)} \tag{18}$$

413 We initialized all parameters, except the input weights (W_{in}) , as samples from the standard 414 Gaussian distribution with zero mean and unit variance, whereas the input weights were drawn 415from Gaussian distribution with zero mean and variance 400. This is because our input signals were bounded within the range [-1, 1], insufficient to bring the membrane voltage from the resting 416417 potential above the action potential threshold. Hence, to accelerate training, it was necessary to 418 make sure units were excited by the input signals in the first place. The synaptic strength matrix 419 (W) was also initialized sparse, with the percentage of connectivity being only 20%. We say the 420 network successfully did the task if the output signal hits above +0.8 (or below -0.8) if the target 421 output is +1 (or -1). We stopped training when the loss (\mathcal{L}) is less than 15 and the accuracy over 422100 trials is above 95%.

423 The method proposed by Song et al. [29] was used to *impose Dale's principle* with separate 424 excitatory and inhibitory populations. The synaptic connectivity matrix (W) in the model was 425 parametrized by

$$W_{i+1} = [W_{i+1}]_+ \cdot D \tag{19}$$

where \tilde{W}_{i+1} is the resulted matrix that encoded separate populations at update step $i + 1, W_{i+1}$ 426is the solution given by the gradient descent algorithm at step i + 1, and $[\cdot]_+$ is the rectified 427428linear unit (ReLU) operation applied at the end of each update step. The ReLU operation is to 429 ensure that entries of the matrix are always non-negative before multiplied by the matrix D, as the negative weight connections update from gradient descent are pruned by the end of each update. 430The diagonal matrix $(D \in \mathbb{R}^{N \times N})$ encode +1 for excitatory units and -1 for inhibitory units. 431432 The value of matrix (D) was randomly assigned before training according to a preset proportion 433 between inhibitory and excitatory units, and the value D was fixed through the whole training 434process. The I/E units proportion in this study was 20% to 80%.

435 In order to capture the biologically realistic dynamics of SNNs, the temporal resolution (Δt) was set to be no longer than the duration of absolute refractory period to ensure that the spiking 436437activities are not affected by the numerical integration process. Therefore, we set $\Delta t = 1$ ms during 438training. Due to the vanishing gradient problem occurring in training RNNs [43], with $\Delta = 1$ ms, it is impossible to train tasks with duration longer than 1 second (i.e., M > 1000). It is notable 439 that in the above formulation, only membrane time constant (τ_m) and synaptic time decay (τ) are 440 dependent on the sampling rate (Δt ; Eq. 10 and Eq. 13). Hence, after the models are trained, 441 442we can make sampling rate (Δt) smaller (i.e., having finer temporal resolution) while still keeping 443the same dynamics of the trained networks. Increasing Δt by a factor is equivalent to decreasing τ 444 and τ_m altogether by the same factor, as τ and τ_m are inversely proportional to Δt in Eq. 10 and Eq. 13. Hence, to train a network performing tasks with duration longer than 1 second, we need 445to make the temporal resolution coarser (i.e., increasing Δt by a factor s) so that with the same 446 trainable range of time steps (i.e., a fixed $M \leq 1000$), the duration of task becomes longer by the 447 same factor s. This "decrease in temporal resolution" can be interpreted as shortening τ and τ_m 448instead of an actual decrease in temporal resolution. Applying this trick enables us to train tasks 449450with arbitrary duration by re-scaling the ranges of τ and τ_m into a smaller one while still making

451 the spiking activities biologically realistic. In practice, we simply scaled down τ and τ_m by a factor 452 s = 3 with a fixed number of time steps (M), and later during the testing stage, we re-scaled M, τ 453 and τ_m up by the same factor s.

454Mollifier gradient approximations. In the above formulation, the Heaviside step function H(x)is not continuous. As a result, the loss function \mathcal{L} is not differentiable. This poses the major problem 455when applying the traditional backpropagation algorithm for training neural networks, because the 456457backpropagation algorithm uses gradient descent methods that require the function being minimized to be differentiable, or at least to be continuous. However, the derivative of Heaviside step function 458459H(x) is Dirac Delta function $\delta(x)$, which is 0 everywhere except at 0, where the function value is 460 ∞ . It is difficult to use this derivative for the gradient descent methods because the value of the 461gradients is 0 almost everywhere.

To address the discontinuity problem, we employed mollifier gradient method proposed by 463 Ermoliev et al. [27]. The method can be applied to any strongly lower semicontinuous functions 464 to find local minima following an iterative gradient descent in which the gradients change over 465 iterations based on averaged functions derived from the original objective function. The family of 466 averaged functions f_{ε} of function f is defined by convolution of f with a mollifier: ψ_{ε}

$$f_{\varepsilon}(x) := \int_{\mathbb{R}^n} f(x-z)\psi_{\varepsilon}(z)dz = \int_{\mathbb{R}^n} f(x)\psi_{\varepsilon}(x-z)dz = f * \psi_{\varepsilon}(x)$$
(20)

467 where $\psi_{\varepsilon} \in \{\psi_{\varepsilon} : \mathbb{R}^n \to \mathbb{R}_+, \varepsilon > 0\}$, a family of compactly supported (generalized) functions named 468 *mollifiers* that satisfy

$$\int_{\mathbb{R}^n} \psi_{\varepsilon}(x) \, dx = 1, \quad \lim_{\varepsilon \to 0} \psi_{\varepsilon}(x) = \lim_{\varepsilon \to 0} \varepsilon^{-n} \psi_{\varepsilon}(x/\varepsilon) = \delta(x) \tag{21}$$

It was shown that for any strongly lower semicontinuous functions f, the averaged functions f_{ε} epi-converge to f as $\varepsilon \to 0$, a type of convergence that preserves the local minima and minimizers. Therefore, it is possible to use the gradients of averaged functions to minimize the original lower semicontinuous functions and find the local minima. We used the conventional family of mollifiers

473 obtained by normalizing a probability density function ψ :

$$\psi_{\varepsilon}(z) := \frac{\psi(z/\varepsilon)}{\varepsilon^n} \tag{22}$$

474 In our case, n = 1 as the domain of H(x) is the real line:

$$H_{\varepsilon}(x) := \frac{1}{\varepsilon} \int_{-\infty}^{\infty} H(x-z) \,\psi(z/\varepsilon) \,dz \tag{23}$$

475 For any $\varepsilon > 0$, the gradient of $H_{\varepsilon}(g(x))$ with respect to parameter p is given by

$$\nabla_p H_{\varepsilon}(g(x)) = \frac{1}{\varepsilon} \psi(g(x)/\varepsilon) \nabla_p g(x) = \psi_{\varepsilon}(g(x)) \nabla_p g(x)$$
(24)

where ψ is some symmetric density function and g(x) is any function with \mathbb{R} as its codomain. Since our goal was not to find a local minimum x^* that satisfies the optimality condition $\lim_{\varepsilon \to 0} ||\nabla f_{\varepsilon}(x^*)|| = 0$ as defined by Ermoliev et al., but rather to minimize the loss function for its value to be sufficiently small so that the network can perform the task correctly, we did not vary the gradients during the minimization process. Instead, we fixed an approximation of the gradient and used the approximation throughout the training process. We chose the normalized box function, i.e., the density function of uniform distribution $\mathcal{U}(-\varepsilon/2, \varepsilon/2)$, as the kernel,

$$\psi(x) := \begin{cases} \frac{1}{\varepsilon} & \text{for } x \in [-\varepsilon/2, \varepsilon/2] \\ 0 & \text{otherwise} \end{cases}$$
(25)

483 and fixed $\varepsilon = 5$.

We found no difference in the trained models with different choices of ε , as long as the value was large enough to keep the gradients active so that gradients did not vanish through time steps. There was also no difference between models trained with fixed ε and those trained with the original scheme in Ermoliev et al. where $\varepsilon \to 0$ as the number of iterations increases. The purpose for fixing the value of ε was to compare the training epochs (iterations) among the retraining paradigms (see Fig. 3) with the same gradient.

490 **Re-training models for DMS task.** To test whether intrinsic properties optimized for one WM 491 task are generalizable to other tasks that also require WM, we re-trained our models to perform 492 the DMS task with all intrinsic properties fixed. In contrast to the training paradigm described 493 in the previous sections, the trainable parameters for re-training only include input weights (W_{in}) , 494 synaptic strengths (W), and readout weights (W_{out}) . Each of the 20 RNNs trained for each of the 495 four tasks (non-WM integration, delayed integration, DMS, and DIS tasks) used in this study was 496 re-trained to perform the DMS task.

497To test whether synaptic decay time constants (τ) and membrane time constants (τ_m) are the 498most crucial parameters for transferability of WM tasks, we repeated the re-training procedure with both τ_m and τ either fixed or optimized for the non-WM RNNs. The RNNs optimized to 499500perform the context-based input integration task were used for re-training under two schemes: the 501tuned scheme and the frozen scheme. For the tuned scheme, the trainable parameters include input 502weights (W_{in}) , synaptic strengths (W), readout weights (W_{out}) , synaptic decay time constants (T), membrane time constants (τ_m), and synaptic decay time constants for firing rate estimates (τ_r). 503504For the frozen scheme, the trainable parameters include input weights (W_{in}) , synaptic strengths (W), readout weights (W_{out}) , input resistances (R), resting potentials (v_{rest}) , reset voltages (v_{reset}) , 505506and action potential thresholds (ϑ) .

Units function analysis. For Fig. 4, we manipulated τ_m and τ during different epochs of the 507508DMS task paradigm to investigate if fast membrane and slow synaptic dynamics are responsible for 509WM maintenance. For each of the RNNs trained from the DMS task, we first divided the population into two subgroups based on their τ_m values. The short τ_m group contained units whose τ_m was 510smaller than the median value of τ_m of all units in the RNN, while the long τ_m group contained 511512units whose τ_m was greater than the median value. The average median value of τ_m across all 20 513models was 19.64 ± 2.45 ms. During each of the four epochs (fixation, first stimulus, delay, and 514second stimulus), we inhibited the two τ_m subgroups separately by hyperpolarizing them and then assessed the task performance. The hyperpolarization was done by setting the membrane voltage 515v = -100 mV for the intended subgroup of units. Similar to the training stage, we say that the 516network successfully did the task if the output signal hits above +0.8 (or below -0.8) if the target 517output is +1 (or -1). If the target output is between -0.8 and +0.8, the network is considered 518

519 having no response. If the output signal is above +0.8 (or below -0.8) while the target output is 520 -1 (or +1), we say that the network gives an incorrect response.

We conducted a similar analysis based on two subgroups of synapses derived from a quartile split of synaptic decay time constant (τ). The short τ group contained synapses whose τ was smaller than the 25th percentile of all τ in the RNN, while the long τ group contained synapses whose τ was greater than the 75th percentile. The average 25th percentile across all 20 models was 25.36 \pm 2.40 ms, and the average 75th percentile was 66.18 \pm 1.17 ms. The targeted subgroup of synapses was suppressed by setting the connection strength w = 0 during each of the four epochs of DMS task.

528 Code availability

529 The implementation of our framework and the codes to generate all the figures in this work are 530 available at https://github.com/y-inghao-li/SRNN/

531 Data availability

532 The trained models used in the present study are available as MATLAB-formatted data at https:

533 //github.com/y-inghao-li/SRNN/

Parameter name	Symbol	Minimum	Maximum
Input resistance	R	$5 M\Omega$	1000 M Ω
Membrane time	τ	$5 \mathrm{ms}$	$50 \mathrm{ms}$
constant	$ au_m$		
Action potential	19	$-50 \mathrm{mV}$	$-30 \mathrm{mV}$
threshold	0		
Resting potential	$v_{\rm rest}$	-80 mV	-60 mV
Reset voltage	v_{reset}	$v_{\rm rest}-10~{\rm mV}$	$v_{\rm rest} - 1 \ {\rm mV}$
value			
Synaptic decay time	au	$5 \mathrm{ms}$	$100 \mathrm{\ ms}$

Supplementary Table

Supplementary Table 1: Parameter values used for this study. To keep the constraint $v_{\text{rest}} > v_{\text{reset}}$, we trained the afterhyperpolarization (AHP) potential with range from -10 mV to -1 mV, so the value of v_{reset} is dependent upon the value of v_{rest} .