
Transformer needs NMDA receptor nonlinearity for long-term memory

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Abstract

The NMDA receptor (NMDAR) in the hippocampus is essential for learning and memory. We find an interesting resemblance between deep models' nonlinear activation function and the NMDAR's nonlinear dynamics. In light of a recent study that compared the transformer architecture to the formation of hippocampal memory, this paper presents new findings that NMDAR-like nonlinearity may be essential for consolidating short-term working memory into long-term reference memory. We design a navigation task assessing these two memory functions and show that manipulating the activation function (i.e., mimicking the Mg^{2+} -gating of NMDAR) disrupts long-term memory formation. Our experimental data suggest that the concept of place cells and reference memory may reside in the feed-forward network and that nonlinearity plays a key role in these processes. Our findings propose that the transformer architecture and hippocampal spatial representation resemble by sharing the overlapping concept of NMDAR nonlinearity.

1 Introduction

In the hippocampus, NMDAR is regarded as an essential component that mediates synaptic plasticity, memory formation, and spatial representation of place cells [1, 2, 3]. It has unique nonlinear dynamics which is modulated by Mg^{2+} -gating [4, 5], serving as a switch for synaptic plasticity and long-term memory formation [6, 7, 8] (Fig. 1a). This work is inspired by 1) the fascinating resemblance of NMDAR with the nonlinear GELU activation function that is widely used in the feed-forward networks of modern transformer architectures (Fig. 1c) [9, 10, 11] and 2) recent models relating transformer's self-attention mechanism to hippocampal formation [12, 13]. These findings motivated us to ask a question; can NMDAR-like nonlinearity in the feed-forward network of transformers enhance the long-term memory formation and spatial place cell representation?

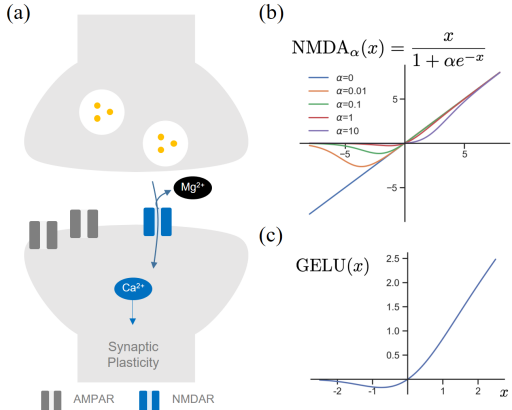


Figure 1: (a) Schematic diagram of Mg^{2+} -gated NMDAR modulating synaptic plasticity. (b) Mg^{2+} -gated NMDAR-like activation function. (c) Gaussian Error Linear Unit (GELU) activation function in transformer's feed-forward layers.

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To address this question, we design a spatial navigation task in a 2D grid environment that assesses two different memory types in neuroscience [14, 15]: working memory and reference memory. Working memory controls the events from a within-trial, while reference memory controls across-trials from the unchanging environment. Our experimental data suggest that NMDAR-like nonlinearity in feed-forward networks of the transformer can enhance the reference memory formation and place cell representation.

2 Methods

Relating activation function in transformers with NMDAR nonlinearities NMDAR’s nonlinear dynamics arises from the voltage-gated Mg^{2+} repulsion at the NMDAR channel’s pore [4, 5] (Fig. 1a). Previously, Mg^{2+} -gated NMDAR open probability p has been shown to follow ion blockade model of Woodhull [16]:

$$p_\alpha(x) = \frac{1}{1 + \alpha e^{-\beta x}}, \quad (1)$$

where x represent an input voltage, $\alpha = [Mg^{2+}] / K_{Mg^{2+}}$ is a parameter determined by $[Mg^{2+}]$, $K_{Mg^{2+}}$ is a dissociation constant, and β is a temperature constant. As experimentally shown, increasing the Mg^{2+} level in the brain can enhance long-term memory formation [7]. We observed the NMDAR’s nonlinear dynamics of the *IV* curve (current-voltage relationship) in the synapse to closely resemble the form of the GELU activation function. GELU is a widely used activation function in transformers (Fig. 1c; $GELU(x) \approx x\sigma(1.702x)$ where σ is the sigmoid function) [9, 10, 11]. Inspired by this resemblance, we define a new nonlinear activation function (Fig. 1b) with α parameter which modulates dynamics as follows:

$$NMDA_\alpha(x) = x p_\alpha(x) = \frac{x}{1 + \alpha e^{-x}}. \quad (2)$$

To investigate this NMDAR-like nonlinearity in transformer memory formation, we replaced the $GELU(x)$ activation function with $NMDA_\alpha(x)$ in a standard transformer model.

Transformers learn spatial navigation tasks

We train the transformer model to predict the subsequent sensory observations of an agent that randomly walks a 2D grid environment [13] (Fig. 2a). A sequence of previous [Action (a), Observation (x)] pairs is an input to the model, and the subsequent observation is masked for prediction (Fig. 2b). Instead of using sinusoidal positional encoding [17] that is commonly used in transformers, we employ the recurrent positional embedding which is encoding the location of an input element by using the recurrent neural network (RNN) [13]³.

We generate the embedding vectors of the sensory observation sequence with a word embedding layer, but the embedding vectors of the action sequence are generated by RNN: $e_{t+1} = \tanh(e_t W_a)$, where e_t is a recurrent positional embedding at step t , and W_a is the action-dependent trainable weight matrix. The input is given by $\{[e_1, x_1], [e_2, x_2], \dots, [e_t, x_t]\}$, where x denotes the embedding vector of sensory observation x ; the initial recurrent positional embedding e_1 is sampled from a normal distribution and we mask the last observation x_t . We generate N maps of 11×11 2D grids. A random sensory observation among ten letters is placed at each position on each map. Agents can move ‘up,’ ‘right,’ ‘down,’ ‘left,’ or ‘stay.’ An agent starts at a random position and initiates a random walk on the map, a randomly selected map among N training maps, for 2,048 steps for each trial.

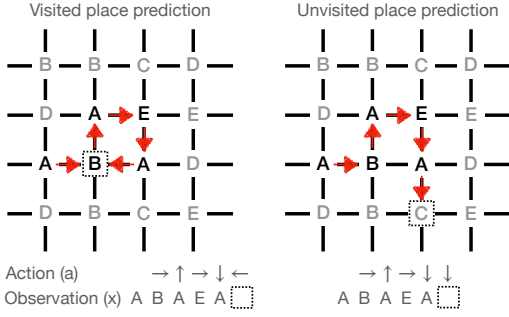


Figure 2: Sensory observation prediction task in a 2D grid, where dotted squares indicate the target position to predict given a sequence of past actions and observations. Gray (black) letters represent the unvisited (visited) places.

³This method is closely related to the most advanced neuroscience model of the hippocampus.

The model is trained with the softmax cross-entropy loss and predicts the subsequent sensory observation (i.e., dotted squares). We evaluate two types of memory: **working memory (WM)** and **reference memory (RM)**⁴. When the prediction on nodes that were previously visited during the random walking is incorrect, it will count as a WM error (see Fig. 2 left). On the other hand, when the prediction on unvisited nodes is incorrect, it will count as a RM error (see Fig. 2 right). Minimizing the RM error by memorizing input sequences is infeasible; the possible number of sequence configurations is exponential since the input sequence is randomly generated at each trial. To solve this task, the model should be able to 1) understand the abstract structure of 2D space, 2) infer which map it is on from input sequence data, and 3) memorize what sensory observation is placed at each position in that map. See Appendix A.1 for training, evaluation, and transformer model details.

3 Results

WM error & RM error To measure the impact of nonlinearity α in the FNNs, we train the transformer models with different values of α in $[0, 0.01, 0.05, 0.1, 0.5, 1, 5, 10]$ and evaluate the WM and RM errors on the train maps (i.e., familiar maps) and test maps (i.e., novel maps). The average number of unvisited nodes in a single trial is 561.

The top left plot in Fig. 3a shows that the RM error on the train maps is rapidly decreased over train trials when α is larger than zero, with a larger improvement for increasing α . The RM error on the novel maps, however, is nearly constant at the chance level of 0.9 ($= 1 - 1/(\text{number of letters})$) for all α (see Fig. 3a top right). Fig. 3a (bottom right) shows that WM is active on novel maps that had not been shown during training. This finding suggests that the WM formation is intact on novel maps. Training the models on different numbers of maps N , Fig. 3b shows that increasing nonlinearity (i.e., α) helps activate the RM, and the trend of improvement is consistently shown for $N = 32, 48$, and 64 cases. Training over more maps leads to bigger RM errors. This is because more maps require the model to store more pairs of ‘what’-‘where’ memory (i.e., each training contains unique ‘what’-‘where’ information).

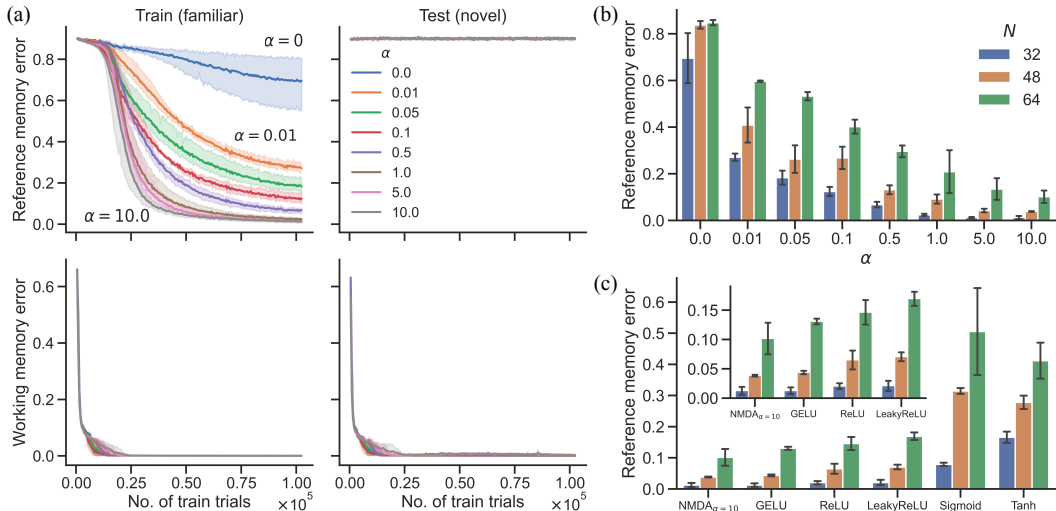


Figure 3: (a) Reference and working memory errors over training trials for training (familiar) maps and testing (novel) maps for $N = 32$ where N is the number of training maps. (b) Reference memory errors evaluated on training maps over different values of α in NMDA_α and N . (c) Reference memory errors comparison between $\text{NMDA}_\alpha = 10$, GELU, ReLU, LeakyReLU, Sigmoid, and Tanh activation functions. Inset: zoom on the top 4 activation functions. Error bars and shaded areas represent the standard deviation of errors from three independently trained models.

⁴Whittington et al. [13] only evaluated the WM error based on our definitions of WM and RM.

In addition, we demonstrate other nonlinear activation functions which are widely used in the machine learning literature. We test GELU ($x\sigma(1.702x)$), ReLU ($\max(0, x)$), LeakyReLU ($\max(0, x) + 0.01 \min(0, x)$), Sigmoid, and Tanh in the FFNs. As can be seen in Fig. 3c, $\text{NMDA}_{\alpha=10}$ shows the lowest RM errors on the training maps.

Place cells in feed-forward networks Place cell is a neuron in the hippocampus which fires at a particular place of the environment [18]. Studies have shown that hippocampal place cells encode the spatial location through localized firing patterns. They have been considered a substrate for long-term memory of the location where specific events occurred (i.e., previously visited position in our navigation task). Selective impairment of NMDAR in hippocampal CA1 disrupts place cell emergence and long-term memory formation [2, 3, 19].

We investigate the role of neurons in the FFNs and self-attention layers by measuring the neuron's place specificity. Given a $K \times K$ 2D grid environment as graph $G = (V, E)$ and a firing rate (cumulative activation value at node i divided by the length of evaluation trial) of node $i \in V$ as a ρ_i , we define maximally firing node as i_{\max} and its firing rate as ρ_{\max} . Where E is directed edges, which connect high to low firing nodes in G . From G , we run depth-first-search from source node, i_{\max} , to build a sub-graph $\mathcal{G} = (\mathcal{V}, \mathcal{E})$ which we call all connected components. Given G and \mathcal{G} , the place cell score is defined as following

$$\text{Place cell score} = \gamma \frac{\sum_{i \in \mathcal{V}} \rho_i}{\sum_{i \in V} \rho_i}, \quad (3)$$

where $\gamma = 1 - |\mathcal{V}^*|/|V|$ is a discount factor and \mathcal{V}^* is \mathcal{V} without node i_{\max} and leaf nodes. To measure place cell score, we record the firing rate ρ_i of neurons over a random walking trajectory with 10^5 steps in one of the training maps; then we measure the place cell scores of neurons in the FFN and self-attention layers. The place cell score is 1 when the neuron is firing only at a certain node; the score is 0 when the neuron is firing homogeneously across all nodes.

Fig. 4b and 4c show the rate maps of neurons with place cell scores in the FFNs and self-attention layers, respectively (Fig. 4a). As can be seen, our metric well represents place specificity. Fig. 4d

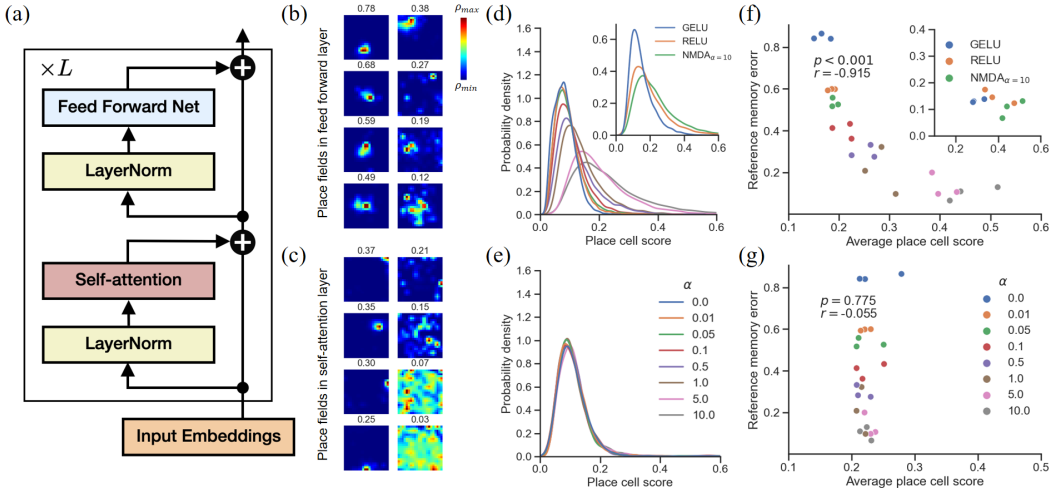


Figure 4: Reference memory-related place cells selectively emerge in the feed-forward layer but not in the self-attention layer along with α increase. (a) The transformer architecture used in the current study. (b, c) Example rate maps with place scores in feed-forward layers and self-attention layers at $\alpha = 10$ and $N = 64$; from top left (high) to bottom right (low) (d) Place cell score distribution in feed-forward layers change along with α modulation. Inset, comparison of $\text{NMDA}_{\alpha=10}$ with ReLU and GELU activation function. (e) Place cell score distribution in self-attention layers does not change along with α modulation. Inset, comparison of $\text{NMDA}_{\alpha=10}$ with ReLU and GELU activation function. (f-g) Scatter plot of average place cell scores and reference memory errors. r and p denote Spearman's rank correlation coefficient and significance score, respectively. All results are evaluated from training maps.

and 4e show the distribution of place cell scores in FFNs and self-attention layers with different values of α . As we increase α , the place cell score distribution found in FFNs gets positively shifted (see Fig. 5 for rate maps for $\alpha = 0, 1.0$, and 10.0 in Appendix A.2), whereas place cell score distribution in the self-attention layers remains. In addition, Fig. 4f and 4g show a relationship between the average place cell score and RM error for each α . While average place cell scores in the self-attention layer show no correlation with RM errors whatsoever, neurons in the FFN layer exhibit substantial correlation. Moreover, compared to two widely used GELU or ReLU activation functions, NMDA $_{\alpha=10}$ shows better place cell representations (Fig. 4f inset). These results imply that the RM formation and place cell emergence can be enhanced by NMDAR-like nonlinearity in the FFN.

4 Discussion and Conclusion

Rigorous previous efforts in finding the optimal nonlinear activation function underlie the great success of modern deep neural network models [20, 9, 21]. However, the neural substrates that mediate nonlinearity in the human brain and their role in intelligence have not been clearly understood. Our work is one of the first to put together the biologically inspired nonlinearity and its effect on long-term memory formation and the place cell representation in the previously described transformer model of the hippocampal formation. This idea was tested on a sensory observation task in the 2D grid environment and with the implementation of NMDAR-like nonlinearity. Our data indicated that NMDAR-like nonlinearity in the FFNs of transformers can enhance the formation of long-term memory and spatial place cell representation. Furthermore, this design choice improves long-term memory more than other commonly used nonlinear functions.

Whittington et al. [13] showed that softmax neurons in the self-attention layer behave like place cells and demonstrated that changing the softmax function to linear slows the WM learning process. However, the role of neurons in FFNs has not been studied. We demonstrate for the first time that place cells could emerge in transformers' FFNs, which we show by testing the emergence of place cells in FFNs with an NMDA-inspired activation function. Even though there are trainable parameters in the self-attention layer, the quantitative analysis of the place cell score indicates that most of the RM is stored in FFNs. Our results agree qualitatively with previous NMDAR impairment experiments from neuroscience: 1) hippocampal CA1 NMDAR perturbation does not impair WM [22], 2) changing NMDAR Mg²⁺-gating (changing α in this work) enhances or disrupts long-term memory formation [7, 8], 3) NMDAR is required for long-term stabilization of newly forming place fields [19, 3]. Our contribution is at showing these patterns experimentally for the first time.

Our research has exciting future directions. The current study only examined what-where memory using a sensory observation task in a static environment. However, our real-world environment is changing dynamically. Unfortunately, modern deep learning systems are generally incapable of adapting to a dynamic environment or reordering sensory inputs. In future work, we intend to explore what-where-when memory, called *episodic memory*, in transformer and other deep models.

Acknowledgments and Disclosure of Funding

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Checklist

1. For all authors...
 - (a) Do the main claims made in the abstract and introduction accurately reflect the paper’s contributions and scope? **[Yes]** See Abstract and Introduction.
 - (b) Did you describe the limitations of your work? **[Yes]** See Discussion section.
 - (c) Did you discuss any potential negative societal impacts of your work? **[N/A]**
 - (d) Have you read the ethics review guidelines and ensured that your paper conforms to them? **[Yes]** Yes, I read it and this paper conforms to them.
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 - (a) Did you state the full set of assumptions of all theoretical results? **[N/A]**
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 - (a) Did you include the code, data, and instructions needed to reproduce the main experimental results (either in the supplemental material or as a URL)? **[Yes]** All training, evaluation, and model details have been specified in the text. The code will be released with the camera-ready version.
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 - (c) Did you report error bars (e.g., with respect to the random seed after running experiments multiple times)? **[Yes]** We ran 3 different random seeds.
 - (d) Did you include the total amount of compute and the type of resources used (e.g., type of GPUs, internal cluster, or cloud provider)? **[Yes]** See Appendix A.1.
4. If you are using existing assets (e.g., code, data, models) or curating/releasing new assets...
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 - (a) Did you include the full text of instructions given to participants and screenshots, if applicable? **[N/A]**
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A Appendix

A.1 Training, evaluation, and model configuration details

In our experiment, the feed-forward network (FFN) in the transformer model consists of two linear layers with the NMDAR-inspired activation function NMDA_α (Eq. 2). We used TransformerXL [23] with an extended memory length of 32 and segment length of 32 so that working memory error is measured within a sequence length of 65($= 64 + 1$; 1 for the masked sensory input); i.e. a node that the agent had never visited within recent 64 steps is treated as an unvisited node. The input embedding is concatenated vector $[e, x]$ of the word embedding x (dimension of 256) and the recurrent positional embedding e (dimension of 256) so that the total input embedding dimension is 512. The number of heads in the self-attention layer is 8 and the number of neurons in the FFN is 2,048. The dropout rate is set to 0.1 and the maximum clip norm of the gradient is set to 0.25. We employ ADAM [24] optimizer and a learning rate schedule with a linear decay from 0.0001 (start) to 0 (end). We run 512 random walk simulations in parallel for collecting training trajectories. The total number of random walking steps is 2,048 for each simulation so the total number of gradient steps for each run is 512 (batch size) \times 2,048 (total number of steps in a trial) \times 200 (number of trials). All runs are performed on a single NVIDIA TITAN V GPU.

A.2 Analysis details of place cell distribution in transformer

We plot each place cell score distribution with neurons from 3 independent experiments. For the self-attention layer, the total number of neurons in the softmax layer is 65 (number of sequence length) \times 8 (number of head) \times 2 (number of layers). For the feed-forward networks, the total number of neurons in the feed-forward layer is 2048 (number of neurons) \times 2 (number of layers). Rate maps of neurons with top-64 place scores in FFNs with varying α are shown in Figure 5.

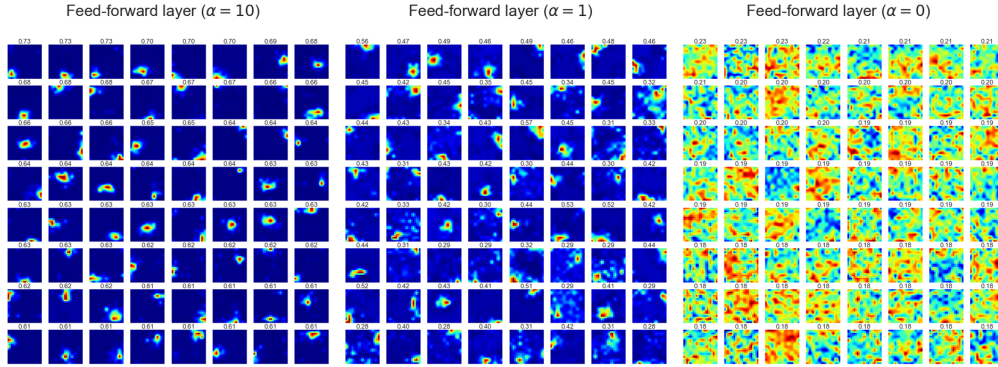


Figure 5: Rate maps of neurons with top-64 place scores in FFNs with varying values of α ; $\alpha = 10$ (left), $\alpha = 1$ (middle), and $\alpha = 0$ (right).