Deep Spiking Neural Networks with High Representation Similarity Model Visual Pathways of Macaque and Mouse

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Abstract

Deep artificial neural networks (ANNs) play a major role in modeling the visual pathways of primate and rodent. However, they highly simplify the computational properties of neurons compared to their biological counterparts. Instead, Spiking Neural Networks (SNNs) are more biologically plausible models since spiking neurons encode information with time sequences of spikes, just like biological neurons do. However, there is a lack of studies on visual pathways with deep SNNs models. In this study, we model the visual cortex with deep SNNs for the first time, and also with a wide range of state-of-the-art deep CNNs and ViTs for comparison. Using three similarity metrics, we conduct neural representation similarity experiments on three neural datasets collected from two species under three types of stimuli. Based on extensive similarity analyses, we further investigate the functional hierarchy and mechanisms across species. Almost all similarity scores of SNNs are higher than their counterparts of CNNs with an average of 6.6\%. Depths of the layers with the highest similarity scores exhibit little differences across mouse cortical regions, but vary significantly across macaque regions, suggesting that the visual processing structure of mice is more regionally homogeneous than that of macaques. Besides, the multi-branch structures observed in some top mouse brain-like neural networks provide computational evidence of parallel processing streams in mice, and the different performance in fitting macaque neural representations under different stimuli exhibits the functional specialization of information processing in macaques. Taken together, our study demonstrates that SNNs could serve as promising candidates to better model and explain the functional hierarchy and mechanisms of the visual system.

Introduction

Originally, the prototype of deep neural networks is inspired by the biological vision system (Hubel and Wiesel 1959, 1962). To date, deep neural networks not only occupy an unassailable position in the field of computer vision (LeCun, Bengio, and Hinton 2015), but also become better models of the biological visual cortex compared to traditional models in the neuroscience community (Khaligh-Razavi and Kriegeskorte 2014; Yamins et al. 2014; Yamins and DiCarlo 2016). They have been successful at predicting the neural responses in primate visual cortex, matching the hierarchy of ventral visual stream (Güçlü and van Gerven 2015; Kubilius et al. 2019; Nayebi et al. 2018; Kietzmann et al. 2019), and even controlling neural activity (Bashivan, Kar, and DiCarlo 2019; Ponce et al. 2019). Moreover, as training paradigms of mice (Zoccolan et al. 2009) and techniques for collecting neural activity (de Vries et al. 2020) have been greatly improved, there is a strong interest in exploring mouse visual cortex. Deep neural networks also play an important role in revealing the functional mechanisms and structures of mouse visual cortex (Shi, Shea-Brown, and Buice 2019; Cadena et al. 2019; Nayebi et al. 2022; Bakhtiari et al. 2021; Conwell et al. 2021).

Compared to biological networks, Artificial Neural Networks discard the complexity of neurons (Pham, Packianather, and Charles 2008). Spiking Neural Networks, incorporating the concept of time and spikes, are more biologically plausible models (Maass 1997). To be more specific, because of their capabilities of encoding information with spikes, capturing the dynamics of biological neurons, and extracting spatio-temporal features, deep SNNs are highly possible to yield brain-like representations (Hodgkin and Huxley 1952; Gerstner and Kistler 2002; Izhikevich 2004; Brette et al. 2007; Kasabov et al. 2013). However, deep SNNs have not been employed to model visual cortex due to the immaturity of training algorithms. Recently, a state-of-the-art directly trained deep SNN (Fang et al. 2021), makes it possible to use deep SNNs as visual cortex models.

Contributions. In this work, we conduct large-scale neural representation similarity experiments on SNNs and other high-performing deep neural networks to study the brain’s visual processing mechanisms, with three datasets and three similarity metrics (Figure 1). Specifically, to the best of our knowledge, we are the first to use deep SNNs to fit complex biological neural representations and explore the biological visual cortex. We summarize our main contributions in four points as follows.

• We find that SNNs outperform their counterparts of CNNs with the same depth and almost the same architectures in almost all experiments. In addition, even with very different depths and architectures, SNNs can achieve top performance in most conditions.
• By making a more direct comparison between macaques
and mice for the first time, we reveal the differences in the visual pathways across the two species in terms of the homogeneity of visual regions and the increases of receptive field sizes across cortical visual pathways, which is consistent with previous physiological work.

- The multi-branch structures in neural networks benefit neural representation similarity to mouse visual cortex, providing computational evidence that parallel information processing streams are widespread between cortical regions in the mouse visual system.

- Comparing the results of two macaque neural datasets under different stimuli, we reveal that the macaque vision system may have functional specialization for processing human faces and other natural scenes.

Altogether, as the first work to apply deep SNNs to fit neural representations, we shed light on visual processing mechanisms in both macaques and mice, demonstrating the potential of SNNs as a novel and powerful tool for research on the visual system. Our codes and appendix are available at https://github.com/Grasshlw/SNN-Neural-Similarity.

Related Work

There are plenty of computational models of macaque and mouse visual systems for exploring the visual processing mechanisms recently. We summarize some of the outstanding work in the following.

The network models of macaque visual system. In the early days, studies basically used simple feedforward neural networks as the models of the macaque visual system (Khaligh-Razavi and Kriegeskorte 2014; Yamins et al. 2014; Yamins and DiCarlo 2016). Recently, some bio-inspired or more complex models achieved better performance in fitting the neural representations of macaque visual cortex (Kubilius et al. 2019; Dapello et al. 2020; Zhuang et al. 2021; Higgins et al. 2021) also effectively matched the neural activity of macaque ventral visual stream. Although the above work developed many bio-inspired structures, the networks are still traditional ANNs in nature. Our work introduces deep SNNs for the first time to explore the visual processing mechanisms of macaque visual system.

The network models of mouse visual system. Large-scale mouse neural dataset provided an experimental basis for model studies of mouse visual system (de Vries et al. 2020; Siegle et al. 2021). (Shi, Shea-Brown, and Buice 2019) conducted comparisons between the representations of mouse visual cortex and the VGG16 trained on the ImageNet dataset. In (Bakhtiari et al. 2021), they developed a single neural network to model both the dorsal and ventral pathways with showing the functional specializations. What’s more, a large survey of advanced deep networks (Conwell et al. 2021) revealed some hierarchy and functional properties of mice. Similar to the studies of macaque visual system, deep SNNs have never been used to model the mouse visual system. In this work, we not only use SNNs as one of the candidates to fit the representations of mouse visual cortex, but also conduct direct comparisons between macaques and mice to further investigate the functional hierarchy and mechanisms of the two species.

Methods

NeuralDatasets

Our work is conducted with three neural datasets. These datasets are recorded from two species under three types of stimuli. More specifically, there are neural responses of mouse visual cortex to natural scene stimuli, and responses of macaque visual cortex to face image and synthetic image stimuli.

Allen Brain mouse dataset. It is part of the Allen Brain Observatory Visual Coding dataset (Siegle et al. 2021) col-
Selected using Neuropixel probes from 6 regions simultaneously in mouse visual cortex. Compared to two-photon calcium imaging, Neuropixel probes simultaneously record the spikes across many cortical regions with high temporal resolution. In these experiments, mice are presented with 118 250-ms natural scene stimuli in random orders for 50 times. Hundreds to thousands of neurons are recorded for each brain region. To get the stable neurons, we first concatenate the neural responses (average number of spikes in 10-ms bins across time) under 118 images for each neuron, and then preserve the neurons whose split-half reliability across 50 trials reaches at least 0.8.

**Macaque-Face dataset.** This dataset (Chang et al. 2021) is composed of neural responses of 159 neurons in the macaque anterior medial (AM) face patch under 2,100 real face stimuli, recorded with Tungsten electrodes. For this dataset, we compute the average number of spikes in a time window of 50-350ms after stimulus onset and exclude eleven neurons with noisy responses by assessing the neurons’ noise ceiling. The details of the preprocessing procedure are the same as (Chang et al. 2021).

**Macaque-Synthetic dataset.** This dataset (Majaj et al. 2015) is also about macaque neural responses which are recorded by electrodes under 3,200 synthetic image stimuli, and used for neural prediction in the initial version of Brain-Score (Schrimpf et al. 2020a). The image stimuli are generated by adding a 2D projection of a 3D object model to a natural background. The objects consist of eight categories, each with eight subclasses. The position, pose, and size of each object are randomly selected. 88 neurons of V4 and 168 neurons of IT are recorded. The neural responses are preprocessed to the form of average firing rate and can be downloaded from Brain-Score.

**Models**

Since the core visual function of macaque and mouse visual cortex is to recognize objects, the basic premise of model selection is that the model has good performance on object recognition tasks (e.g. classification on ImageNet). Based on this premise, we employ 12 SNNs, 43 CNNs, and 26 vision transformers, all of which are pretrained on the ImageNet dataset and perform well in the classification task. As for SNNs, we use SEW ResNet as the base model, which is the deepest and SOTA directly trained SNN (Fang et al. 2021). Furthermore, by combining the residual block used in SEW ResNet and the hierarchy of the visual cortex, we build several new SNNs and train them on the ImageNet using SpikingJelly (Fang et al. 2020) (see Appendix A for model structures and the details of model training). As for CNNs and vision transformers, we use 44 models from the Torchvision model zoo (Paszke et al. 2019), 22 models from the Timm model zoo (Wightman 2019) and 3 models from the brain-like CNNs, CORnet family (Kubilius et al. 2019). In the feature extraction procedures of all models, we feed the same set of images used in biological experiments to the pretrained models and obtain features from all chosen layers. Different from CNNs and vision transformers, the features of SNNs are spikes in multiple time steps.

**Similarity Metrics**

To obtain the representation similarity between biological visual cortex and computational models, we apply three similarity metrics to computing similarity scores: representational similarity analysis (RSA) (Kriegeskorte et al. 2008; Kriegeskorte, Mur, and Bandettini 2008), regression-based encoding method (Carandini et al. 2005; Yamins et al. 2014; Schrimpf et al. 2020a,b) and singular vector canonical correlation analysis (SVCCA) (Raghu et al. 2017; Morcos, Raghu, and Bengio 2018). RSA has already been widely used to analyze neural representations of a model and a brain to different stimuli at the population level, while the regression-based encoding method directly fits the model features to neural activity data. SVCCA is originally proposed to compare features of deep neural networks, and then (Shi, Shea-Brown, and Buice 2019) used it to compare representation matrices from mouse visual cortex and DNNs, which demonstrated its effectiveness.

With the same model and same cortical region, we use these metrics for a layer-by-layer comparison to compute the similarity scores. The maximum similarity score across layers for a given cortical region is considered to be the level of representation similarity between the model and the cortical region. Finally, in a given dataset, we take the average score of all cortical regions as the final similarity score for each model, which gives the overall model rankings. The implementation of each similarity metric is as follows.

**RSA.** For two response matrices $R \in \mathbb{R}^{n \times m}$ from each layer of models and each cortical region, where $n$ is the number of units/neurons and $m$ is the number of stimuli, we calculate the representational similarity between the responses to each pair of image stimuli using the Pearson correlation coefficient $r$, yielding two representative dissimilarity matrices ($RDM \in \mathbb{R}^{m \times m}$, where each element is the correlation distance $1 - r$). Then, the Spearman rank correlation coefficient between the flattened upper triangles of these two matrices is the metric score.

**Regression-Based Encoding Method.** Firstly, we run truncated singular value decomposition (TSVD) to reduce the feature dimension of model layers to 40. Secondly, the features after dimensionality reduction are fitted to the representations of each neuron by ridge regression. Finally, we compute the Pearson correlation coefficient between the predicted and ground-truth representations of each neuron and take the mean of all correlation coefficients as the metric score. More specifically, we apply leave-one-out cross-validation to obtain predicted representations of each neuron. For simplicity, we name this method 'TSVD-Reg'.

**SVCCA.** For both the responses of model layers and cortical regions, we use TSVD to reduce the dimension of unit/neuron to 40, yielding two reduced representation matrices. Then we apply canonical correlation analysis (CCA) to these two matrices to obtain a vector of correlation coefficients (the length of the vector is 40). The metric score is the mean of the vector. Because of the invariance of CCA to affine transformations (Raghu et al. 2017), in this procedure, we only need to ensure that the stimulus dimension is consistent and aligned, even if the unit/neuron dimension is different. Dimensionality reduction plays an important role.
in this method to make the number of model features comparable to the number of neurons in cortical regions, since the former usually far exceeds the latter. In addition, dimensionality reduction helps to determine which features are important to the original data, while CCA suffers in important feature detection. Using just CCA performs badly, which has been proven by (Raghu et al. 2017).

Results

Comparisons of Representation Similarity Scores between SNNs and Other Types of Models

To check how similar the models are to the visual cortex’s mechanisms in visual processing, we rank the final similarity scores of all models and conduct comparisons among three types of models (CNNs, SNNs, and vision transformers). Specially, we focus on comparing SNN (SEW ResNet) and CNN (ResNet) with the same depth and almost the same architectures (Figure 2). The final similarity score of a model is the average similarity score across all cortical regions. (The overall rankings can be found in Appendix B and the comparisons among three types of models are shown in Appendix C.)

Allen brain mouse dataset. No single model achieves the highest final similarity scores with all three metrics. For a fair comparison, we apply the paired t-test to SEW ResNet and ResNet with the same depth. For all three metrics, SEW ResNet performs better than ResNet by a large margin ($t = 5.857$, $p = 0.004$; $t = 7.666$, $p = 0.002$; $t = 7.592$, $p = 0.002$).

Macaque-Face dataset. For both SVCCA and TSVD-Reg, Wide-SEW-ResNet14 and Wide-SEW-ResNet8 achieve the first and second highest final similarity scores respectively. But for RSA, TNT-S and Inception-ResNet-V2 take their place and outperform other models by a large margin. As for SEW ResNet and ResNet, the former performs significantly better than the latter for both SVCCA and TSVD-Reg ($t = 8.195$, $p = 0.001$; $t = 7.528$, $p = 0.002$). However, the difference is not significant for RSA ($t = 1.117$, $p = 0.327$). Specifically, the similarity score of SEW ResNet152 is only slightly higher than that of ResNet152, and at the depth of 50 and 101, SEW ResNet’s scores are lower than ResNet’s.

Macaque-Synthetic dataset. Similar to the results of Allen Brain dataset, no model performs best for all three metrics. SEW ResNet performs moderately better than ResNet ($t = 3.354$, $p = 0.028$; $t = 3.824$, $p = 0.019$; $t = 2.343$, $p = 0.079$). The only contrary is that SEW ResNet18 performs worse than ResNet18 for RSA.

Further, to check the details of comparison between the SNNs and their CNN counterparts, we analyze the trajectories of similarity score across model layers (Figure 3). As for ResNet and SEW ResNet with the same depth, the trends of their similarities across model layers are almost the same, but the former’s trajectory is generally below the latter’s. In other words, the similarity scores of SEW ResNet are higher than those of ResNet at almost all layers.

Taken together, the results suggest that when the overall results that appear below also correspond to the three metrics in this order, unless the correspondence is stated in the text.
For mouse visual cortex (Figure 4 (a)), taking the deep model group as an example, ANOVA shows overall significant changes in depth across cortical regions for TSVD-Reg and RSA (Friedman’s $\chi^2 = 49.169, p = 2.0 \times 10^{-9}$, $\chi^2 = 19.455, p = 0.002$). But there is no significant change for SVCCA ($\chi^2 = 8.689, p = 0.122$). According to these results, the differences in depth across regions are indeterminacy and irregular. Meanwhile, the trends of layer depth between some regions contradict the hierarchy observed in physiological experiments of mice (those between VISp and VISl for TSVD-Reg and between VISal and VISpm for RSA). However, for macaque visual cortex (Figure 4 (b)), there are significant differences ($t = -5.451, p = 6.5 \times 10^{-6}; t = -8.312, p = 2.8 \times 10^{-9}; t = -3.782, p = 6.9 \times 10^{-4}$, also taking the deep model group as an example) between V4 and IT, and the trend is consistent with the information processing hierarchy in primate visual cortex.

The comparative analyses of the best layer depths of the shallow and deep model groups also exhibit the differences between macaques and mice. For mouse visual cortex, the best layer depths of shallow models are significantly higher than those of deep models. Compared to deep models, most shallow models achieve the top similarity scores in intermediate and even later layers. Differently, for macaque visual cortex, the depth of models has little effect on the depth of the most similar layer. What’s more, we find that the most similar layer of mouse visual cortex always occurs after the $28 \times 28$ feature map is downsampled to $14 \times 14$, which leads to the layer depths’ difference between shallow and deep models. Nevertheless, the best layer of macaque IT appears in the last part of networks, where the feature map has been downsampled more times.

In summary, our results might reveal two distinctions in the functional hierarchy between macaques and mice. First, there is a distinct functional hierarchical structure of macaque ventral visual pathway, while there might be no clear sequential functional hierarchy in mouse visual cortex. One explanation is that the mouse visual cortex is organized into a parallel structure and the function of mouse cortical regions are more generalized and homogeneous than those of macaques. Another possibility would be that even though the sequential relations exist among mouse cortical regions as proposed in anatomical and physiological work, they are too weak for the current deep neural networks to capture. Additionally, mice perform more complex visual tasks than expected with a limited brain capacity (Djurdjevic et al. 2018). Consequently, the neural responses of mouse visual cortex may contain more information not related to object recognition that neural networks focus on. Secondly, it is well known that the units in the neural networks get larger in later layers. This means that the in-determinacy and irregularity of the neural responses due to the downsampling of the receptive fields may be less pronounced.

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reduce the number of parameters and improve its similarity. Any detailed structures in the neural networks help to reduce the number of model parameters but not with the depth of models. It calls into the question whether other hand, the similarity scores of the mouse dataset are consistent with Spearman’s rank correlation and are shown in Table 1 and 2. We first measure the correlation between the similarity scores and the model size. The results are consistent with Spearman’s rank correlation and are shown in Appendix E. Based on these results, we further investigate more detailed properties of neural networks to explain the processing mechanisms in the visual cortex. For the mouse dataset, on the one hand, the best layer depths show non-significant changes across the mouse cortical regions as mentioned in the previous section. On the other hand, the similarity scores of the mouse dataset are only correlated with the number of model parameters but not with the depth of models. It calls into the question whether any detailed structures in the neural networks help to reduce the number of parameters and improve its similarity to mouse visual cortex. Therefore, we explore the commonalities between models that have the top 20% representation similarities (see Appendix D) for Allen Brain dataset. As expected, the top models contain similar structures, such as fire module, inception module, and depthwise separable convolution. All these structures essentially process information through multiple branches/channels and then integrate the features from each branch. The models with this type of structure outperform other models ($t = 2.411, p = 0.024; t = 3.030, p = 0.007; t = 1.174, p = 0.247$). Moreover, we apply the depthwise separable convolution to SNNs, which yields a positive effect. The representation similarity of Spiking-MobileNet is higher than SEW-ResNet50 with a similar depth (+0.8%; +3.9%; +12.1%). In fact, some studies using multiple pathways simulate the functions of mouse visual cortex to some extent (Shi et al. 2022; Nayebi et al. 2022). Our results further suggest that not only the mouse visual cortex might be an organization of parallel structures, but also there are extensive parallel information processing streams between each pair of cortical regions (Wang, Sporns, and Burkhalter 2012; Siegle et al. 2021).

For the two macaque datasets with different stimuli, not only are the model rankings significantly different, but also the correlations between the similarity scores and the model depth are totally opposite. These results corroborate the following two processing mechanisms in macaques: the ventral visual stream of primate visual cortex possesses canonical coding principles at different stages; the brain exhibits a high degree of functional specialization, such as the visual recognition of faces and other objects, which is reflected in the different neural responses of the corresponding region (although the face patch AM is a sub-network of IT, they differ in the neural representations). Besides, as shown in Figure 5,
the similarity scores of vision transformers reach the maximum in the early layers and then decrease. Differently, the scores of CNNs and SNNs keep trending upwards, reaching the maximum in almost the last layer. On the other hand, Appendix C shows that vision transformers perform well in Macaque-Face dataset but poorly in Macaque-Synthetic dataset. Considering the features extraction mechanism of vision transformers, it divides the image into several patches and encodes each patch as well as their internal relation by self-attention. This mechanism is effective for face images that are full of useful information. However, the synthetic image consists of a central target object and a naturalistic background. When vision transformers are fed with this type of stimuli, premature integration of global information can lead to model representations containing noise from the unrelated background. What’s more, when we take all models with the top 20% representation similarities as a whole for analyses, as described in the above paragraph, the properties that enable networks to achieve higher neural similarity are not yet clear. Taken together, the computational mechanism of the better models may reveal core processing divergence to different types of stimuli in the visual cortex.

Discussion

In this work, we take large-scale neural representation similarity experiments as a basis, aided by analyses of the similarities across models and the visual cortical regions. Compared to other work, we introduce SNNs in the similarity analyses with biological neural responses for the first time, showing that SNNs achieve higher similarity scores than CNNs that have the same depth and almost the same architectures. As analyzed in Section 3.1, two properties of SNNs might serve as the explanations for their high similarity scores.

The subsequent analyses of the models’ simulation performance and structures indicate significant differences in functional hierarchies between macaque and mouse visual cortex. As for macaques, we observed a clear sequential hierarchy. However, as for mouse visual cortex, some work (Conwell et al. 2021) exhibits that the trend of the model feature complexity roughly matches the processing hierarchy, but other work suggests that the cortex (Shi, Shea-Brown, and Buice 2019; Nayebi et al. 2022) is organized into a parallel structure. Our results are more supportive of the latter. Furthermore, we provide computational evidence not only that the increased ratio of the receptive field size in cortical regions across the mouse visual pathway is smaller than those across the macaque visual pathway, but also that there may be multiple pathways with parallel processing streams between mouse cortical regions. Our results also clearly reveal that the processing mechanisms of macaque visual cortex differ to various stimuli. These findings provide us with new insights into the visual processing mechanisms of macaque and mouse, which are the two species that dominate the research of biological vision systems and differ considerably from each other.

Compared to CNNs, the study of task-driven deep SNNs is just in its initial state. Although we demonstrate that SNNs outperform their counterparts of CNNs, SNNs exhibit similar properties as CNNs in the further analyses. In this work, we only build several new SNNs by taking the hints from the biological visual hierarchy, while many well-established structures and learning algorithms in CNNs have not been applied to SNNs yet. In addition, the neural datasets used in our experiments are all collected under static image stimuli, lacking rich dynamic information to some certain, which may not fully exploit the properties of SNNs. Given that SNNs perform well in the current experiments, we hope to explore more potential of SNNs in future work.

In conclusion, as more biologically plausible neural networks, SNNs may serve as a shortcut to explore the biological visual cortex. With studies on various aspects of SNNs, such as model architectures, learning algorithms, processing mechanisms, and neural coding methods, it’s highly promising to better explain the sophisticated, complex, and diverse vision systems in the future.
Ethics Statement

The biological neural datasets used in our experiments are obtained from public datasets or from published papers with the authors’ consent.

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References


