# Scaling Properties for Artificial Neural Network Models of a Small Nervous System

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Abstract—The nematode worm C. elegans provides a unique opportunity for exploring in silico data-driven models of a whole nervous system, given its transparency and well-characterized nervous system facilitating a wealth of measurement data from wet-lab experiments. This study explores the scaling properties that may govern learning the underlying neural dynamics of this small nervous system by using artificial neural network (ANN) models. We investigate the accuracy of self-supervised next timestep neural activity prediction as a function of data and models. For data scaling, we report a monotonic log-linear reduction in mean-squared error (MSE) as a function of the amount of neural activity data. For model scaling, we find MSE to be a nonlinear function of the size of the ANN models. Furthermore, we observe that the dataset and model size scaling properties are influenced by the particular choice of model architecture but not by the precise experimental source of the C. elegans neural data. Our results fall short of producing long-horizon predictive and generative models of C. elegans whole nervous system dynamics but suggest directions to achieve those. In particular our data scaling properties extrapolate that recording more neural activity data is a fruitful near-term approach to obtaining better predictive ANN models of a small nervous system.

*Index Terms*—nervous system, neural networks, machine learning, neuroscience, scaling laws

## I. INTRODUCTION

Exploring neural system dynamics is crucial in neuroscience and artificial intelligence (AI). This intersection has spurred the evolution of artificial neural network (ANN) models, inspired by biological neural systems. ANNs offer the potential to emulate diverse animal behaviors, providing advantages like detailed specification, causal manipulability, and increasing analytical accessibility, reflecting key aspects of biological nervous systems ([1], [2]). The nematode *Caenorhabditis elegans* (*C. elegans*) is an exemplary model in this context, offering a valuable platform for comparing real and artificial neural dynamics.

*C. elegans* is an excellent model organism for neural dynamics research due to its well-mapped connectome and

capabilities for non-invasive neuronal activity tracking via advanced imaging techniques ([3], [4]). The organism's compact size, transparency, and well-annotated genome allow for intricate optical measurements and deep insights into neural activity. NeuroPAL, a multicolor atlas, allows precise *in vivo* neuron identification, enhancing the capabilities for measurement and analysis of the *C. elegans* nervous system [5].

We formulate the problem of *in silico* nervous system modeling as a teacher-student framework. A real biological neural network (that of *C. elegans*, in our case) is the teacher, ANN models are the students, and self-supervised next time-step neural activity prediction is the curriculum. We consider different instances of the nervous system to be phenotypically matched animals (in our case, adult hermaphrodite worms).

Predicting future neural activity based on historical neural data is not new but the machine learning approach to do it has seen increasing adaption ever since advancements in models like LSTMs demonstrating some success in mammals [6]. In *C. elegans*, the simplified behavioral repertoire and consistent biology offer a unique setting for in-depth model analysis. Self-supervised learning, predicting future states from intrinsic neural patterns, reduces dependence on behaviorally annotated data. While acknowledging the importance of behavior in neural dynamics, our study concentrates on the inherent predictability within neural activity, exploring how neural dynamics can be predicted without direct behavioral reference, similar to how large language models (LLMs) uncover intricate structures in language data [7].

Research into ANNs' scaling properties has shown that improvements in model size, data volume, and computational resources significantly enhance performance ([8], [9]). The relationship between data size and model capacity is critical in optimizing model performance. However, this relationship in the context of predicting neural dynamics in biological organisms like *C. elegans* is not well-explored. Our study aims to fill this gap by analyzing the impact of data volume, model architecture, and size on ANN performance in neural activity prediction in *C. elegans*. These insights are crucial for optimizing experimental and modeling strategies in neuroscience, contributing to the development of more accurate predictive models for biological nervous systems.

### II. Methods

### A. Neural Activity Data

**Data sources**. We obtained 8 open-source datasets ([5], [10]–[16]) measuring neural activity in *C. elegans* (Table I). These datasets, each recorded under varying experimental conditions, quantify neural activity through the measurement of changes in calcium fluorescence ( $\Delta F/F_0$ ) within subsets of the worm's 302 neurons. The 8 experimental datasets contained different numbers of recorded instances of the nervous system (i.e. worms) as well as variable lengths of time for neural activity measurements (Fig. 1A). The experimental conditions were variable across datasets, ranging from freely moving [10], immobilized [12], and asleep [15] states, to optogenetically stimulated scenarios [11]. However, our modeling strategy is agnostic to the potentially differing conditions and protocols under which the data was acquired as long as the measured system (*C. elegans*) and metric ( $\Delta F/F_0$ ) is consistent.

 TABLE I

 Open-source C. elegans neural activity datasets.

Paper Link		Database Link		Files w/ Data	
tinyurl.com/Leifer2023		osf.io/e2syt/		exported_data.tar.gz	
tinyurl.com/Flavell2023		wormwideweb.org		YYYY-MM-DD-	
				*.json/h5	
tinyurl.com/Uzel2022		osf.io/3vkxn/		Uzel_WT.mat	
tinyurl.com/Yemini2021		zenodo.org/records/3906530		_Activity_OH*.mat	
tinyurl.com/Kaplan2020a		osf.io/9nfhz/		Neuron2019_Datamat	
tinyurl.com/Skora2018		osf.io/za3gt/		WTmat	
tinyurl.com/Nichols2017		osf.io/kbf38/		let.mat	
tinyurl.com/Kato2015		osf.io/2395t/		WT_Stim.mat	
Num. worms	Mean num. neurons ID'd		Num. ID'd neurons (min, max)		
103	69/122		(26, 102)		
42	88/136		(64, 115)		
6	50/138		(46, 58)		
49	110/125		(33, 179)		
19	36/114		(23, 51)		
12	46/129		(39, 55)		
44	34/108		(23, 43)		
12	42/127		(31, 51)		

Summary of key details such as publication references, data sources, content format, worm counts, and neuron labeling statistics.

**Standard data format**. Each dataset  $\mathcal{D}^n$  includes individual recordings from *n* worms, each consisting of neural activity and a mask indicating the subset of the 302 neurons that were measured and labelled. This mask ensures models are trained only on neural activity recorded from NeuroPAL labelled neurons (Fig. 1B). Specifically:

$$\mathcal{D}^n = \{\mathbf{X}^1, \mathbf{X}^2, \dots, \mathbf{X}^n\} \times \{\mathbf{y}^1, \mathbf{y}^2, \dots, \mathbf{y}^n\}$$
(1)

where  $n = |\mathcal{D}^n|$ .



Fig. 1. Worm neural datasets curation. Eight open-source *C. elegans* neural activity datasets were downloaded, preprocessed and assimilated. (A) The distribution of the number of worms and the total recording time in each dataset. (B) The neural activity data of all worms is organized a standard format involving a muti-dimensional time series and a boolean feature mask.

Each worm, indexed by k, has a data matrix  $\mathbf{X}^k \in \mathbb{R}^{302 \times T_k}$ and a binary vector  $\mathbf{y}^k \in \{0, 1\}^{302}$  specifying which neurons were recorded and labelled. Each row  $\mathbf{x}_i^k$  of the matrix  $\mathbf{X}^k$ contains the time series of neural activity for the *i*<sup>th</sup> neuron for  $T_k$  time steps. The rows of  $\mathbf{X}^k$  and are ordered according to the alphabetical canonical names of the neurons <sup>1</sup> with rows ordered analogously to  $\mathbf{y}^k$ .

**Preprocessing.** The data, denoted as  $\mathbf{X}^k$ , is processed from the original raw data. First, we normalized the calcium data of each worm by *z*-scoring the full time series independently for each neuron. We then smoothed the signal using a causal exponential kernel (smoothing parameter  $\alpha = 0.5$ ). Finally, the neural data was resampled to a fixed time step interval ( $\Delta t \approx 0.667s$ ). Fig. 2 steps through the preprocessing pipeline for a handful of neurons from one worm in the Kato dataset  $\mathcal{D}_{Kato}$  [16].

**Train-Test split.** For each worm's neural activity data matrix  $\mathbf{X}^k$ , we performed a temporal split to create a training set  $\mathbf{X}^k_{\text{train}}$  and a testing set  $\mathbf{X}^k_{\text{test}}$ . A balanced 50:50 split was adopted, allocating the first half of the neural activity recording to the training set and the second half to the testing set. One might create equally sized train and test sets containing  $n_s$  sequences of length *L* by sampling their start indices uniformly from (or equidistantly within) the range  $[0, \lfloor T/2 \rfloor - L - 1)$  and  $\lfloor \lceil T/2 \rceil, T - L - 1 \rfloor$  for train and test, respectively

Amount of Data. The ability to vary the amount of training data is central to our investigation of the effects of data scaling on the ability of self-supervised models to do future neural

<sup>1</sup>https://www.wormatlas.org/NeuronNames.htm



Fig. 2. Example of preprocessing worm neural data. We use the example of the Kato (2015) dataset to illustrate steps of the preprocessing the neural activity data. Starting with the raw calcium fluorescence signals (A); we first standardize or *z*-score each neuron independently (B); then smooth using a causal filter (C); and finally resample to a fixed time step (D).

activity prediction. However, we are data constrained in our setting since the collective dataset of all worms  $\mathcal{D}_{ALL}$  has only 284 worms (Fig. 3).

**Source Datasets.** The experimental sources contributing to  $\mathcal{D}_{ALL} = \mathcal{D}^{284}$  are Kato  $\mathcal{D}_{Kato}$  ( $|\mathcal{D}_{Kato}| = 12$ ), Nichols  $\mathcal{D}_{Nichols}$  ( $|\mathcal{D}_{Nichols}| = 44$ ), Skora  $\mathcal{D}_{Skora}$  ( $|\mathcal{D}_{Skora}| = 12$ ), Kaplan  $\mathcal{D}_{Kaplan}$  ( $|\mathcal{D}_{Kaplan}| = 19$ ), Yemini  $\mathcal{D}_{Yemini}$  ( $|\mathcal{D}_{Yemini}| = 49$ ), Uzel  $\mathcal{D}_{Uzel}$  ( $|\mathcal{D}_{Uzel}| = 6$ ), Flavell  $\mathcal{D}_{Flavell}$  ( $|\mathcal{D}_{Flavell}| = 42$ ), Leifer  $\mathcal{D}_{Leifer}$  ( $|\mathcal{D}_{Leifer}| = 103$ ).

**Mixed Datasets.** To create mixed datasets combining worms from the different experimental sources, we randomly sample from the combined pool of all available worms  $\mathcal{D}_{ALL}$ .

Let  $i \in [8] := \{1, 2, ..., 8\}$  index into the list of experimental sources (sorted by publication date): [Kato, Nichols, ..., Leifer]. We denote a mixed worm dataset containing  $n_w$  worms from



Fig. 3. Distribution of neurons in  $\mathcal{D}_{ALL}$ . Since the train-validation split is along the temporal (not feature) dimension, the neuron distribution is the same in the train and validation sets. The combined dataset has 284 worms from the 8 experimental dataset sources, with 247 out the 302 neuron classes in *C.elegans* represented (i.e. recorded in at least 1 worm). Of the recorded neurons, some are over-represented whereas others have been recorded in only 1 worm.

any combination of experimental sources as  $\mathcal{D}^{n_w}$ .

$$\mathcal{D}^{n_w} = \bigcup_{i=1}^8 \mathcal{D}_i^{(k_i)} \quad \text{s.t.} \quad \sum_{i=1}^8 k_i = n_w \tag{2}$$

Here,  $\mathcal{D}_i^{(k_i)}$  stands for  $k_i$  worms sampled specifically from the experimental dataset indexed by *i* (Fig. 4A). Note that there are multiple assignments that can achieve a dataset with  $n_w$  worms. Therefore,  $D^{n_w}$  is a random variable. Our mixed dataset sampling process is akin to sampling from a multinomial distribution where the probabilities correspond to the proportion of available worms from each experimental source:

$$\mathcal{D}^{n_w} \sim \text{Multinomial}(n_w; p_1, \dots, p_8)$$
 (3)

where  $p_i$  reflects the relative contribution of the experimental dataset indexed by *i* to the pool.

This methodical approach allows us to create increasingly larger mixed datasets up to the largest one containing all worms  $\mathcal{D}_{ALL} := \mathcal{D}^{284}$  (for which there is only 1 possible assignment). The result is a series of mixed datasets, each with a unique composition of worms, yet collectively spanning the full range of neural dynamics present in the collective data. The mixed datasets  $\mathcal{D}^{n_w}$  thus represent a diverse cross-section of neural activities encompassing variations in experimental conditions.

We could also generate subsets from any single experimental dataset using this approach by simply restricting our random sampling to that source. This allows us to create increasingly larger data subsets constrained to a particular experimental context.



Fig. 4. Sampling datasets and dataset scaling experiment. (A) Creating mixed worm datasets of various sizes by sampling from the pool of all available worms across different experimental datasets. (B) The validation loss from optimizing for next time step prediction is a decreasing function of the number of training time steps and depends on the model architecture. All models are approximately matched in size at 580K (K=thousand) trainable parameters. (C) Within a model class, the data scaling slopes exhibit remarkable similarity across experimental dataset sources, despite the diversity of the experimental and behavioral conditions.

**Synthetic Datasets.** We created synthetic datasets that mirror the complexity and challenges of the processed real *C.elegans* neural activity data, such as partial observability, sparsity, and noise, with 200 synthetic 'worms' each having 1500 time steps of activity from 50 randomly chosen neurons. These datasets allow for robust model validation against known dynamics, offering clear benchmarks for performance evaluation.

The **Sines** dataset models each neuron's activity as an independent sinusoid with a random phase but consistent frequency across worms, exploring an assumption of neuron-specific 'fingerprints' in the network dynamics. This dataset tests models on their ability to capture simple, uncoupled oscillatory patterns, a task suited for recurrent models like LSTMs.

Conversely, the **Random Walk** dataset challenges models with inherently unpredictable neural activity simulated through a random walk process, aiming to benchmark against the naive predictor's performance which is provably the best estimator in this case. This setup serves as a critical test of model implementations, ensuring no model unjustifiably surpasses this baseline.

# B. Model Structure

**Model architectures.** Our study utilizes three distinct classes of neural networks to harness different inductive biases for the prediction of future neural activity in *C. elegans.* These include Long-Short Term Memory (LSTM) networks, Transformer networks, and Feedforward networks. These architectures were chosen to represent a fundamental set of mechanisms—recurrence, attention, and feedforward processing—allowing us to assess the impact of structural and mechanistic differences on the task at hand.

**Shared model structure.** Each architecture is implemented within a common structural framework comprising an embedding block, a hidden 'core' module, and a linear readout layer to enable a consistent training and evaluation procedure.

This shared structure is modular allowing for the comparison of different ANN architectures by substituting only the core module (Fig. 5A).

- Embedding: The basic embedding layer linearly projects from the 302-dimensional neural state space to a higher or lower *H*-dimensional latent space. Certain architectures like the Transformer may add a positional encoding and other architectures may apply layer normalization to stabilize the learning process.
- 2) Core: The core module is architecture-specific and constitutes the primary computational engine of the model. It is restricted to use a single layer to maintain simplicity and facilitate interpretability. Besides the single layer restriction, we place no constraints on the class of computations the core is allowed to use (e.g. recurrence, parallel-processing, etc.), making our approach highly modular and scalable. The Baseline and LinearRegression prediction models are 'shallow' models, which means that they lack the 'core' module.
- 3) *Output Mapping:* The final component of the model is a linear projection from the latent space back to the original neural state space. The output of this layer is the predicted future neural activity  $\hat{\mathbf{X}} = f_{ho}(\mathbf{Z})$ , where  $f_{ho}$  is the linear transformation from the hidden to the output space.

**Causal Predictions and Temporal Memory.** Our models are tasked with making causal predictions, where future predictions do not rely on future inputs. We use a causal attention mask in the Transformer and Attention models. The LSTM and CTRNN models are inherently causal by definition. The Linear and Feedforward models lack access to temporal context beyond the current time step (i.e. they process each time point in a sequence independently). This essentially restricts the computation of the Linear and Feedforward models to feature regression (linear and nonlinear, respectively), providing a baseline for the importance of temporal information in self-supervised neural prediction.

**Baseline Model.** We use the naive predictor which posits that the next neural state will be identical to the current one as our baseline model. This baseline is a commonly used one for time series prediction tasks and it is the known optimal predictor for a random walk. Despite its simplicity, this baseline is not trivial to beat in the context of neural activity data which can often resemble, at a first-order approximation, a random process. Beating this baseline requires our ANN models to uncover and leverage complex, higher-order structures in the neural activity data beyond what is expected from a purely stochastic process. To maintain consistency with our models superstructure (Fig. 5A left), our Baseline model is implemented as 'thunk' model class with no trainable parameters that simply copies its masked input as its output.

#### C. Training Objective and Loss Function

**Training Objective.** The models are trained under a selfsupervised objective to predict the 1-time step shifted sequence of neural activity given a input sequence of neural activity of



Fig. 5. Model architecture and model scaling experiment. (A) The six (6) model classes we investigate share a common backbone but have differentiated core architectures: Feedforward, Linear, LSTM, CTRNN, Transformer, and Attention. Asterisks indicate modules appear in some architectures but not others. \*Positional encoding present here in the Transformer and Attention models. \*\*Layer normalization absent here in the Feedforward and Attention models. (B) The next time step prediction validation accuracy (inverse relation to loss) improves non-linearly with increasing the model sizes with some model architectures scaling better than others. Only the recurrent (CTRNN and LSTM) models beat the baseline loss. All models are trained and validated on the fixed largest training and validation datasets, respectively, made from  $\mathcal{D}_{ALL}$ .

length *L*. The training objective is simply to minimize the meansquared error (MSE) between the predicted and true neural activity sequences. The loss function further incorporates the boolean neuron mask to ensure that only neurons with measured data contribute to the loss computation. The mean-squared error (MSE) loss function with the boolean mask is defined as:

$$\mathcal{L}(\mathbf{X}, \mathbf{\hat{X}}; \mathbf{y}) = \frac{1}{N \times L} \sum_{i=1}^{N} \sum_{t=0}^{L-1} \mathbf{y} \odot (\mathbf{X}_i(t+1) - \mathbf{\hat{X}}_i(t+1))^2$$
(4)

where  $\mathbf{X}_i(t+1)$  is the true activity of the *i*<sup>th</sup> neuron at time t+1,  $\mathbf{\hat{X}}_i(t+1)$  is the predicted activity at time t+1,  $\mathbf{y} \in \{0, 1\}^{302}$  is the boolean feature mask indicating the presence of data for neuron *i*, *L* is the sequence length used for training the model, and  $N = \mathbb{1}^T \mathbf{y}$  is the number of masked neurons (i.e. the number of labelled neurons with data).

**Data Sampling and Model Evaluation.** We construct the training and validation sets by sampling from each worm  $n_s = 32$  sequences of length L = 180 time steps according to the method in subsection II-A **Train-Test split**. For example, a dataset  $\mathcal{D}^{n_w}$  containing  $n_w$  worms would produce training and validation sets each with  $n_w \times 32$  sequences. Train and validation data loaders use a batch size of 128.

**Training Protocol.** Models are trained up to a maximum of 500 epochs using the AdamW optimizer, with an initial learning rate of 0.001. A learning rate scheduler reduces the rate upon a validation loss plateau, with a decay factor of 0.1. Early stopping with a patience of 100 epochs is employed for efficiency. Training for all experiments was run with the same computing resources and device specifications (1 NVIDIA A100 80GB GPU).

#### III. RESULTS

## A. Data Scaling

1) Mixed Dataset Scaling: To assess how increasing the amount of training data influences the self-supervised next time step neural activity prediction.

We trained models of every architecture on incrementally larger training sets from a sequence of mixed worm datasets ranging in size from 1 to 284 worms, sampled according to subsection III-A1 **Mixed Dataset Scaling**. At each training set size, the models were evaluated against the fixed maximum sized validation set made from  $\mathcal{D}_{ALL}$ . Since there are multiple possible combinatorial assignments for a mixed dataset  $\mathcal{D}^{n_w}$ of size  $n_w < 284$  worms, we plot the independent variable as the number time-steps in the train dataset. We controlled for the hidden size of each model architecture class so that all models were approximately matched at 580K (K=thousand) trainable parameters (Table II).

TABLE II Model Hidden Sizes and Parameter Counts.

A: Parameter Counts for Hidden Size 300						
Model Name	Model Alias	Hidden Size	Parameter Count			
FeatureFFNN	Feedforward	300	272702			
NetworkCTRNN	CTRNN	300	363302			
NetworkLSTM	LSTM	300	904802			
NeuralTransformer	Transformer	300	1268402			
PureAttention	Attention	300	543602			
A comparison of the size in number of trainable parameters						

for different model architectures.

<sup>A</sup> The models are matched for the hidden size of their internal 'core' module to compare their parameter counts.

B: Matched Parameter Count (Approx. 580K)							
Model Name	Model Alias	Hidden Size	Parameter Count				
FeatureFFNN	Feedforward	516	580286				
NetworkCTRNN	CTRNN	408	582110				
NetworkLSTM	LSTM	234	582260				
NeuralTransformer	Transformer	196	584186				
PureAttention	Attention	312	580310				

<sup>B</sup> The models are approximately matched for their

parameters count to compare their hidden sizes.

The results shown in Fig 4B indicate that, at a fixed model size 0.580M trainable parameters, the CTRNN models scale to scale the best (slope = -0.65) with dataset size, whereas the Feedforward models scale the worst (slope = -0.40), out of the architecture classes investigated. We also validated our dataset scaling on the synthetic datasets introduced in subsection II-A **Synthetic Datasets** (Fig. 6A-B).

2) Individual Dataset Scaling: To determine if models trained on mixed datasets maintained consistent scaling properties when evaluated on the individual experimental source datasets.

Utilizing the best model from the mixed dataset scaling experiment at each training dataset size, we evaluated on the largest validation set made from each of the experimental sources (refer to last paragraph of subsection II-A **Mixed Datasets**).

Fig. 4C presents the results for scaling the sizes of the individual experimental source datasets.

### B. Model Scaling

To determine the effect of model complexity, as determined by the number of trainable parameters and architecture, on the performance of self-supervised neural activity prediction in *C. elegans.* 

We varied the hidden size of the 'core' architecture as a knob with which to vary the number of parameters of our models. Since each architecture (Linear, Feedforward, CTRNN, LSTM, Transformer, Attention) has a different number of trainable parameters than the others at any given hidden size, we plot the number of trainable parameters as the independent variable. The various sized models of each architecture/class were trained on the same, fixed maximum sized training dataset made from  $\mathcal{D}_{ALL}$ . We also validated our model scaling on the synthetic datasets introduced in subsection II-A **Synthetic Datasets** (Fig. 6C-D).

Fig. 5B shows the results of increasing the model size for the different architecture classes. As discussed in subsection II-B **Shared model structure**, the Baseline and Linear models have no hidden size dimension to vary. The Baseline model simply has no trainable parameters at all, whereas the Linear model has no trainable parameters in its 'core' module – which is just an Identity layer (Fig. 5A left).

The Baseline model achieved a minimum validation loss of 0.03541 which, by definition, is the baseline loss. The Linear model achieved a minimum validation loss of 0.03533. The minimum validation loss and parameter counts of the other trained models is presented in Fig. 7, along with a qualitative comparison of their auto-regressive generation capability.

## C. Synthetic Experiments

To investigate the degree to which the scaling properties observed for the real worm datasets are a function of the underlying system (*C elegans*) versus of the networks being trained.

We trained the same ANN classes and sizes used to model the real *C. elegans* neural datasets (sections III-A and III-B) on two synthetic neural datasets generated by known dynamical systems: a sinusoidal oscillator (Sines) and a stochastic process (Random Walk).

To scale the dataset sizes, for each architecture class, we sampled increasingly larger subsets of worms from each synthetic dataset (Sines or Random Walk) ranging from 1 worm to all 200 worms. Varying the number of worms in



Fig. 6. Dataset and model size scaling on synthetic datasets. (A-B) Dataset size scaling experiments. (A) Dataset size scaling of the ANN models used in 4B on the Sines dataset. Beyond a modest dataset size threshold, the sequence models – recurrent and transformer – outperform the naive predictor baseline loss. (B) Dataset size scaling of the ANN models used in 4B on the Random Walk dataset. As expected, no model beats the naive predictor baseline loss since that is the optimal predictor for a random walk. (C-D) Model size scaling experiments. (C) Model size scaling of the ANN models used in 5B on the Sines dataset. Beyond modest models used in 5B on the Random Walk dataset. No model can beat the naive predictor baseline loss. (D) Model size scaling of the ANN models used in 5B on the Random Walk dataset. No model can beat the naive predictor baseline loss, even after training on all the available data.

a dataset was an indirect way to vary the number of time steps of neural activity in the dataset since those two variables are positively correlated. The number of time steps, in turn, causally determines the amount of data used for training. To scale the model sizes, for each architecture class, we increased the number of parameters by varying the hidden size in the range [8, 4096] at log-spaced intervals. The hidden size was a proxy variable that could be manipulated to directly vary the number of trainable parameters in the models.

For the dataset size scaling experiments, all models were approximately matched in size at 580K (K=thousand) trainable parameters and the validation set used was the fixed largest possible one made from all 200 worms in the synthetic dataset. For the model size scaling experiments, the training and validation sets used were the fixed largest possible ones made from all 200 worms in the synthetic dataset.

Fig.6 presents the results for scaling the sizes of the synthetic datasets.

# IV. DISCUSSION

This study's exploration into the scaling properties of ANNs in predicting neural activity within *C. elegans* reveals critical insights into data-driven modeling of biological neural networks. Our results demonstrate that the predictive accuracy of ANNs is significantly influenced by both the volume of training data and the complexity of the model. We observed a logarithmic decrease in mean-squared error with increased training data, a consistency that persisted across various experimental datasets. This suggests a pivotal role for data volume in enhancing model performance, with further gains possible through optimal model complexity.

Model architecture emerged as a decisive factor; recurrent models, such as LSTMs and CTRNNs, outperformed others, underscoring the importance of temporal dynamics in neural activity prediction. The experiments indicated that different datasets, despite their diverse origins, share common underlying dynamics, as evidenced by the similar scaling slopes in prediction accuracy. This finding substantiates the pooling of data from multiple sources to improve model robustness.

Challenges encountered include determining appropriate model sizes for varying data volumes and incorporating behavioral contexts into predictions, which present promising avenues for future research. Our work thus far has not fully realized the long-horizon predictive capabilities for *C. elegans* neural dynamics. However, it paves the way for more comprehensive models by highlighting the potential benefits of incorporating richer datasets and nuanced model architectures.

Future efforts will focus on refining models to capture



Fig. 7. Auto-regressive generation of trained models on  $\mathcal{D}_{Uzel}$ . Models attaining the lowest validation loss for some architecture classes in Fig. 5B are seeded with a length L = 180 time step sequence from the validation set of one worm in  $\mathcal{D}_{Uzel}$  as context. The models are then made to autoregressively generate the next 180 time-steps by repeatedly predicting 1-time step ahead and appending this prediction to the end of a sliding context window.

the complexity of neural dynamics more accurately. This includes considering the integration of behaviorally-annotated data to provide additional context and investigating architectures capable of leveraging larger datasets without succumbing to the diminishing returns of over-complexity. Extension of these approaches to more complex nervous systems could offer valuable comparative insights and further the understanding of neural dynamics prediction.

In conclusion, this research contributes foundational knowledge towards the development of ANN models that more accurately reflect the intricacies of biological neural networks, bridging the gap from model organisms to broader biological contexts.

## V. Reproducibility

We have made the combined *C. elegans* neural activity dataset publicly available on the Hugging Face platform here: qsimeon/celegans\_neural\_data. All the code written for this study has been released publicly on GitHub at this repository: metaconsciousgroup/worm-graph.

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