Goal-Driven Recurrent Neural Network Models of the Ventral Visual Stream

Aran Nayebi∗, Javier Sagastuy-Brena1, Daniel M. Bear1, Kohitij Kar2, Jonas Kubilius2,3, Surya Ganguli1, David Sussillo1, James J. DiCarlo2, Daniel L. K. Yamins1

1 Stanford University, Stanford, USA
2 Massachusetts Institute of Technology, Cambridge, USA
3 KU Leuven, Leuven, Belgium

*To whom correspondence should be addressed; E-mail: anayebi@stanford.edu

The ventral visual stream (VVS) is a hierarchically connected series of cortical areas known to underlie core object recognition behaviors, enabling humans and non-human primates to effortlessly recognize objects across a multitude of viewing conditions. While recent feedforward convolutional neural networks (CNNs) provide quantitatively accurate predictions of temporally-averaged neural responses throughout the ventral pathway, they lack two ubiquitous neuroanatomical features: local recurrence within cortical areas and long-range feedback from downstream areas to upstream areas. As a result, such models are unable to account for the temporally-varying dynamical patterns thought to arise from recurrent visual circuits, nor can they provide insight into the behavioral goals that these recurrent circuits might help support. In this work, we augment CNNs with local recurrence and long-range feedback, developing convolutional RNN (ConvRNN) network models that more correctly mimic the gross neuroanatomy of the ventral pathway. Moreover,
when the form of the recurrent circuit is chosen properly, ConvRNNs with comparatively small numbers of layers can achieve high performance on a core recognition task, comparable to that of much deeper feedforward networks. We then compared these models to temporally fine-grained neural and behavioral recordings from primates to thousands of images. We found that ConvRNNs better matched these data than alternative models, including the deepest feedforward networks, on two metrics: 1) neural dynamics in V4 and inferotemporal (IT) cortex at late timepoints after stimulus onset, and 2) the varying times at which object identity can be decoded from IT, including more challenging images that take longer to decode. Moreover, these results differentiate within the class of ConvRNNs, suggesting that there are strong functional constraints on the recurrent connectivity needed to match these phenomena. Finally, we find that recurrent circuits that attain high task performance while having a smaller network size as measured by number of units, rather than another metric such as the number of parameters, are overall most consistent with these data. Taken together, our results evince the role of recurrence and feedback in the ventral pathway to reliably perform core object recognition while subject to a strong total network size constraint.

1 Introduction

The visual system of the brain must discover meaningful patterns in a complex physical world[1].

Within 200ms, primates can quickly identify objects despite changes in position, pose, contrast, background, foreground, and many other factors from one occasion to the next: a behavior known as “core object recognition”[2,3]. It is known that the ventral visual stream (VVS) underlies this ability by transforming the retinal image of an object into a new internal representation, in which high-level properties, such as object identity and category, are more explicit[3].

Task-optimized, deep convolutional neural networks (CNNs) are currently the most quanti-
tatively accurate models of encoding in primate visual cortex\textsuperscript{4,5,6}. This observed correspondence is due to their cortically-inspired architecture, which consists of a cascade of spatially-tiled linear and nonlinear operations; and their being optimized to perform the same behaviors that animals must perform to survive, such as object recognition\textsuperscript{7}. CNNs trained to recognize objects in the ImageNet dataset predict the \emph{time-averaged} neural responses of cortical neurons better than any other model class. Model units from early, intermediate, and higher convolutional layers, respectively, provide the best-known linear predictions of time-averaged visual responses in neurons of early (area V1\textsuperscript{5,8}), intermediate (area V4\textsuperscript{4}), and higher visual cortical areas (inferotemporal cortex, IT\textsuperscript{4,5}).

While these results are promising, it is not obvious how to extend the architecture and task-optimization of CNNs to the case where responses change over time. Non-trivial dynamics result from biological features not present in purely feedforward CNNs, including synapses that facilitate or depress, dense local recurrent connections within each cortical region, and long-range connections between different regions, such as feedback from higher to lower visual cortex\textsuperscript{9}. Furthermore, the behavioral roles of recurrence and dynamics in the visual system are not well understood. Several conjectures are that recurrence “fills in” missing data\textsuperscript{10,11,12,13}, such as object parts occluded by other objects; that it “sharpens” representations by top-down attentional feature refinement, allowing for easier decoding of certain stimulus properties or performance of certain tasks\textsuperscript{14,15,16,17}, that it allows the brain to “predict” future stimuli (such as the frames of a movie\textsuperscript{18,19,20}), or that recurrence “extends” a feedforward computation, reflecting the fact that an unrolled recurrent network is equivalent to a deeper feedforward network that conserves on neurons (and learnable parameters) by repeating transformations several times\textsuperscript{21,22,23,12}. Formal computational models are needed to test these hypotheses: if optimizing a model for a certain task leads to accurate predictions of neural dynamics, then that task may be a primary reason those dynamics occur in the brain.
We therefore broaden the method of goal-driven modeling from solving tasks with feedforward CNNs\(^7\) or RNNs\(^{24}\) to explain dynamics in the primate visual system, resulting in convolutional recurrent neural networks (ConvRNNs). We show that with the appropriate choice of layer-local recurrence and feedback connections, ConvRNNs can match the performance of much deeper feedforward CNNs on ImageNet but with far fewer parameters and a more anatomically consistent number of layers. Furthermore, we found that such task-optimized ConvRNNs better match the VVS than feedforward CNNs by two metrics: 1) they are able to match the fine-timescale trajectories of neural responses in the visual pathway across the entirety of stimulus presentation with a fixed linear mapping, and 2) they provide a better match to primate behavior in the form of object solution times. Specifically, we observe that ConvRNNs that attain high task performance but low network size, as measured by number of units, are most consistent with both of these metrics. These results indicate that very deep feedforward models are overall a less consistent match to primate VVS than shallower feedforward networks with added recurrence. This in turn implies that the role of recurrence in core object recognition is consistent with the hypothesis of extending a shallower feedforward network across time in order to perform a categorization task while obeying a physical size constraint.

2 Results

2.1 An evolutionary architecture search yields specific layer-local recurrent circuits and long-range feedbacks that improve task performance.

We first tested whether augmenting CNNs with standard RNN circuits from the machine learning community, SimpleRNNs and LSTMs, could improve performance on ImageNet object recognition (Figure 2a). We found that these recurrent circuits\(^{1}\) added a small amount of accuracy when introduced into the convolutional layers of a 6-layer feedforward backbone (“FF”

\(^{1}\)Adapting other recurrent cell structures to ConvRNNs from the literature, including the UGRNN and IntersectionRNN\(^{25}\), had similar effects.
in Figure 2b) based off of the AlexNet architecture, which we will refer to as a “BaseNet” (see Section A.3 for architecture details). However, there were two problems with these resultant recurrent architectures: first, these ConvRNNs did not perform substantially better than parameter-matched, minimally unrolled controls – defined as the minimum number of timesteps after the initial feedforward pass whereby all recurrence connections were engaged at least once. This control comparison suggested that the observed performance gain was due to an increase in the number of unique parameters added by the implanted ConvRNN cells rather than temporally-extended recurrent computation. Second, making the feedforward model wider or deeper yielded an even larger performance gain than adding these standard RNN cells, but with fewer parameters. This suggested that standard RNN circuits, although well-suited for a range of temporal tasks, are less well-suited for inclusion within deep CNNs to solve challenging object recognition tasks.

We speculated that this was because standard circuits lack a combination of two key properties, each of which on their own have been successful either purely for RNNs or for feedforward CNNs: (1) **Gating**, in which the value of a hidden state determines how much of the bottom-up input is passed through, retained, or discarded at the next time step; and (2) **Bypassing**, where a zero-initialized hidden state allows feedforward input to pass on to the next layer unaltered, as in the identity shortcuts of ResNet-class architectures (Figure 2a; top left). For example, LSTMs employ gating, but no bypassing, as their inputs must pass through several nonlinearities to reach their output; whereas SimpleRNNs do bypass a zero-initialized hidden state, but do not gate their input (Figure 2a).

We thus implemented recurrent circuits with both features to determine whether they function better than standard cells within CNNs. One example of such a structure is the “Reciprocal Gated Cell” (RGC), which bypasses its zero-initialized hidden state and incorporates LSTM-style gating (Figure 2a, bottom right; see Section A.3.7 for the cell equations). Adding this
cell to the 6-layer BaseNet improved performance substantially relative to both the feedforward baseline and minimally unrolled, parameter-matched control version of this model. Moreover, the RGC used substantially fewer parameters than the standard cells to achieve greater accuracy (Figure 2b).

However, it has been shown that different RNN structures can succeed or fail to perform a given task because of differences in trainability rather than differences in capacity\textsuperscript{25}. Therefore, we designed an evolutionary search to jointly optimize over both discrete choices of recurrent connectivity patterns as well as continuous choices of learning hyperparameters and weight initializations (search details in Section A.4). While a large-scale search over thousands of convolutional LSTM architectures did yield a better purely gated LSTM-based ConvRNN (“LSTM Opt”), it did not eclipse the performance of the smaller RGC ConvRNN. In fact, applying the same hyperparameter optimization procedure to the RGC ConvRNNs equally increased that architecture class’s performance and further reduced its parameter count (Figure 2b, “RGC Opt”).

If the primate visual system uses recurrence in lieu of greater network depth to perform object recognition, then a shallower recurrent model with a suitable form of recurrence should achieve recognition accuracy equal to a deeper feedforward model, albeit with temporally-fixed parameters\textsuperscript{21}. We therefore tested whether our search (depicted in Figure 2c) had identified such well-adapted recurrent architectures by fully training a representative ConvRNN, the model with the median five-epoch performance after 7000 samples. This median model (“RGC Median”) reached a final ImageNet top-1 validation accuracy nearly equal to a ResNet-34 model with nearly twice as many layers, even though the ConvRNN used only \(\sim 75\%\) as many parameters. The fully unrolled model from the random phase of the search (“RGC Random”) did not perform substantially better than the BaseNet, though the minimally unrolled control did (Figure 2d). This observation suggests that our evolutionary search strategy yielded effective recurrent architectures beyond the initial random phase of the search.
We also considered a control model (“Time Decay”) that produces temporal dynamics by learning time constants on the activations independently at each layer, rather than by learning connectivity between units. In this ConvRNN, unit activations have exponential rather than immediate falloff once feedforward drive ceases. These dynamics could arise, for instance, from single-neuron biophysics (e.g. synaptic depression) rather than interneuronal connections. However, this model did not perform any better than the feedforward BaseNet, implying that ConvRNN performance is not a trivial result of outputting a dynamic time course of responses. We further implanted other more sophisticated forms of ConvRNN cells into the BaseNet, and while this improved performance over the Time Decay model, it did not outperform the RGC Median ConvRNN despite having many more parameters (Figure 2d). Together, these results demonstrate that the RGC Median ConvRNN uses recurrent computations to subserve object recognition behavior and that particular motifs in its recurrent architecture (Figure S1), found through an evolutionary search, are required for its improved accuracy. Thus, given suitable local recurrent circuits and patterns of long-range feedback connectivity, a physically more compact, temporally-extended ConvRNN can do the same challenging object recognition task as a deeper feedforward CNN.

2.2 ConvRNNs provide an improved explanation of neural dynamics. ConvRNNs naturally produce a dynamic time series of outputs given an unchanging input stream, unlike feedforward networks. While these recurrent dynamics could be used for tasks involving time, here we optimized the ConvRNNs to perform the “static” task of object classification on ImageNet. It is possible that the primate visual system is optimized for such a task, because even static images produce reliably dynamic neural response trajectories at temporal resolutions of tens of milliseconds. The object content of some images becomes decodable from the neural population significantly later than the content of other images, even though an-
imals recognize both object sets equally well. Interestingly, late-decoding images are not well characterized by feedforward CNNs, raising the possibility that they are encoded in animals through recurrent computations \cite{17}. If this were the case, we reason then that recurrent networks trained to perform a difficult, but static object recognition task might explain neural dynamics in the primate visual system, just as feedforward models explain time-averaged responses\cite{4,5}.

Prior studies\cite{28} have directly fit recurrent parameters to neural data, as opposed to optimizing them on a task. While it is natural to try to fit recurrent parameters to the temporally-varying neural responses directly, we found that this approach suffers from a fundamental overfitting issue to the particular image statistics of the neural data collected. Specifically, we directly fit these recurrent parameters (implanted into the task-optimized feedforward BaseNet) to the dynamic firing rates of primate neurons recorded during encoding of visual stimuli. However, while these non-task optimized dynamics generalized to held-out images and neurons (Figure S2a, b), they had no longer retained performance to the original object recognition task (Figure S2c). Therefore, to avoid this issue, we instead asked whether fully task-optimized ConvRNN models (including the ones introduced in Section 2.1) could predict these dynamic firing rates from multi-electrode array recordings from the ventral visual pathway of rhesus macaques\cite{29}.

We began with the feedforward BaseNet and added a variety of ConvRNN cells, including the RGC Median ConvRNN and its counterpart generated at the random phase of the evolutionary search (“RGC Random”). All of the ConvRNNs were presented with the same images shown to the primates and collected the time series of features from each model layer. To decide which layer should be used to predict which neural responses, we fit linear models from each feedforward layer’s features to the neural population and measured where explained variance on held-out images peaked (see Section A.6 for more details). Units recorded from distinct arrays – placed in the successive V4, posterior IT (pIT), and central/anterior IT (cIT/aIT) cortical areas
of the macaque – were fit best by the successive layers of the feedforward model, respectively.

Finally, we measured how well ConvRNN features from these layers predicted the dynamics of each unit. In contrast with feedforward models fit to temporally-averaged neural responses, the linear mapping in the temporal setting must be fixed at all timepoints. The reason for this choice is that the linear mapping yields “artificial units” whose activity can change over time, but the identity of these units should not change over the course of 260 ms, as would be the case if a separate linear mapping was fit at each 10 ms timebin. This choice of a temporally-fixed linear mapping therefore maintains the physical relationship between real neurons and model neurons.

As can be seen from Figure 3, with the exception of the RGC Random ConvRNN, the ConvRNN feature dynamics fit the neural response trajectories as well as the feedforward baseline features on early phase responses (Wilcoxon test $p$-values in Table 1) and better than the feedforward baseline features for late phase responses (Wilcoxon test with Bonferroni correction $p < 0.001$), across V4, pIT, and cIT/aIT on held-out images. This observation is due to the fact that any feedforward model has the same square wave dynamics as its 100 ms visual input, so it cannot predict neural responses after image offset plus a fixed delay, corresponding to the number of layers (Figure S3, purple lines). In contrast, the activations of ConvRNN units have persistent dynamics, yielding predictions of the entire neural response trajectories.

Crucially, these predictions result from the task-optimized nonlinear dynamics from ImageNet, as both models are fit to neural data with the same form of temporally-fixed linear model with the same number of parameters. Since the initial phase of neural dynamics was well-fit by feedforward models, we asked whether the later phase could be fit by a much simpler model than any of the ConvRNNs we considered, namely the Time Decay ConvRNN with ImageNet-trained time constants at convolutional layers. If the Time Decay ConvRNN were to explain neural data as well as the other ConvRNNs, it would imply that interneuronal recurrent con-
nections are not needed to account for the observed dynamics; however, this model did not fit the late phase dynamics of intermediate areas (V4 and pIT) as well as the other ConvRNNs. Thus, the more complex recurrence found in ConvRNNs is generally needed both to improve object recognition performance and to account for neural dynamics in the ventral stream, even when animals are only required to fixate on visual stimuli. In fact, not all forms of complex recurrence are equally predictive of temporal dynamics. We found among these that the RGC Median, UGRNN, and GRU ConvRNNs attained the highest median neural predictivity for each visual area in both early and late phases, but in particular significantly outperformed the SimpleRNN ConvRNN at the late phase dynamics of these areas. We will explore this observation further in Section 2.3.

A natural follow-up question to ask is whether recurrent processing explains any more of the variance at any individual timebins than feedforward models, especially in light of the observation that there is a drop in explained variance for feedforward models from early to late timebins. It is well-known that recurrent neural networks can be viewed as very deep feedforward networks with weight sharing across layers that would otherwise be recurrently connected. In Figure 3, we present the feedforward BaseNet with a constant stream of inputs in order for it to have a consistent output throughout time, despite the ConvRNNs and the primates only being provided with a 100ms stimulus presentation. We find that while the BaseNet in this setting can slightly outperform the best ConvRNNs at the late phase dynamics (Wilcoxon test with Bonferroni correction $p < 0.001$), it underperforms relative to itself and the best ConvRNNs at the early phase dynamics (Wilcoxon test with Bonferroni correction $p < 0.001$), providing not as consistent predictions under a temporally-fixed mapping. Thus, to address this question, we compare feedforward models of varying depths (with a constant input stream) to

---

*Wilcoxon test with Bonferroni correction $p < 0.001$ for each ConvRNN vs. Time Decay, except for the SimpleRNN $p \approx 0.46$ for pIT.*

*Wilcoxon test with Bonferroni correction between each of these ConvRNNs vs. the SimpleRNN on late phase dynamics, $p < 0.001$ per visual area.*
ConvRNNs across the entire temporal trajectory under a varying linear mapping at each time-bin, in contrast to the above. Specifically, as can be seen in Figure S4a, we observe a drop in explained variance from early (130-140ms) to late (200-210ms) timebins for the shallower BaseNet and ResNet-18 models, across multiple neural datasets. Models with increased feedforward depth (such as ResNet-101 or ResNet-152), along with our performance-optimized RGC Median ConvRNN, exhibit a similar drop in median population explained variance as the shallower feedforward models. The benefit of model depth with respect to increased explained variance of late IT responses might be only noticeable while comparing very shallow models (< 7 nonlinear transforms) to much deeper (> 15 nonlinear transforms) models\textsuperscript{17}. Our results suggest that the amount of variance explained in the late IT responses is not a monotonically increasing function of model depth.

As a result, an alternative hypothesis is that the drop in explained variance from early to late timebins could instead be attributed to task-orthogonal dynamics specific to an individual primate as opposed to iterated nonlinear transforms, resulting in variability unable to be captured by any task-optimized model (feedforward or recurrent). To explore this possibility, we examined whether the model’s neural predictivity at these early and late timebins was relatively similar in ratio to that of one primate’s IT neurons mapped to that of another primate (see Section A.7 for more details). As shown in Figure S4b, across various hyperparameters of the linear mapping, we observe a ratio close to one between the neural predictivity (of the target primate neurons) of the feedforward BaseNet to that of the source primate mapped to the same target primate. Therefore, as it stands, temporally-varying linear mappings to neural responses collected from an animal during rapid visual stimulus presentation (RSVP) may not sufficiently separate feedforward models from recurrent models any better than one animal to another – though more investigation is needed to ensure tight estimates of the inter-animal consistency measure we have introduced here with neural data recorded from many primates.
Nonetheless, this observation motivates us to look beyond neural response predictions and turn to temporally-varying behavioral metrics in order to further separate these model classes, which we do next.

2.3 ConvRNNs better match temporal dynamics of primate behavior than feedforward models.

To address whether recurrent processing is engaged during core object recognition behavior, we turn to behavioral data collected from behaving primates. There is a growing body of evidence that current feedforward models fail to accurately capture primate behavior. We therefore reasoned that if recurrence is critical to core object recognition behavior, then recurrent networks should be more consistent with suitable measures of primate behavior compared to the feedforward model family. Given that the identity of different objects is decoded from the IT population at different times, we considered the first time at which the IT neural decoding accuracy reaches the (pooled) primate behavioral accuracy of a given image, known as the “object solution time (OST)”. Given that our ConvRNNs also have an output at each 10 ms timebin, the procedure for computing the OST for these models is computed from its “IT-preferred” layers, and we report the “OST consistency” which we define as the Spearman correlation between the model OSTs and the IT population’s OSTs on the common set of images solved by the given model and IT.

Unlike our ConvRNNs, which exhibit more biologically plausible temporal dynamics, evaluating the temporal dynamics in feedforward models poses an immediate problem. Given that recurrent networks repeatedly apply nonlinear transformations across time, we can analogously map the layers of a feedforward network to timepoints, observing that a network with $k$ distinct layers can produce $k$ distinct OSTs in this manner. Thus, the most direct definition of a feedforward model’s OST is to uniformly distribute the timebins between 70-260 ms across its $k$
layers. For very deep feedforward networks such as ResNet-101 and ResNet-152, this number of distinct layers will be as fine-grained as the 10 ms timebins of the IT responses; however, for most other shallower feedforward networks this will be much coarser. Therefore to enable shallow feedforward models to be maximally temporally expressive, we also randomly sample units from consecutive feedforward layers to produce a more graded temporal mapping, depicted in Figure 4b. This graded mapping is ultimately what we use for the feedforward models in Figure 4d, providing the highest OST consistency for that model class.

With model OST defined across both model families, we compared various ConvRNNs and feedforward models to the IT population’s OST in Figure 4d. Among shallower and deeper models, we found that ConvRNNs were generally able to better explain IT’s OST than their feedforward counterparts. Specifically, we found that ConvRNN cells without any multi-unit interaction such as the Time Decay ConvRNN only marginally, and not always significantly, improved the OST consistency over its respective BaseNet model. On the other hand, consistent with our prior observation in Figure 3, ConvRNNs with multi-unit interactions generally provided the greatest match to IT OSTs than even the deepest feedforward models.

Consistent with our observations in Figures 2 and 3 that different recurrent cells with multi-unit interactions were not all equally effective when embedded in CNNs (despite outperforming the simple Time Decay model), we similarly found that this observation held for the case of matching IT’s OST. Given recent observations that inactivating parts of macaque ventrolateral PFC (vlPFC) results in behavioral deficits in IT for late-solved images, we reasoned that additional decoding procedures employed at the categorization layer during task optimization

\[\text{Wilcoxon test on uniform vs. graded mapping OST consistencies across feedforward models, } p < 0.001; \text{ see also Figure S5.}\]

\[^d\text{Paired t-test with Bonferroni correction: Shallow Time Decay vs. “BaseNet” in blue, } t(9) \approx 3.23, p < 0.025; \text{ Deeper Time Decay vs. “BaseNet” in red, } t(9) \approx 1.73, p \approx 0.11.}\]

\[^e\text{Paired t-test with Bonferroni correction: Shallow RGC vs. “BaseNet” in blue, } t(9) \approx 6.08, p < 0.001; \text{ Deeper UGRNN vs. ResNet-152, } t(9) \approx 7.55, p < 0.001; \text{ Deeper GRU vs. ResNet-152, } t(9) \approx 7.71, p < 0.001; \text{ RGC Median vs. ResNet-152, } t(9) \approx 3.44, p < 0.01.}\]

13
might meaningfully impact the model’s OST consistency, in addition to the choice of recurrent cell used. We designed several decoding procedures (defined in Section A.5), motivated by prior observations of accumulation of relevant sensory signals during decision making in primates\(^{32}\). Overall, we found that ConvRNNs with different decoding procedures, but with the same layer-local recurrent cell (RGC Median) and long-range feedback connectivity patterns, yielded significant differences in final consistency with the IT population OST (Friedman test, \(p < 0.05\)). Moreover, the simplest decoding procedure of outputting a prediction at the last timepoint, a strategy commonly employed by the computer vision community, had a lower OST consistency than each of the more nuanced Max Confidence\(^{g}\) and Threshold decoding procedures\(^{h}\) that we considered. Taken together, our results suggest that the type of multi-unit layer-wise recurrence and downstream decoding strategy are important features for OST consistency with IT, suggesting that specific, non-trivial connectivity patterns further downstream the ventral stream may be important to core object recognition behavior over timescales of a couple hundred milliseconds.

### 2.4 ConvRNNs mediate a tradeoff between task performance and network size.

Why might a suitably shallower feedforward network with temporal dynamics be desirable for the ventral visual stream? We reasoned that recurrence mediates a tradeoff between network size and task performance; a tradeoff that the ventral stream also maintains. To examine this possibility, in Figure 5 we compared each network’s task performance versus its size, measured either by parameter count or unit count. Across models, we found unit count (related to the number of neurons) to be more consistent with task performance than parameter count (related to the number of synapses). In fact, there are many models with a large parameter

\(^{g}\)Paired \(t\)-test with Bonferroni correction, \(t(9) \approx -4.52, p < 0.01\).

\(^{h}\)Paired \(t\)-test with Bonferroni correction, \(t(9) \approx -4.41, p < 0.01\).
count but not very good task performance, indicating that adding synapses is not necessarily as useful for performance as adding neurons. For shallower recurrent networks, task performance seemed to be more strongly associated with OST consistency than network size, though generally having fewer parameters at a given performance level resulted in higher OST consistency (e.g., UGRNN vs. SimpleRNN and RGC vs. IntersectionRNN). This tradeoff became more salient for deeper feedforward models and the deeper ConvRNNs, as the very deep ResNets (ResNet-34 and deeper) attained an overall lower OST consistency compared to the deeper ConvRNNs, using both much more units and parameters compared to small relative gains in task performance. Similarly, deeper ConvRNNs with high task performance and minimal unit count, such as the UGRNN, GRU, and RGCs attained both the highest OST consistency overall (Figures 4 and 5) along with providing the best match to neural dynamics across visual areas (Figure 3). This observation indicates that suitably-chosen recurrence can provide a means for maintaining this fundamental tradeoff.

Given that specific forms of task-optimized recurrence are more consistent with IT’s OST than iterated feedforward transformations (with unshared weights), we asked whether it was possible to approximate the effect of recurrence with a feedforward model. This approximation would allow us to better describe the additional “action” that recurrence is providing in its improved OST consistency. Furthermore, a crucial difference between this metric and the explained variance metric evaluated on neural data in the prior section is that the latter uses a linear transform from model features to neural responses, whereas the former operates directly on the original model features. Therefore, a related question is whether the use of a linear transform for mapping from model units to neural responses masks the improvement that recurrent processing can have over deep feedforward models in their original feature space.

To address these questions, we trained a separate linear mapping from each model layer to the corresponding IT response at the given timepoint, on a set of images distinct from those on
which the OST consistency metric is evaluated on. Overall, as depicted in Figure S5, we found that models with less temporal variation in their source features (namely those under a uniform mapping with few “IT-preferred” layers) had significantly improved OST consistency with their linearly transformed features (Wilcoxon test, \( p < 0.001 \)), whereas models with the maximum amount of temporal variation such as ResNet-101, ResNet-152, and the ConvRNNs had a significant reduction in OST consistency with their linearly transformed features (Wilcoxon test, \( p < 0.001 \)), indicating the harmful dimensionality reduction of the linear mapping. On the other hand, the linearly transformed shallower variants of deeper feedforward models were not significantly different from task-optimized ConvRNNs that achieved high OST consistency\(^1\), suggesting that the action of suitable task-optimized recurrence approximates that of a shallower feedforward model with linearly induced neural dynamics.

**Discussion**

The overall goal of this study is to determine what role recurrent circuits may have in the execution of core object recognition behavior in the ventral stream. By broadening the method of goal-driven modeling from solving tasks with feedforward CNNs to ConvRNNs that include layer-local recurrence and feedback connections, we first demonstrate that appropriate choices of these recurrent circuits which incorporate specific principles of “gating” and “bypassing” lead to matching the task performance of much deeper feedforward CNNs with fewer units and parameters. Moreover, unlike deep feedforward CNNs, the mapping from the early, intermediate, and higher layers of these shallower ConvRNNs to corresponding cortical areas is neuroanatomically consistent and reproduces prior quantitative properties of the ventral stream.

\(^1\)Paired \( t \)-test with Bonferroni correction: RGC Median vs. PLS Uniform BaseNet, \( t(9) \approx -0.86, p \approx 0.41 \); RGC Median with Threshold Decoder vs. PLS Uniform ResNet-18, \( t(9) \approx 0.82, p \approx 0.43 \); RGC Median with Max Confidence Decoder vs. PLS Uniform ResNet-34, \( t(9) \approx 0.02, p \approx 0.99 \).
response trajectories at temporal resolutions of tens of milliseconds throughout the ventral visual hierarchy, unlike feedforward models or certain other choices for recurrence adapted to solving visual recognition problems such as Temporal Decay or the LSTM circuit. In fact, ConvRNNs with high task performance but small network size (as measured by number of neurons rather than synapses) are not only the most quantitatively accurate models of neural response trajectories during passive viewing but also are most consistent with the temporal evolution of primate IT object identity solutions during active task performance. Taken together, our results suggest that recurrence in the ventral stream mediates a tradeoff between task performance and neuron count, suggesting that the computer vision community’s solution of stacking more feedforward layers to solve challenging visual recognition problems approximates what is compactly implemented in the primate visual system by leveraging additional nonlinear temporal transformations to the initial feedforward IT response. This work therefore provides a quantitative prescription for the next generation of dynamic ventral stream models, addressing the call to action in a recent previous study\textsuperscript{17} for a change in architecture from feedforward models.

Many hypotheses about the role of recurrence in vision have been put forward, particularly in regards to overcoming certain challenging image properties\textsuperscript{10,11,12,13,9,14,15,16,17,18,19,20}. We believe this is the first work to address the role of recurrence at scale by connecting novel \textit{task-optimized} recurrent models to temporal metrics defined on high-throughput neural and behavioral data. Moreover, these metrics are well-defined for feedforward models (unlike prior work\textsuperscript{13}) and therefore meaningfully demonstrate a separation between the two model classes.

Though our results help to clarify the role of recurrence during core object recognition behavior, many major questions remain. Our work addresses why the visual system may leverage recurrence to subserve visually challenging behaviors, replacing a physically implausible architecture (deep feedforward CNNs) with one that is ubiquitously consistent with anatomical ob-
servations (shallower ConvRNNs). However, our work does not address gaps in understanding either the loss function or the learning rule of the neural network. Specifically, we intentionally implant layer-local recurrence and long-range feedback connections into feedforward networks that have been useful for supervised learning via backpropagation on ImageNet. A natural next step would be to connect these ConvRNNs with unsupervised objectives, as has been done for feedforward models of the ventral stream in concurrent work\textsuperscript{34}. The question of biologically plausible learning targets is similarly linked to biologically plausible mechanisms for learning such objective functions. Recurrence could play a separate role in implementing the propagation of error-driven learning, obviating the need for some of the issues with backpropagation (such as weight transport), as has been recently demonstrated at scale\textsuperscript{35,36}. Therefore, building ConvRNNs with unsupervised objective functions optimized with biologically-plausible learning rules would be essential towards a more complete goal-driven theory of visual cortex.

Additionally, high-throughput experimental data will also be critical to further separate hypotheses about recurrence. While we see evidence of recurrence as mediating a tradeoff between network size and task performance for core object recognition, it could be that recurrence plays a more task-specific role under more temporally dynamic behaviors. Not only would it be an interesting direction to optimize ConvRNNs on more temporally dynamic visual tasks than ImageNet, but to compare to neural and behavioral data collected from such stimuli, potentially over longer timescales than 200\textit{ms} while the animal is performing a task. Such models and experimental data would synergistically provide great insight into how rich visual behaviors proceed, while also inspiring better computer vision algorithms.

References

[1] James, W. The principles of psychology (vol. 1). \textit{New York: Holt} \textbf{474} (1890).
[2] Pinto, N., Cox, D. D. & DiCarlo, J. J. Why is real-world visual object recognition hard? *PLoS Computational Biology* (2008).

[3] DiCarlo, J. J., Zoccolan, D. & Rust, N. C. How does the brain solve visual object recognition? *Neuron* 73, 415–434 (2012).

[4] Yamins, D. L. K. *et al.* Performance-optimized hierarchical models predict neural responses in higher visual cortex. *Proceedings of the National Academy of Sciences* 111, 8619–8624 (2014). [http://www.pnas.org/content/111/23/8619.full.pdf](http://www.pnas.org/content/111/23/8619.full.pdf).

[5] Khaligh-Razavi, S.-M. & Kriegeskorte, N. Deep supervised, but not unsupervised, models may explain it cortical representation. *PLoS computational biology* 10, e1003915 (2014).

[6] Gückü, U. & van Gerven, M. A. Deep neural networks reveal a gradient in the complexity of neural representations across the ventral stream. *The Journal of Neuroscience* 35, 10005–10014 (2015).

[7] Yamins, D. L. & DiCarlo, J. J. Using goal-driven deep learning models to understand sensory cortex. *Nature neuroscience* 19, 356 (2016).

[8] Cadena, S. A. *et al.* Deep convolutional models improve predictions of macaque v1 responses to natural images. *PLoS computational biology* 15, e1006897 (2019).

[9] Gilbert, C. D. & Wu, L. Top-down influences on visual processing. *Nat. Rev. Neurosci.* 14, 350–363 (2013).

[10] Spoerer, C. J., McClure, P. & Kriegeskorte, N. Recurrent convolutional neural networks: a better model of biological object recognition. *Front. Psychol.* 8, 1–14 (2017).
[11] Michaelis, C., Bethge, M. & Ecker, A. One-shot segmentation in clutter. In *International Conference on Machine Learning*, 3549–3558 (PMLR, 2018).

[12] Rajaei, K., Mohsenzadeh, Y., Ebrahimpour, R. & Khaligh-Razavi, S.-M. Beyond core object recognition: Recurrent processes account for object recognition under occlusion. *PLoS computational biology* 15, e1007001 (2019).

[13] Linsley, D., Kim, J., Veerabadran, V., Windolf, C. & Serre, T. Learning long-range spatial dependencies with horizontal gated recurrent units. In Bengio, S. *et al.* (eds.) *Advances in Neural Information Processing Systems*, vol. 31 (Curran Associates, Inc., 2018). URL [https://proceedings.neurips.cc/paper/2018/file/ec8956637a99787bd197eacd77acce5e-Paper.pdf](https://proceedings.neurips.cc/paper/2018/file/ec8956637a99787bd197eacd77acce5e-Paper.pdf).

[14] Lindsay, G. W. Feature-based attention in convolutional neural networks. *arXiv preprint arXiv:1511.06408* (2015).

[15] McIntosh, L., Maheswaranathan, N., Sussillo, D. & Shlens, J. Recurrent segmentation for variable computational budgets. In *Proceedings of the IEEE Conference on Computer Vision and Pattern Recognition Workshops*, 1648–1657 (2018).

[16] Li, X., Jie, Z., Feng, J., Liu, C. & Yan, S. Learning with rethinking: recurrently improving convolutional neural networks through feedback. *Pattern Recognition* 79, 183–194 (2018).

[17] Kar, K., Kubilius, J., Schmidt, K., Issa, E. B. & DiCarlo, J. J. Evidence that recurrent circuits are critical to the ventral stream’s execution of core object recognition behavior. *Nature neuroscience* 22, 974–983 (2019).

[18] Rao, R. P. & Ballard, D. H. Predictive coding in the visual cortex: a functional interpretation of some extra-classical receptive-field effects. *Nature neuroscience* 2, 79–87 (1999).
[19] Lotter, W., Kreiman, G. & Cox, D. Deep predictive coding networks for video prediction and unsupervised learning. In ICLR (2017).

[20] Issa, E. B., Cadieu, C. F. & DiCarlo, J. J. Neural dynamics at successive stages of the ventral visual stream are consistent with hierarchical error signals. Elife 7, e42870 (2018).

[21] Liao, Q. & Poggio, T. Bridging the gaps between residual learning, recurrent neural networks and visual cortex. arXiv preprint arXiv:1604.03640 (2016).

[22] Zamir, A. R. et al. Feedback networks. In CVPR (2017).

[23] Leroux, S. et al. Iamnn: iterative and adaptive mobile neural network for efficient image classification. In ICLR Workshop 2018 (2018).

[24] Mante, V., Sussillo, D., Shenoy, K. V. & Newsome, W. T. Context-dependent computation by recurrent dynamics in prefrontal cortex. Nature 503, 78–84 (2013).

[25] Collins, J., Sohl-Dickstein, J. & Sussillo, D. Capacity and trainability in recurrent neural networks. In ICLR (2017).

[26] Krizhevsky, A., Sutskever, I. & Hinton, G. E. Imagenet classification with deep convolutional neural networks. In Pereira, F., Burges, C. J. C., Bottou, L. & Weinberger, K. Q. (eds.) Advances in Neural Information Processing Systems, vol. 25 (Curran Associates, Inc., 2012). URL https://proceedings.neurips.cc/paper/2012/file/c399862d3b9d6b76c8436e924a68c45b-Paper.pdf

[27] Nayebi, A. et al. Task-driven convolutional recurrent models of the visual system. In Bengio, S. et al. (eds.) Advances in Neural Information Processing Systems, vol. 31 (Curran Associates, Inc., 2018). URL https://proceedings.neurips.cc/paper/2018/file/6be93f7a96fed60c477d30a1dfe032fd-Paper.pdf
[28] Kietzmann, T. C. et al. Recurrence is required to capture the representational dynamics of the human visual system. *Proceedings of the National Academy of Sciences* **116**, 21854–21863 (2019).

[29] Majaj, N. J., Hong, H., Solomon, E. A. & DiCarlo, J. J. Simple learned weighted sums of inferior temporal neuronal firing rates accurately predict human core object recognition performance. *Journal of Neuroscience* **35**, 13402–13418 (2015).

[30] Rajalingham, R. et al. Large-scale, high-resolution comparison of the core visual object recognition behavior of humans, monkeys, and state-of-the-art deep artificial neural networks. *Journal of Neuroscience* **38**, 7255–7269 (2018).

[31] Kar, K. & DiCarlo, J. J. Fast recurrent processing via ventrolateral prefrontal cortex is needed by the primate ventral stream for robust core visual object recognition. *Neuron* **109**, 164–176 (2021).

[32] Shadlen, M. N. & Newsome, W. T. Neural basis of a perceptual decision in the parietal cortex (area lip) of the rhesus monkey. *Journal of neurophysiology* **86**, 1916–1936 (2001).

[33] Kubilius, J. et al. Brain-like object recognition with high-performing shallow recurrent anns. In Wallach, H. et al. (eds.) *Advances in Neural Information Processing Systems*, vol. 32 (Curran Associates, Inc., 2019). URL https://proceedings.neurips.cc/paper/2019/file/7813d1590d28a7dd372ad54b5d29d033-Paper.pdf.

[34] Zhuang, C. et al. Unsupervised neural network models of the ventral visual stream. *Proceedings of the National Academy of Sciences* **118** (2021).

[35] Akrout, M., Wilson, C., Humphreys, P., Lillicrap, T. & Tweed, D. B. Deep learning without weight transport. In Wallach, H. et al. (eds.) *Advances*
in Neural Information Processing Systems, vol. 32 (Curran Associates, Inc., 2019). URL https://proceedings.neurips.cc/paper/2019/file/f387624df552cea2f369918c5e1e12bc-Paper.pdf.

[36] Kunin, D. et al. Two routes to scalable credit assignment without weight symmetry. In International Conference on Machine Learning, 5511–5521 (PMLR, 2020).

[37] Abadi, M. et al. Tensorflow: A system for large-scale machine learning. In OSDI, vol. 16, 265–283 (2016).

[38] Mizuseki, K., Sirota, A., Pastalkova, E. & Buzsáki, G. Theta oscillations provide temporal windows for local circuit computation in the entorhinal-hippocampal loop. Neuron 267–280 (2009).

[39] Sutskever, I., Martens, J., Dahl, G. & Hinton, G. On the importance of initialization and momentum in deep learning. In International conference on machine learning, 1139–1147 (PMLR, 2013).

[40] He, K., Zhang, X., Ren, S. & Sun, J. Deep residual learning for image recognition. In Proceedings of the IEEE conference on computer vision and pattern recognition, 770–778 (2016).

[41] Clevert, D.-A., Unterthiner, T. & Hochreiter, S. Fast and accurate deep network learning by exponential linear units (elus). In ICLR (2016).

[42] Ioffe, S. & Szegedy, C. Batch normalization: Accelerating deep network training by reducing internal covariate shift. In International Conference on Machine Learning, 448–456 (PMLR, 2015).
[43] Bengio, Y., Simard, P. & Frasconi, P. Learning long-term dependencies with gradient
descent is difficult. *IEEE transactions on neural networks* **5**, 157–166 (1994).

[44] Ba, J. L., Kiros, J. R. & Hinton, G. E. Layer normalization. *arXiv preprint*
arXiv:1607.06450 (2016).

[45] Cho, K., van Merriënboer, B., Bahdanau, D. & Bengio, Y. On the properties of neural ma-
machine translation: Encoder–decoder approaches. In *Proceedings of SSST-8, Eighth Work-
shop on Syntax, Semantics and Structure in Statistical Translation*, 103–111 (Association
for Computational Linguistics, Doha, Qatar, 2014). URL https://www.aclweb.
org/anthology/W14-4012.

[46] Hochreiter, S. & Schmidhuber, J. Long short-term memory. *Neural Computation* **9**, 1735–
1780 (1997).

[47] Jozefowicz, R., Zaremba, W. & Sutskever, I. An empirical exploration of recurrent
network architectures. In *International Conference on Machine Learning*, 2342–2350
(PMLR, 2015).

[48] Gers, F. A., Schraudolph, N. N. & Schmidhuber, J. Learning precise timing with lstm
recurrent networks. *Journal of machine learning research* **3**, 115–143 (2002).

[49] Bergstra, J., Bardenet, R., Bengio, Y. & Kégl, B. Algorithms for hyper-parameter opti-
mization. In Shawe-Taylor, J., Zemel, R., Bartlett, P., Pereira, F. & Weinberger, K. Q.
(eds.) *Advances in Neural Information Processing Systems*, vol. 24 (Curran Associates,
Inc., 2011). URL https://proceedings.neurips.cc/paper/2011/file/
86e8f7ab32cfd12577bc2619bc635690-Paper.pdf.

[50] Bergstra, J., Komer, B., Eliasmith, C., Yamins, D. & Cox, D. D. Hyperopt: a python
library for model selection and hyperparameter optimization. *Computational Science &

*Discovery* 8 (2015).

[51] He, K., Zhang, X., Ren, S. & Sun, J. Delving deep into rectifiers: Surpassing human-

level performance on imagenet classification. In *Proceedings of the IEEE international

cference on computer vision*, 1026–1034 (2015).

[52] Nesterov, Y. A method of solving a convex programming problem with convergence rate

\(o(1/k^2)\). In *Sov. Math. Dokl*, vol. 27.

[53] Klindt, D., Ecker, A. S., Euler, T. & Bethge, M. Neural system identification

for large populations separating “what” and “where”. In Guyon, I. *et al.* (eds.)

*Advances in Neural Information Processing Systems*, vol. 30 (Curran Associates,

Inc., 2017). URL [https://proceedings.neurips.cc/paper/2017/file/8c249675aea6c3cbd91661bbae767ff1-Paper.pdf](https://proceedings.neurips.cc/paper/2017/file/8c249675aea6c3cbd91661bbae767ff1-Paper.pdf).

[54] Kingma, D. P. & Ba, J. Adam: A method for stochastic optimization. In *ICLR* (2015).

[55] Srivastava, N., Hinton, G., Krizhevsky, A., Sutskever, I. & Salakhutdinov, R. Dropout: a

simple way to prevent neural networks from overfitting. *The journal of machine learning

research* 15, 1929–1958 (2014).
Acknowledgements

We thank Tyler Bonnen and Eshed Margalit for comments on this manuscript. We thank the Google TensorFlow Research Cloud (TFRC) team for generously providing TPU hardware resources for this project. D.L.K.Y is supported by the James S. McDonnell Foundation (Understanding Human Cognition Award Grant No. 220020469), the Simons Foundation (Collaboration on the Global Brain Grant No. 543061), the Sloan Foundation (Fellowship FG-2018-10963), the National Science Foundation (RI 1703161 and CAREER Award 1844724), the DARPA Machine Common Sense program, and hardware donation from the NVIDIA Corporation. This work is also supported in part by Simons Foundation grant SCGB-542965 (J.J.D. & D.L.K.Y.). This project has received funding from the European Union’s Horizon 2020 research and innovation programme under grant agreement No. 70549 (J.K.). J.S. is supported by the Mexican National Council of Science and Technology (CONACYT).

Author Contributions

A.N. and D.L.K.Y. designed the experiments. A.N., J.S., and D.B. conducted the experiments, and A.N. analyzed the data. K.K. contributed neural data, and J.K. contributed to initial code development. K.K. and J.J.D. provided technical advice on neural predictivity metrics. D.S. and S.G. provided technical advice on recurrent neural network training. A.N. and D.L.K.Y. interpreted the data and wrote the paper.

Competing Interest Declaration

The authors declare no competing interests.
ConvRNN Cells

1. Performance-Optimized Recurrence

2. Linear Mapping to Temporal Averages

3. Temporally-Fixed Linear Mapping to 10ms Binned Dynamics

Figure 1: ConvRNNs as models of the primate ventral visual stream. Performance-optimized recurrence. Convolutional recurrent networks (ConvRNNs) have a combination of local recurrent cells (green) and long-range feedback connections (red) added on top of a feedforward CNN “BaseNet” backbone (blue). In our implementation displayed on the top, propagation along each arrow takes one time step (10ms) to mimic conduction delays between cortical layers. In addition, we consider particular choices of “light-weight” (in terms of parameter count) decoding strategy that determines the final object category of that image. Linear mapping to temporal averages. We stipulated that units from each multi-unit array must be fit by features from a single model layer. To determine which one, we fit the features from the feedforward backbone to the unit’s time-averaged response, and counted how many units had minimal loss for a given model layer, detailed in Section A.6.2. Temporally-fixed linear mapping to 10ms binned dynamics. The ConvRNN model features produce a temporally-varying output that is mapped linearly to temporally-varying neural responses in V4 and IT, under a temporally-fixed mapping whose parameters are reused throughout the entire timecourse of stimulus presentation.
Figure 2: **Suitably-chosen ConvRNNs can match the object recognition performance of much deeper feedforward models.** (a) Architectural differences between ConvRNN cells. Standard ResNet blocks and SimpleRNN cells have bypassing but no gating. The LSTM cell has gating, denoted by T-junctions, but no bypassing. The Reciprocal Gated Cell has both. (b) Performance of various ConvRNN and feedforward models as a function of number of parameters. Colored points incorporate the respective ConvRNN cell into the the 6-layer feedforward BaseNet architecture (“FF”). Here “T” denotes number of timesteps the model is unrolled for, corresponding to the propagation of a single layer to the next. Hyperparameter-optimized versions of the LSTM (“LSTM Opt”) and Reciprocal Gated Cell ConvRNNs (“RGC Opt”) are connected to their non-optimized versions by black lines. (c) ConvRNN cell search. Each blue dot represents a model, sampled from hyperparameter space, trained for 5 epochs. The orange line is the average performance of the last 50 models up to that time. The red line denotes the top performing model at that point in the search. Search space schematic: Question marks denote optional connections, which may be conventional or depth-separable convolutions with a choice of kernel size. (d) Performance of models fully trained on ImageNet. We compared the performance of an 11-layer feedforward base model (“BaseNet”) modeled after ResNet-18, a control ConvRNN model with trainable time constants (“Time Decay”), along with various other common RNN architectures implanted into this BaseNet, as well as the median Reciprocal Gated Cell (RGC) model from the search (“RGC Median”) with or without global feedback connectivity, and its minimally-unrolled control ($T = 12$). The “RGC Random” model was selected randomly from the initial, random phase of the model search. Parameter and unit counts (total number of neurons in the output of each layer) in millions are shown on top of each bar.
Figure 3: **Suitably-chosen ConvRNN circuits provide consistent predictions of primate ventral stream neural dynamics.** The y-axis indicates the median across neurons of the explained variance between predictions and ground-truth responses on held-out images divided by the square root of the internal consistencies of the neurons, defined in Section A.6.3. Error bars indicates the s.e.m across neurons (N = 88 for V4, N = 88 for pIT, N = 80 for cIT/aIT) averaged across 10ms timebins (N = 4 each for the “Early” and “Late” designations). As can be seen, the feedforward BaseNet model (first bars) is incapable of generating a response beyond the feedforward pass, but certain types of ConvRNN cells (such as “RGC Median”, “UGRNN”, and “GRU”) added to the feedforward model are overall best predictive across visual areas at late timepoints (Wilcoxon test (with Bonferroni correction) with feedforward BaseNet, p < 0.001 for each visual area). “BaseNet (Constant)” refers to the same feedforward model but presented with a constant image presentation, in contrast to the other models which are given the same 100ms stimulus presentation as the primate. See Figure S3 for the full timecourses at the resolution of 10ms bins.
Figure 4: ConvRNNs explain the object solution times (OST) of IT across images. (a) Mapping model layers to timepoints. In order to compare to primate IT object solution times, namely the first time at which the neural decode accuracy for each image reached the level of the (pooled) primate behavioral accuracy, we first need to define object solution times for models. This procedure involves identification of the “IT-preferred” layer(s) via a standard linear mapping to temporally averaged IT responses. (b) Choosing a temporal mapping gradation. These “IT-preferred” model layer(s) are then mapped to 10ms timebins from 70-260ms in either a uniform or graded fashion, if the model is feedforward. For ConvRNNs, this temporal mapping is always one-to-one with these 10ms timebins. (c) Defining model OSTs. Once the temporal mapping has been defined, we train a linear SVM at each 10ms model timebin and compute the classifier’s $d'$ (displayed in each of the black dots for a given example image). The first timebin at which the model $d'$ matches the primate’s accuracy is defined as the model OST for that image (obtained via linear interpolation). (d) Proper choices of recurrence best match IT OSTs. Mean and s.e.m. are computed across train/test splits ($N = 10$) when that image (of 1320 images) was a test-set image, with the Spearman correlation computed with the IT object solution times (analogously computed from the IT population responses) across the imageset solved by both the given model and IT, constituting the “Fraction of IT Solved Images” on the x-axis. We start with either a shallower base feedforward model consisting of 5 convolutional layers and 1 layer of readout (“BaseNet” in blue) as well as a deeper variant with 10 feedforward layers and 1 layer of readout (“BaseNet” in red), detailed in Section A.2.1. From these base feedforward models, we embed recurrent cells, resulting in either “Shallow ConvRNNs” or “Deeper ConvRNNs”.
Figure 5: **ConvRNNs with highest OST consistency conserve on network size while maintaining task performance.** Across all models considered, the Deeper ConvRNNs (denoted by “x”) that attain high categorization performance (x-axis) while maintaining a low unit count (panel B) rather than parameter count (panel A) for their given performance level, achieve the highest mean OST consistency (Spearman correlation with IT population OST, averaged across $N = 10$ train/test splits). The colorbar indicates this mean OST consistency (monotonically increasing from light to dark), binned into 6 equal ranges ($0.04478253 − 0.0769489$, $0.0769489 − 0.10911527$, $0.10911527 − 0.14128164$, $0.14128164 − 0.17344802$, $0.17344802 − 0.20561439$, and $0.20561439 − 0.23778076$). Models with a larger size at a fixed performance level are less consistent with primate object recognition behavior (e.g. deep feedforward models, denoted by boxes), with recurrence maintaining a fundamental tradeoff between network size and task performance.
A Methods

A.1 Model framework

A.1.1 Software package

To explore the architectural space of ConvRNNs and compare these models with the primate visual system, we used the Tensorflow library\textsuperscript{37} to augment standard CNNs with both local and long-range recurrence (Figure 1). Conduction from one area to another in visual cortex takes approximately 10\textit{ms}\textsuperscript{38}, with signal from photoreceptors reaching IT cortex at the top of the ventral stream by 70-100\textit{ms}. Neural dynamics indicating potential recurrent connections take place over the course of 100-260\textit{ms}\textsuperscript{20}. A single feedforward volley of responses thus cannot be treated as if it were instantaneous relative to the timescale of recurrence and feedback. Hence, rather than treating each entire feedforward pass from input to output as one integral time step, as is normally done with RNNs\textsuperscript{10}, each time step in our models corresponds to a single feedforward layer processing its input and passing it to the next layer. This choice required an unrolling scheme different from that used in the standard Tensorflow RNN library, the code for which (and for all of our models) can be found at in our TNN Github repository: [https://github.com/neuroailab/tnn](https://github.com/neuroailab/tnn).

A.1.2 Defining ConvRNNs

Within each ConvRNN layer, feedback inputs from higher layers are resized to match the spatial dimensions of the feedforward input to that layer. Both types of input are processed by standard 2-D convolutions. If there is any local recurrence at that layer, the output is next passed to the recurrent cell as input. Feedforward and feedback inputs are combined within the recurrent cell by spatially resizing the feedback inputs (via bilinear interpolation) and concatenating these with the feedforward input across the channel dimension. We let $\oplus$ denote this concatenation along the channel dimension with appropriate resizing to align spatial dimensions. Finally, the
output of the cell is passed through any additional nonlinearities, such as max-pooling. The generic update rule for the discrete-time trajectory of such a network is thus:

\[ h_\ell^t = C_\ell \left( F_\ell \left( \bigoplus_{j \neq \ell} r_j^t \right), h_{\ell-1}^t \right) \]

\[ r_\ell^t = A_\ell (h_\ell^t), \]

where \( r_\ell^t \) is the output of layer \( \ell \) at time \( t \), \( h_{\ell-1}^t \) is the hidden state of the locally recurrent cell \( C_\ell \) at time \( t - 1 \), and \( A_\ell \) is the activation function and any pooling post-memory operations. The learned parameters of such a network consist of \( F_\ell \), comprising any feedforward and feedback connections coming into layer \( \ell = 1, \ldots, L \), and any of the learned parameters associated with the local recurrent cell \( C_\ell \).

In this work, all forms of recurrence add parameters to the feedforward base model. Because this could improve task performance for reasons unrelated to recurrent computation, we trained two types of control model to compare to ConvRNNs:

1. Feedforward models with more convolution filters (“wider”) or more layers (“deeper”) to approximately match the number of parameters in a recurrent model.

2. Replicas of each ConvRNN model unrolled for a minimal number of time steps, defined as the number that allows all model parameters to be used at least once. A minimally unrolled model has exactly the same number of parameters as its fully unrolled counterpart, so any increase in performance from unrolling longer can be attributed to recurrent computation. Fully and minimally unrolled ConvRNNs were trained with identical learning hyperparameters.

### A.1.3 Training Procedure

All models (both feedforward and ConvRNN) used the standard ResNet preprocessing provided by TensorFlow here: [https://github.com/tensorflow/tpu/blob/master/models/](https://github.com/tensorflow/tpu/blob/master/models/).
Furthermore, they were trained on 224 pixel ImageNet with stochastic gradient descent with momentum (SGDM)\textsuperscript{39}, using a momentum value of 0.9.

We allowed the base learning rate, batch size, and L2 regularization strength to vary for each model, depending on what was optimal in terms of top-1 validation accuracy for that model. All models (except for AlexNet) used the ResNet training schedule\textsuperscript{40}, whereby the base learning rate is decayed by 90\% at 30, 60, and 80 epochs, training for 90 epochs total. The AlexNet had its base learning rate of 0.01 subsequently decayed to 0.005, 0.001, and 0.0005, at 30, 60, and 80 epochs, respectively. We list these values for each model in the table below:

| Model Class         | Base Learning Rate | Batch Size | L2 Regularization |
|---------------------|--------------------|------------|-------------------|
| AlexNet             | 0.01               | 1024       | $5 \times 10^{-4}$ |
| 6-layer BaseNet     | 0.01               | 256        | $1 \times 10^{-4}$ |
| Shallow ConvRNNs    | 0.01               | 256        | $1 \times 10^{-4}$ |
| 11-layer BaseNet    | 0.0025             | 64         | $1 \times 10^{-4}$ |
| ResNets             | 0.025              | 64         | $1 \times 10^{-4}$ |
| Deeper ConvRNNs     | 0.0025             | 64         | $1 \times 10^{-4}$ |

The only exceptions to the above are the models that are the result of the large-scale hyperparameter searches, detailed in Section \textsuperscript{A.4}. Here the learning rate and batch size are allowed to vary, and the L2 regularization is not uniform across the model, but is also allowed to vary for both the feedforward backbone and each layer’s ConvRNN cell. We list the learning rates and batch sizes for these models below:

| Model                              | Base Learning Rate       | Batch Size |
|------------------------------------|--------------------------|------------|
| Shallow LSTM (“LSTM Opt” in Figure 2b) | $7.587 \times 10^{-3}$ | 64         |
| RGC Random                         | $5.184 \times 10^{-3}$  | 64         |
| RGC Median                         | $6.736 \times 10^{-3}$  | 64         |

Since these model hyperparameters are non-standard, we manually drop the learning rate (using the same decay factor of 90\%) once the top-1 validation accuracy saturates at that given
learning rate.

A.2 Feedforward model architectures

A.2.1 BaseNet architectures

Here we provide the architectures of the feedforward CNNs we developed in this paper, referred to as “BaseNet” when they are later implanted with ConvRNN cells. For all of these architectures, we use ELU nonlinearities\(^{41}\).

The 6-layer BaseNet (into which we implanted ConvRNN cells to form the orange “Shallow ConvRNN” model class in Figure 4d), referenced as “FF” in Figure 2b, referred to as “BaseNet” among the blue “Shallow Feedforward” models in Figure 4d, and “Feedforward” in Figure S2c, had the following architecture:

| Layer | Kernel Size | Channels | Stride | Max Pooling |
|-------|-------------|----------|--------|-------------|
| 1     | 7 × 7       | 64       | 2      | 2 × 2       |
| 2     | 3 × 3       | 128      | 1      | 2 × 2       |
| 3     | 3 × 3       | 256      | 1      | 2 × 2       |
| 4     | 3 × 3       | 256      | 1      | 2 × 2       |
| 5     | 3 × 3       | 512      | 1      | 2 × 2       |
| 6     | 2 × 2       | 1000     | 1      | No          |

The 11-layer BaseNet used for the “Deeper ConvRNNs” (green models in Figure 4h) and modeled after ResNet-18\(^{40}\) (but using MaxPooling rather than stride-2 convolutions to perform downsampling) is given below:
| Block | Kernel Size | Depth | Stride | Max Pooling | Repeat |
|-------|-------------|-------|--------|-------------|--------|
| 1     | 7 × 7       | 64    | 2      | 2 × 2       | ×1     |
| 2     | 3 × 3       | 64    | 1      | None        | ×2     |
| 3     | 3 × 3       | 64    | 1      | None        | ×2     |
| 4     | 3 × 3       | 128   | 1      | 2 × 2       | ×2     |
| 5     | 3 × 3       | 128   | 1      | None        | ×2     |
| 6     | 3 × 3       | 256   | 1      | 2 × 2       | ×2     |
| 7     | 3 × 3       | 256   | 1      | 2 × 2       | ×2     |
| 8     | 3 × 3       | 512   | 1      | None        | ×2     |
| 9     | 3 × 3       | 512   | 1      | None        | ×2     |
| 10    | 3 × 3       | 512   | 1      | 2 × 2       | ×2     |
| 11    | None (Avg. Pool FC) | 1000 | None | None | ×1 |

This is the BaseNet used in Figure 3, among the red “Deeper Feedforward” models in Figure 4d, Figure S4, and Figure S5.

The variant of the above 6-layer feedforward CNN, referenced in Figure 2b as “FF Wider” is given below:

| Layer | Kernel Size | Channels | Stride | Max Pooling |
|-------|-------------|----------|--------|-------------|
| 1     | 7 × 7       | 128      | 2      | 2 × 2       |
| 2     | 3 × 3       | 512      | 1      | 2 × 2       |
| 3     | 3 × 3       | 512      | 1      | 2 × 2       |
| 4     | 3 × 3       | 512      | 1      | 2 × 2       |
| 5     | 3 × 3       | 1024     | 1      | 2 × 2       |
| 6     | 2 × 2       | 1000     | 1      | None        |

The “FF Deeper” model referenced in Figure 2b is given below:
| Layer | Kernel Size | Depth | Stride | Max Pooling |
|-------|-------------|-------|--------|-------------|
| 1     | 7 × 7       | 64    | 2      | 2 × 2       |
| 2     | 3 × 3       | 64    | 1      | None        |
| 3     | 3 × 3       | 64    | 1      | None        |
| 4     | 3 × 3       | 128   | 1      | 2 × 2       |
| 5     | 3 × 3       | 128   | 1      | None        |
| 6     | 3 × 3       | 256   | 1      | 2 × 2       |
| 7     | 3 × 3       | 256   | 1      | 2 × 2       |
| 8     | 3 × 3       | 512   | 1      | None        |
| 9     | 3 × 3       | 512   | 1      | None        |
| 10    | 3 × 3       | 512   | 1      | 2 × 2       |
| 11    | None (Avg. Pool FC) | 1000  | None   | None        |

### A.2.2 AlexNet

We use the standard AlexNet architecture, which uses local response normalization\(^\text{[26]}\). We note that we are able to attain a higher than reported top-1 validation accuracy of 63.9\% (compared to 57\% accuracy) by using the ResNet preprocessing mentioned in Section \[A.1.3\].

### A.2.3 ResNet Architectures

For the ResNet architectures, we used the original v1 versions\(^\text{[40]}\) for ResNet-18 and ResNet-34. For deeper ResNets (ResNet-50, ResNet-101, and ResNet-152), we used the v2 variants of ResNets, as this gave them a slightly higher increase in top-1 ImageNet validation accuracy. Specifically, the v2 variants of ResNets use the pre-activation of the weight layers rather than the post-activation used in the original versions. Furthermore, the v2 variants of ResNets apply batch normalization\(^\text{[42]}\) and ReLU to the input prior to the convolution, whereas the original variants apply these operations after the convolution. We use the TensorFlow Slim implementations for these two variants provided here: [https://github.com/tensorflow/models/tree/master/research/slim](https://github.com/tensorflow/models/tree/master/research/slim).
A.3 ConvRNN Cell Equations

Here we provide the explicit update equations for each of the ConvRNN cells referenced in the barplot in Figure 2d ($C_\ell$ in (1)), where $\sigma$ denotes the sigmoid function.

Throughout these sections, we let $\circ$ denote Hadamard (elementwise) product, let $\ast$ denote convolution, let $h_\ell^t$ denote the output of the cell, let $s_\ell^t$ denote the propagated memory of the cell (also known as the hidden state), and let $x_\ell^t = \bigoplus_{j \neq \ell} r_j^t$ denote the input to the cell at layer $\ell$ (this is the concatenation of feedforward and feedback inputs to layer $\ell$, defined in Section A.1.2).

In the following table, we provide the number of timesteps the ConvRNNs were unrolled for, and the timestep at which the image presentation was replaced by a mean gray stimulus during model training:

| Model Class       | Unroll Timesteps | Image Presentation Off Timestep |
|-------------------|------------------|---------------------------------|
| Shallow ConvRNNs  | 16               | 12                              |
| Deeper ConvRNNs   | 17               | 12                              |
| RGC Random        | 26               | 10                              |

These parameters were chosen based on what yielded high performance for that model class and also what was able to feasibly fit into TPU memory for training (more unroll timesteps requires more memory, but can also lead to instability during training, as is common with training RNNs). For the “Shallow ConvRNNs”, ConvRNN cells were implanted into convolutional layers 3, 4, and 5 of the 6-layer BaseNet. For the “Deeper ConvRNNs”, ConvRNN cells were implanted into convolutional layers 4, 5, 6, 7, 8, 9, and 10 of the 11-layer BaseNet.
A.3.1 Time Decay

This is the simplest form of recurrence that we consider and has a discrete-time trajectory given by

\[ s_\ell^t = F_\ell(x_\ell^t) + \tau_\ell s_\ell^{t-1} \]
\[ h_\ell^t = s_\ell^t, \]

where \( \tau_\ell \) is the learned time constant at a given layer \( \ell \). This model is intended to be a control for simplicity, where the time constants could model synaptic facilitation and depression in a cortical layer.

For the TensorFlow implementation of this cell, see the GenFuncCell() class and its associated memory() function in the cell.py file of our TNN Github repository.

A.3.2 SimpleRNN

The update equations in this case are given by:

\[ a_\ell^t = W_s^\ell * s_\ell^{t-1} + b_s^\ell \]
\[ i_\ell^t = W_i^\ell * x_\ell^t + b_i^\ell \]
\[ s_\ell^t = \text{elu}(\text{LN}(i_\ell^t + a_\ell^t)) \]
\[ h_\ell^t = s_\ell^t, \]

where LN denotes the layer normalization operation with offset parameter \( \beta \) initialized to 0 and scale parameter \( \gamma \) initialized to 1. For the shallow SimpleRNN (among the orange “Shallow ConvRNN” models in Figure), we use layer normalization but omit its usage in the deeper ConvRNN as it was not able to train with that operation.

For the TensorFlow implementation of this cell, see the ConvNormBasicCell() class in the convrnn.py file of our TNN Github repository.
A.3.3 GRU

We adapt the standard GRU cell\(^{15}\) to the convolutional setting:

\[
\begin{align*}
\ell_t^r &= \sigma(W^r_{\ell} \ast x^\ell_t + U^r_{\ell} \ast s^\ell_{t-1} + b^r_{\ell} + 1) \\
u^\ell_t &= \sigma(W^u_{\ell} \ast x^\ell_t + U^u_{\ell} \ast s^\ell_{t-1} + b^u_{\ell}) \\
c^\ell_t &= \tanh(W^c_{\ell} \ast x^\ell_t + U^c_{\ell} \ast (r^\ell_t \circ s^\ell_{t-1}) + b^c_{\ell}) \\
s^\ell_t &= u^\ell_t \circ s^\ell_{t-1} + (1 - u^\ell_t) \circ c^\ell_t \\
h^\ell_t &= s^\ell_t.
\end{align*}
\]

(4)

For the TensorFlow implementation of this cell, see the ConvGRUCell() class in the convrnn.py file of our TNN Github repository.

A.3.4 LSTM

We adapt the standard LSTM cell\(^{15}\) to the convolutional setting, with some slight modifications such as added layer normalization for stability in training.

We first make the gates convolutional as follows:

\[
\begin{align*}
i^\ell_t &= LN(W^i_{\ell} \ast x^\ell_t + U^i_{\ell} \ast h^\ell_{t-1} + b^i_{\ell}) \\
j^\ell_t &= LN(W^j_{\ell} \ast x^\ell_t + U^j_{\ell} \ast h^\ell_{t-1} + b^j_{\ell}) \\
f^\ell_t &= LN(W^f_{\ell} \ast x^\ell_t + U^f_{\ell} \ast h^\ell_{t-1} + b^f_{\ell}) \\
o^\ell_t &= LN(W^o_{\ell} \ast x^\ell_t + U^o_{\ell} \ast h^\ell_{t-1} + b^o_{\ell}),
\end{align*}
\]

(5)

where LN denotes the layer normalization operation\(^{44}\) with offset parameter \(\beta\) initialized to 0 and scale parameter \(\gamma\) initialized to 1.

Next, the LSTM update equations are as follows:

\[
\begin{align*}
s^\ell_t &= LN(s^\ell_{t-1} \circ \sigma(f^\ell_t + f^f_{b}) + \sigma(i^\ell_t) \circ \tanh(j^\ell_t)) \\
h^\ell_t &= \tanh(s^\ell_t) \circ \sigma(o^\ell_t),
\end{align*}
\]

(6)
where $f^\ell_b$ is the forget gate bias, typically set to 1, as recommended by others. When peephole connections are allowed, these update equations are augmented to become:

$$s^\ell_t = LN(s^\ell_{t-1} \circ \sigma(f^\ell_t + f^\ell_b + V^\ell_f \circ s^\ell_{t-1}) + \sigma(i^\ell_t + V^\ell_i \circ s^\ell_{t-1}) \circ \tanh(j^\ell_t))$$

$$h^\ell_t = \tanh(s^\ell_t) \circ \sigma(o^\ell_t + V^\ell_o \circ s^\ell_{t-1}).$$

(7)

In the shallow LSTM (among the orange “Shallow ConvRNN” models in Figure 4d), we use peepholes and layer normalization, as that was found in the LSTM search for shallow models (described in Section A.4.1) to be useful for performance. We found, however, that neither of these augmentations are needed in the deeper variant (among the green “Deeper ConvRNN” models in Figure 4d) in order to achieve high top-1 validation accuracy on ImageNet.

For the TensorFlow implementation of this cell, see the ConvLSTMCell() class in the convrnn.py file of our TNN Github repository.

A.3.5 UGRNN

We adapt the UGRNN to the convolutional setting. The update equations are as follows:

$$c^\ell_t = \tanh(W^\ell_c \ast x^\ell_t + U^\ell_c \ast s^\ell_{t-1} + b^\ell_c)$$

$$g^\ell_t = \sigma(W^\ell_g \ast x^\ell_t + U^\ell_g \ast s^\ell_{t-1} + b^\ell_g + 1)$$

$$s^\ell_t = g^\ell_t \circ s^\ell_{t-1} + (1 - g^\ell_t) \circ c^\ell_t$$

$$h^\ell_t = s^\ell_t.$$  

(8)

For the TensorFlow implementation of this cell, see the ConvUGRNNCell() class in the convrnn.py file of our TNN Github repository.
A.3.6 IntersectionRNN

We adapt the IntersectionRNN to the convolutional setting. The update equations are as follows:

\[
\begin{align*}
m^\ell_t &= \tanh(W^m_\ell \ast x^\ell_t + U^m_\ell \ast s^\ell_{t-1} + b^m_\ell) \\
n^\ell_t &= \text{relu}(W^n_\ell \ast x^\ell_t + U^n_\ell \ast s^\ell_{t-1} + b^n_\ell) \\
p^\ell_t &= \sigma(W^p_\ell \ast x^\ell_t + U^p_\ell \ast s^\ell_{t-1} + b^p_\ell + 1) \\
y^\ell_t &= \sigma(W^y_\ell \ast x^\ell_t + U^y_\ell \ast s^\ell_{t-1} + b^y_\ell + 1) \\
s^\ell_t &= p^\ell_t \circ s^\ell_{t-1} + (1 - p^\ell_t) \circ m^\ell_t \\
h^\ell_t &= y^\ell_t \circ x^\ell_t + (1 - y^\ell_t) \circ n^\ell_t.
\end{align*}
\]

(9)

For the TensorFlow implementation of this cell, see the ConvIntersectionRNNCell() class in the convrnn.py file of our TNN Github repository.

A.3.7 Reciprocal Gated Cell (RGC)

Here we provide the explicit update equations for the Reciprocal Gated Cell, diagrammed in Figure 2a (bottom right). The update equation for the output of the cell, \( h^\ell_t \), is given by a gating of both the input and memory \( s^\ell_t \):

\[
a^\ell_{t+1} = (1 - \sigma(W^r_{sh} \ast s^\ell_t)) \circ x^\ell_t + (1 - \sigma(W^r_{hh} \ast h^\ell_t)) \circ h^\ell_t
\]

\[
h^\ell_t = \text{elu} \left( a^\ell_t \right).
\]

(10)

The update equation for the memory \( s^\ell_t \) is given by a gating of the input and the output of the cell \( h^\ell_t \):

\[
\tilde{s}^\ell_{t+1} = (1 - \sigma(W^r_{hs} \ast h^\ell_t)) \circ x^\ell_t + (1 - \sigma(W^r_{ss} \ast s^\ell_t)) \circ s^\ell_t
\]

\[
s^\ell_t = \text{elu} \left( \tilde{s}^\ell_t \right).
\]

(11)

For the TensorFlow implementation of this cell, see the ReciprocalGateCell() class in the reciprocalgaternn.py file of our TNN Github repository.
We employed a form of Bayesian optimization, a Tree-structured Parzen Estimator (TPE), to search the space of continuous and categorical hyperparameters\cite{10.1145/3065388.3065455}. This algorithm constructs a generative model of $P[\text{score} \mid \text{configuration}]$ by updating a prior from a maintained history $H$ of hyperparameter configuration-loss pairs. The fitness function that is optimized over models is the expected improvement, where a given configuration $c$ is meant to optimize $EI(c) = \int_{x \leq t} P[x \mid c, H]$. This choice of Bayesian optimization algorithm models $P[c \mid x]$ via a Gaussian mixture, and restricts us to tree-structured configuration spaces.

Models were trained synchronously 100 models at a time using the HyperOpt package\cite{2015arXiv150204999P}, which implements the above Bayesian optimization. Each model was trained on its own Tensor Processing Unit (TPUv2), and during the search, ConvRNN models were trained by stochastic gradient descent on 128 pixel ImageNet for efficiency. The top performing ConvRNN models were then fully trained out on 224 pixel ImageNet.

### A.4.1 LSTM search

The search for better LSTM architectures involved searching over training hyperparameters and common structural variants of the LSTM to better adapt this local structure to deep convolutional networks, using hundreds of second generation Google Tensor Processing Units (TPUv2s). We searched over learning hyperparameters (e.g. gradient clip values, learning rate) as well as structural hyperparameters (e.g. gate convolution filter sizes, channel depth, whether or not to use peephole connections, etc.).

Specifically, we implanted LSTMs into convolutional layers 3, 4, and 5, of the 6-layer BaseNet described in Section A.2. At each of these layers, the parameters of the LSTM cell (defined in Section A.3.4) were allowed to vary per layer, as follows:

- The discrete number of convolutional channels was chosen from $\{64, 128, 256\}$. 
• The discrete choice of convolutional filter sizes were chosen from \{1, 4\}.

• The binary choice of whether or not to use layer normalization.

• The strength of the L2 regularization of all LSTM parameters in that layer $\in [10^{-7}, 10^{-3}]$, sampled log-uniformly.

• The scale of the He-style initialization\(^\text{51}\) of the convolutional filter weights $\in [0.25, 2]$, sampled uniformly.

• The value of the constant initialization of the biases $\in [-2, 2]$, sampled uniformly.

• The forget gate bias $f^\ell_b \in [0, 6]$, sampled uniformly (defined in (6)).

• The binary choice of whether or not to use peephole connections (as defined in (7)).

Outside of the LSTM cell at each layer, we additionally searched over the following parameters as well:

• The number of discrete timesteps the model is unrolled $\in [12, 26]$, sampled uniformly in consecutive groups of size 2.

• The timestep at each the image presentation is “turned off” and replaced with a mean gray stimulus $\in [8, 12]$, sampled uniformly in consecutive groups of 2.

• The discrete choice of batch size used for the training the entire model $\in \{64, 128, 256\}$.

• The learning rate for training the entire model $\in [10^{-3}, 10^{-1}]$, sampled log-uniformly.

• The binary choice of whether or not to use Nesterov momentum\(^\text{52}\).

• The gradient clipping value $\in [0.3, 3]$, sampled log-uniformly.
• The scale of the He-style initialization of the convolutional filter weights of the feed-forward base model $\in [0.25, 2]$, sampled uniformly.

• The strength of the L2 regularization of the feedforward base model parameters $\in [10^{-7}, 10^{-3}]$, sampled log-uniformly.

Each search point is a sampled value from the above described search space and trained for 1 epoch on ImageNet, in order to sample as many models as much as possible with the computational resources available. More than 1600 models were sampled in total, and we trained out the top ones and the median performing one after 1 epoch were trained out fully on 224 pixel ImageNet. The median model from this search attained the best top-1 validation accuracy on ImageNet, which is the resultant “LSTM Opt” model in Figure 2b.

A.4.2 Reciprocal Gate Cell (RGC) search

From the Reciprocal Gated Cell equations in (10) and (11), there are a variety of possibilities for how $h_{t-1}^{\ell}, x_t^{\ell}, s_t^{\ell},$ and $h_t^{\ell}$ can be connected to one another (schematized in Figure 2c).

Mathematically, the search in Figure 2c can be formalized in terms of the following update equations. First, we define our input sets and building block functions:

$$\text{minin} = \{h_{t-1}^{\ell}, x_t^{\ell}, s_t^{\ell}, h_t^{\ell} \}$$

$$\text{minin}_a = \text{minin} \cup \{s_t^{\ell} \}$$

$$\text{minin}_b = \text{minin} \cup \{h_t^{\ell} \}$$

$$S_a \subseteq \text{minin}_a$$

$$S_b \subseteq \text{minin}_b$$

$$\text{Affine}(x) \in \{ +, 1 \times 1 \text{ conv}, K \times K \text{ conv}, K \times K \text{ depth-separable conv} \}$$

$$K \in \{3, \ldots, 7 \}$$
With those in hand, we have the following update equations:

\[
\tau_a = v_1^\tau + v_2^\tau \sigma(\text{Affine}(S_a))
\]

\[
\tau_b = v_1^\tau + v_2^\tau \sigma(\text{Affine}(S_b))
\]

\[
\text{gate}_a = v_1^q + v_2^q \sigma(\text{Affine}(S_a))
\]

\[
\text{gate}_b = v_1^q + v_2^q \sigma(\text{Affine}(S_b))
\]

\[
a_t^\ell = \{\text{gate}_a\} \cdot \text{in}_t^\ell + \{\tau_a\} \cdot h_{t-1}^\ell
\]

\[
h_t^\ell = f(a_t^\ell)
\]

\[
b_t^\ell = \{\text{gate}_b\} \cdot \text{in}_t^\ell + \{\tau_b\} \cdot s_{t-1}^\ell
\]

\[
s_t^\ell = f(b_t^\ell)
\]

\[
f \in \{\text{elu}, \tanh, \sigma\}.
\]

For clarity, the following matrix summarizes the connectivity possibilities (with ? denoting the possibility of a connection), schematized in Figure 2c:

\[
\begin{pmatrix}
    h_{t-1}^\ell & x_t^\ell & s_{t-1}^\ell & s_t^\ell & h_{t-1}^\ell & h_t^\ell \\
    0 & 1 & 0 & ? & 0 & ? \\
    x_t^\ell & 0 & 0 & 0 & ? & 0 & ? \\
    s_{t-1}^\ell & 0 & 0 & 0 & ? & 0 & ? \\
    s_t^\ell & 0 & 0 & 0 & 0 & 0 & ? \\
    h_{t-1}^\ell & 0 & 0 & 0 & ? & 0 & ? \\
    h_t^\ell & 0 & 0 & 0 & ? & 0 & 0
\end{pmatrix}
\]

Each search point is a sampled value from the above described search space and trained for 5 epochs on ImageNet, in order to sample as many models as much as possible with the computational resources available. Around 6000 models were sampled in total over the course of the search. The top and median models from this search were then fully trained out on 224 pixel ImageNet with a batch size of 64 (which was maximum that we could fit into TPU memory). Moreover, as explicated in the table in Section A.1.3, the ResNet models were also trained using this same batch size, with the standard ResNet learning rate of 0.1 for a batch size of 256 linearly rescaled to accommodate, to ensure fair comparison between these two model
classes. The median model from this search attained the best top-1 validation accuracy on ImageNet of all models selected to be trained out fully on ImageNet from the search, producing the resultant “RGC Median” model in Figure 2d (note that this designation also includes the long-range feedback connections). The “RGC Random” model is from the random phase of this search (400th sampled model, since models sampled earlier than that failed to train out fully on ImageNet).

A.5 Decoders

In addition to choice of ConvRNN cell, we consider particular choices of “light-weight” (in terms of parameter count) decoding strategy that determines the final object category of that image. By construction, the model will output category logit probabilities at each timestep, given by the softmax function

$$\text{softmax}(z; \beta) = \frac{e^{\beta z_i}}{\sum_{j=1}^{C} e^{\beta z_j}}$$

where $C = 1000$ is the number of ImageNet categories. This will then be passed to a decoding function which can take one of several forms:

1. **Default**: Use the logits at the last timestep and discard the remaining, with $\beta = 1$.

2. **Threshold Decoder**: Select the logits from the first timepoint at which the maximum logit value at that timepoint crosses a fixed threshold (set to 0.9), with $\beta = 1$.

3. **Max Confidence Decoder**: For the most confident category, find the timepoint at which that confidence peaks, and return the logits at that timepoint, where $\beta$ is a trainable scalar parameter initialized to 1.

“RGC Median” therefore refers to the model trained using the default decoder, but when using the other two decoders with the “RGC Median” model, we append it to the name (as is done in Figures 4d, S4a, and S5).
A.6 Model prediction of neural responses

A.6.1 Neural data

Neural responses came from three multi-unit arrays per primate (rhesus macques): one implanted in V4, one in posterior IT (pIT), and one in central and anterior IT (cIT/aIT)\(^\text{[29]}\). Each image was presented approximately 50 times, using rapid visual stimulus presentation (RSVP). Each stimulus was presented for 100\(\text{ms}\), followed by a mean gray stimulus interleaved between images. Each trial lasted 260\(\text{ms}\). The image set consisted of 5120 images based on 64 object categories. These objects belonged to 8 high-level categories (tables, planes, fruits, faces, chairs, cars, boats, animals), each of which consisted of 8 unique objects. Each image consisted of a 2D projection of a 3D model added to a random background. The pose, size, and \(x\)- and \(y\)-position of the object was varied across the image set, whereby 2 levels of variation were used (corresponding to medium and high variation\(^\text{[29]}\)). Multi-unit responses to these images were binned in 10\(\text{ms}\) windows, averaged across trials of the same image, and normalized to the average response to a blank image. This produced a set of 5120 images \(\times 256\) units \(\times 25\) timebins responses, which were the targets for our model features to predict. There were 88 units from V4, 88 units from pIT, and 80 units from cIT/aIT.

A.6.2 Fitting procedure

Generating train/test split. The 5120 images were split 75\%-25\% within each object category into a training set and a held-out testing set. All images were presented to the models for 10 time steps (corresponding to 100\(\text{ms}\)), followed by a mean gray stimulus for the remaining 15 time steps, to match the image presentation to the primates. The images are matched to the procedure when used to validate the models on ImageNet, namely they are bilinearly resized to 224 \(\times\) 224 and normalized by the ImageNet mean ([0.485, 0.456, 0.406]) and standard deviation ([0.229, 0.224, 0.225]), applied per channel.
Model layer determination. We stipulated that units from each multi-unit array must be fit by features from a single model layer. To determine which one, we fit the features from the relevant feedforward BaseNet (either the 6-layer BaseNet or 11-layer BaseNet) to unit’s time-averaged response, and counted how many units had minimal loss for a given model layer, schematized in Step 2 of Figure 1. This yielded a mapping from the V4 array to model layer 3 of the 6-layer BaseNet and model layers 5 & 6 of the 11-layer BaseNet, pIT mapping to model layer 4 of the 6-layer BaseNet and model layers 7 & 8 of the 11-layer BaseNet, and cIT/aIT mapping to layer 5 of the 6-layer BaseNet and model layers 9 & 10 of the 11-layer BaseNet.

Mapping transform from models to neural responses. Model features from each image (i.e. the activations of units in a given model layer) were linearly fit to the neural responses by stochastic gradient descent with a standard L2 loss using a spatially factored mapping, where each of the 256 units was fit independently. This spatially factored mapping is defined as follows: Given a model feature $f^\ell \in \mathbb{R}^{x,y,c}$ from layer $\ell$, where $x$ and $y$ are the number of units in the spatial extent and $c$ is the number of channels, we fit a spatial mask $w_{\text{space}} \in \mathbb{R}^{x,y}$ and a channel mask $w_{\text{channels}} \in \mathbb{R}^c$ for each neuron $n$ to predict the ground-truth neuron’s response $r_{i,n,t}$ at image $i$ and timebin $t$. The predicted response can be written as:

$$\hat{r}_{i,n,t;w} = \sum_{i=1}^{x} \sum_{j=1}^{y} \sum_{k=1}^{c} w_{\text{space}}[i, j] w_{\text{channels}}[k] f^\ell[i, j, k]. \quad (12)$$

This mapping is implemented in the `factored_fc()` function of the `cell.py` file of our TNN Github repository.

Loss function. After these layers were determined, model features were then fit to the entire set of 25 timebins for each unit using a shared linear model: that is, a single set of regression coefficients was used for all timebins, as schematized in Step 3 of Figure 1. The loss for this fitting was the average L2 loss across training images and 25 timebins for each unit, given by

$$\mathcal{L}(\hat{r}_{i,n,t;w}, r_{i,n,t}) = \frac{1}{|B|} \sum_{t=6}^{25} \sum_{i \in B} \sum_{n=1}^{256} (\hat{r}_{i,n,t;w} - r_{i,n,t})^2. \quad (13)$$
Note that $t$ indexes model timesteps, which correspond to 10ms timebins, so $t = 6$ refers to the 60-70ms timebin, $t = 7$ refers to the 70-80ms timebin, and so forth.

We trained the temporally-fixed parameters $w = [w_{\text{space}}; w_{\text{channels}}]$ of the mapping using the Adam optimizer\(^{53}\) with a learning rate of $1 \times 10^{-4}$ and a training batch size $|\mathcal{B}| = 64$ images. Additionally, we used a dropout\(^{53}\) level of 0.5 on the model features, prior to the mapping, as further regularization.

### A.6.3 Metrics

To estimate a noise ceiling for each neuron’s response at each timebin, we computed the Spearman-Brown corrected split-half reliability $\rho_n$ of neuron $n$, averaged across 900 bootstrap iterations of split-half trials.

Let “Neural Predictivity” (used in Figure S2) refer to

$$\text{Corr}(\hat{r}_{\text{test}}^n, r_{\text{test}}^n),$$

namely the Pearson correlation across test set images of the model’s response $\hat{r}_{\text{test}}^n$ to the of any neuron $n$’s response $r_{\text{test}}^n$ at a given timebin (or time-averaged).

The “Neural Predictivity (Noise Corrected)” (used in Figure 3 and Figure S4) for neuron $n$ is given by

$$\frac{\text{Corr}(\hat{r}_{\text{test}}^n, r_{\text{test}}^n)}{\sqrt{\rho_n}}.$$  

### A.7 Inter-animal consistency

We provide the definition and justification of the inter-animal consistency metric mentioned in Figure S4b. Suppose we have neural responses from two primates $A$ and $B$. Let $t_i^p$ be the vector of true responses (either at a given timebin or averaged across a set of timebins) of primate $p \in \{A, B\}$ on stimulus set $i \in \{\text{train, test}\}$. Of course, we only receive noisy observations of $t_i^p$, so let $s_{j,i}^p$ be the $j$-th set of $n$ trials of $t_i^p$. Finally, let $M(x)_i$ be the predictions of a mapping.
When trained on input $x$ and tested on stimulus set $i$. For example, $M(t_{train}^p)_{test}$ is the prediction of the mapping $M$ on the test stimulus trained on the true neural responses from primate $p$ on the train stimulus, and correspondingly, $M(s_{1,train}^p)_{test}$ is the prediction of the mapping $M$ on the test stimulus trained on the (trial-average) of noisy sample 1 on the train stimulus from primate $p$.

With these definitions in hand, the inter-animal mapping consistency from one primate $A$ to another primate $B$ corresponds to the following true quantity to be estimated:

$$\text{Corr}(M(t_{train}^A)_{test}, t_{test}^B),$$  \quad (16)

where Corr is the Pearson correlation across test stimuli. In what follows, we argue that this true quantity can be approximated with the following ratio of measurable quantities where we divide the noisy trial observations into two sets of equal samples:

$$\text{Corr}(M(t_{train}^A)_{test}, t_{test}^B) \sim \frac{\text{Corr}(M(s_{1,train}^A)_{test}, s_{2,test}^B)}{\sqrt{\text{Corr}(M(s_{1,train}^A)_{test}, s_{2,train}^A) \times \text{Corr}(s_{1,test}^B, s_{2,test}^B)}}.$$

(17)

In words, the inter-animal consistency corresponds to the predictivity of the mapping on the test set stimuli from primate $A$ to $B$ on two different (averaged) halves of noisy trials, corrected by the square root of the mapping reliability on primate $A$’s test stimuli responses on two different halves of noisy trials and the internal consistency of primate $B$.

We justify the approximation in (17) by gradually eliminating the true quantities by their measurable estimates, starting from the original quantity in (16). First, we make the approximation that

$$\text{Corr}(M(t_{train}^A)_{test}, s_{2,test}^B) \sim \text{Corr}(M(t_{train}^A)_{test}, t_{test}^B) \times \text{Corr}(t_{test}^B, s_{2,test}^B).$$

(18)

by transitivity of positive correlations (which is reasonable assumption when the number of stimuli is large). Next, by normality assumptions in the structure of the noisy estimates and
since the number of trials \((n)\) between the two sets is the same, we have that

\[
\text{Corr} \left( s_{1, \text{test}}^B, s_{2, \text{test}}^B \right) \sim \text{Corr} \left( t_2^B, s_{2, \text{test}}^B \right)^2. \tag{19}
\]

Namely, the correlation between the average of two sets of noisy observations of \(n\) trials each is approximately the square of the correlation between the true value and average of one set of \(n\) noisy trials. Therefore, from (18) and (19) it follows that

\[
\text{Corr} \left( M \left( t_{\text{train}}^A \right)_{\text{test}}^B, t_2^B \right) \sim \frac{\text{Corr} \left( M \left( t_{\text{train}}^A \right)_{\text{test}}^B, s_{2, \text{test}}^B \right)}{\sqrt{\text{Corr} \left( s_{1, \text{test}}^B, s_{2, \text{test}}^B \right)}}. \tag{20}
\]

We have gotten rid of \(t_2^B\), but we still need to get rid of the \(M \left( t_{\text{train}}^A \right)_{\text{test}}^B\) term. We apply the same two steps by analogy though these approximations may not always be true (though are true for additive Gaussian noise):

\[
\text{Corr} \left( M \left( s_{1, \text{train}}^A \right)_{\text{test}}^B, s_{2, \text{test}}^B \right) \sim \text{Corr} \left( s_{2, \text{test}}^B, M \left( t_{\text{train}}^A \right)_{\text{test}}^B \right) \times \text{Corr} \left( M \left( t_{\text{train}}^A \right)_{\text{test}}^B, M \left( s_{1, \text{train}}^A \right)_{\text{test}}^B \right)
\]

\[
\text{Corr} \left( M \left( s_{1, \text{train}}^A \right)_{\text{test}}^B, M \left( s_{2, \text{train}}^A \right)_{\text{test}}^B \right) \sim \text{Corr} \left( M \left( s_{1, \text{train}}^A \right)_{\text{test}}^B, M \left( t_{\text{train}}^A \right)_{\text{test}}^B \right)^2,
\]

which taken together implies

\[
\text{Corr} \left( M \left( t_{\text{train}}^A \right)_{\text{test}}^B, s_{2, \text{test}}^B \right) \sim \frac{\text{Corr} \left( M \left( s_{1, \text{train}}^A \right)_{\text{test}}^B, s_{2, \text{test}}^B \right)}{\sqrt{\text{Corr} \left( M \left( s_{1, \text{train}}^A \right)_{\text{test}}^B, M \left( s_{2, \text{train}}^A \right)_{\text{test}}^B \right)}}. \tag{21}
\]

Equations (20) and (21) together imply the final estimated quantity given in (17).

**A.8 Object solution times (OSTs)**

**A.8.1 Generating model OSTs**

Here we describe how we defined object solution times from both feedforward models and ConvRNNs. As depicted in Figure 1, this is a multi-stage process that involves first identifying the most “IT-preferred” layers of each model.
Determining “IT-preferred” model layers. These are identified by a standard linear mapping using 25 component partial least squares regression (PLS), from model layer units to time-averaged IT (namely, pIT/cIT/aIT) responses from the neural data described in Section A.6.1, and corroborates the results obtained by the same procedure described in Section A.6.2. We use this neural data as it has both V4 and IT responses, and demonstrates a disjoint set of layers between the preferred V4 model layers and preferred IT layers.

Mapping model timepoints to IT timepoints. Once these “IT-preferred” model layers are identified, we then map these model timepoints to 10 ms timebins as in the IT data. For ConvRNNs with intrinsic temporal dynamics, this mapping is one-to-one, we simply concatenate the model layers at each timepoint to construct an entire IT pseudopopulation, and each timepoint of the ConvRNN corresponds to a 10 ms timebin between 70-260 ms. For feedforward models, we map each “IT-preferred” layer to a 10 ms timebin between 70-260 ms. If the number of “IT-preferred” layers for a feedforward model matches the total number of timebins (19), then there is only one admissible mapping, corresponding to the “uniform” mapping, whereby the earliest (in the feedforward hierarchy) layer is matched to the earliest 10 ms timebin of 70 ms, and so forth. On the other hand, if the number of “IT-preferred” layers is strictly less than the total number of timebins, then we additionally consider a “graded” mapping that picks a random sample of units from one layer to the next so that the number of feedforward layers exactly matches the total number of timebins.

Obtaining model $d'$ values. Once a timepoint mapping is selected, we compute the model object solution time (OST) in the same manner as the OST is computed for IT. Specifically, we train an SVM ($C = 5 \times 10^4$) separately for each model timepoint after it has been dimension reduced through PCA (with 1000 components) to solve the ten-way categorization task for each image. The ten categories are apple, bear, bird, car, chair, dog, elephant, person, plane, and zebra. 1000 images constitute the training set of the SVM (100 images per category) and 320
images are randomly chosen to be in the test set. We perform 20 trials each of 10 train/test splits to get errorbars, where each image is in the test set at least once. The (model or IT) $d'$ for that image is only computed from the SVM when it has been in the test set, and is bounded between -5 and 5. Since this dataset consists of 1,320 grayscale images presented centrally to behaving primates for 100 ms, there are therefore 1,320 $d'$ values (one for each image) for any given model, constituting its “I1” vector.

**Correlating model OST with IT OST.** The OST of the model therefore is the first model timepoint in which the $d'$ reaches the recorded primate $d'$ for that image. Using the Levenberg–Marquardt algorithm, we further linearly interpolate between 10 ms bins to determine the precise millisecond that the response surpassed the primate’s behavioral output for that image (as was done analogously with the IT population’s OST). Finally, we compare the model OST to the IT OST via a Spearman correlation across the common set of images solved by both the model and IT.

### A.8.2 Relating the linear mapping to neural responses with the OST behavioral metric

The IT population OST was computed from primarily anterior IT (aIT) responses. Therefore, to isolate the interaction a linear mapping of model features to neural responses (as we do in neural response prediction described in Section A.6) might have compared to directly computing the OST from the original model features, we turned to neural data collected from 486 aIT units on 1100 greyscale images.

For each model, we train a linear mapping on this dataset, with 550 images used for training the mapping and 550 images are held-out for the test set. We observe similar conclusions as with the original neural data in Section A.6 for both the temporally-fixed linear mapping in Figure S3 (in the “aIT” panel), and with a temporally-varying PLS mapping in Figure S4 (“aIT” in panel (a) as well as the data used in panel (b)), all from layer 10 of the 11-layer BaseNet/ConvRNNs.
With these observations, we then proceeded to evaluate the effect of the linear mapping on OST correlations in Figure S5. Crucially, in this setting, we train a 100 component PLS mapping on the 526 images for which an IT $d'$ is not defined, in order to ensure that the images from Section A.8.1 that the OST correlation is evaluated on are not the same images the PLS mapping was trained with.
Table 1: Wilcoxon test (with Bonferroni correction) p-values for comparing each Deeper ConvRNN’s neural predictivity at the “early” timepoints (Figure 3) to the (11-layer) BaseNet.

| Model               | Visual Area | Wilcoxon test p-value |
|---------------------|-------------|-----------------------|
| Time Decay          | V4          | < 0.001               |
| IntersectionRNN     | V4          | < 0.001               |
| LSTM                | V4          | < 0.001               |
| UGRNN               | V4          | < 0.001               |
| GRU                 | V4          | < 0.001               |
| SimpleRNN           | V4          | < 0.001               |
| RGC Random          | V4          | < 0.001               |
| RGC Median          | V4          | < 0.01                |
| Time Decay          | pIT         | 0.022                 |
| IntersectionRNN     | pIT         | < 0.001               |
| LSTM                | pIT         | < 0.001               |
| UGRNN               | pIT         | < 0.001               |
| GRU                 | pIT         | < 0.001               |
| SimpleRNN           | pIT         | < 0.001               |
| RGC Random          | pIT         | 0.31                  |
| RGC Median          | pIT         | < 0.001               |
| Time Decay          | aIT         | < 0.001               |
| IntersectionRNN     | aIT         | < 0.001               |
| LSTM                | aIT         | 0.47                  |
| UGRNN               | aIT         | 0.09                  |
| GRU                 | aIT         | 0.16                  |
| SimpleRNN           | aIT         | < 0.001               |
| RGC Random          | aIT         | < 0.001               |
| RGC Median          | aIT         | < 0.01                |

Supplementary Figures
Figure S1: **Optimal local recurrent cell motif and global feedback connectivity.** (a) RNN Cell structure from the top-performing search model. Red lines indicate that this hyperparameter choice (connection and filter size) was chosen in each of the top unique models from the search. \( K \times K \) denotes a convolution and \( dSK \times K \) denotes a depth-separable convolution with filter size \( K \times K \). (b) Long-range feedback connections from the search. (Top) Each trace shows the proportion of models in a 100-sample window that have a particular feedback connection. (Bottom) Each bar indicates the difference between the median performance of models with a given feedback and the median performance of models without that feedback. Colors correspond to the same feedbacks as above.
Figure S2:  (a) **Both local recurrence and global feedbacks are needed to best fit neural data.** Among a wide range of architectures with different local recurrent motifs and global feedback patterns, the best architecture was one with both gated local recurrence and a global feedback. Local recurrent circuits were particularly useful for improving fits to IT neurons ($N = 168$), whereas both local recurrence and global feedback were critical for improving fits to V4 neurons ($N = 88$). Except for “temporally-varying mapping”, fixed model-unit-to-neuron linear mappings were fixed across all time bins, constraining trajectories to be produced by actual dynamics of the network. In contrast, “temporally-varying mapping” indicates an independent PLS regression for each time bin. The fact that models with local recurrence and global feedbacks are better than “temporally-varying mapping” suggests that some nonlinear dynamics at earlier layers contributed meaningfully to network fits. S.e.m. across four splits of held-out test images. (b) **Held-out neural predictivity.** As both 100ms and 200ms, this direct fitting procedure to the dynamics generalizes to neurons held-out (right bars) in the fitting procedure, a stronger test of generalization than held-out images depicted in the left bars. (c) **Underfitting to the task.** However, a subtle overfitting to the neural image distribution occurs, whereby the task-optimized network whose dynamics are trained on the V4 and IT neural dynamics no longer transfers to ImageNet.
Figure S3: **Modeling primate ventral stream neural dynamics with ConvRNNs.** Fitting model features of ConvRNNs with a temporally-fixed linear mapping to neural dynamics approaches the noise ceiling of these responses in most cases. The $y$-axis indicates the median across neurons of the explained variance between predictions and ground-truth responses on held-out images. Error bars indicate the s.e.m across neurons ($N = 88$ for V4, $N = 88$ for pIT, $N = 80$ for cIT/aIT, and $N = 486$ for aIT). Note that “aIT” refers to a separate neural dataset from primarily anterior IT neurons, detailed in Section A.8.2. As can be seen, the feedforward BaseNet model (purple) is incapable of generating a response beyond the feedforward pass, and certain types of ConvRNN cells added to the feedforward model are less predictive than others.
Figure S4: (a) Increasing feedforward depth does not account for drop in median explained variance from early to late timepoints. We observe a similar drop in median explained variance from 130-140\(\text{ms}\) to 200-210\(\text{ms}\), between the ConvRNN and deeper feedforward models, where we fix each model’s training image size and batch size to be able compare across depths. To compare these two models, we subselect for high reliability neurons (above 0.3 split-half consistency) and use a temporally-varying mapping (PLS 25 components). We plot the median and s.e.m. predictivity in both panels per timebin (\(N = 108, 113, 117, 123, 118, 118, 116, 115, 108, 99, 86\) neurons for each timebin in the “pIT/cIT/aIT” panel, and \(N = 247, 313, 378, 441, 437, 411, 397, 391, 392, 384, 380\) neurons for each timebin in the “aIT” panel). (b) Drop in explained variance may be exhibited in inter-animal consistency. Using the neural data described in Section \[A.8.2\], we see a similar inter-animal consistency (metric detailed in Section \[A.7\]) at 130-140\(\text{ms}\) and 200-210\(\text{ms}\), as we do with the 11-layer BaseNet. Median and s.e.m. across aIT neurons (\(N = 441\) at 130-140\(\text{ms}\) and \(N = 380\) at 200-210\(\text{ms}\)) from the dataset described in Section \[A.8.2\].
Figure S5: **Behaviorally harmful effect of dimensionality reduction due to linear transform.** Mean and s.e.m. are computed across train/test splits ($N = 10$) when that image (of 1320 images) was a test-set image, with the Spearman correlation computed with the IT solution times across the imageset mutually solved by the given model and IT. As can be seen, a temporally-graded mapping directly from the model features of feedforward models always attains higher OST consistency than a uniform one (“Graded” vs. “Uniform” comparison). We additionally train a 100 component PLS regression to IT responses at each defined model timepoint, where the responses are to a different set of images than used to evaluate the OST metric. This procedure, detailed in Section A.8.2, results in an image-computable model on which the OST metric is evaluated on and corresponds to “PLS” prepended to the name of each point on this plot, for any given model and associated temporal mapping. As can be seen, “PLS Uniform” for the BaseNet and ResNet-34 match the OST consistency of the RGC Median ConvRNNs from their original model features. However, “PLS Uniform” for the ConvRNNs and ResNet-101 and ResNet-152 have a significant decrease in OST consistency compared to when evaluated on their original model features, indicating the behaviorally harmful effect of dimensionality reduction due to PLS.