Check for updates

A neural network trained for prediction mimics diverse features of biological neurons and perception

William Lotter ^{1,6}^K, Gabriel Kreiman ^{1,2,3} and David Cox^{1,4,5}

Recent work has shown that convolutional neural networks (CNNs) trained on image recognition tasks can serve as valuable models for predicting neural responses in primate visual cortex. However, these models typically require biologically infeasible levels of labelled training data, so this similarity must at least arise via different paths. In addition, most popular CNNs are solely feedforward, lacking a notion of time and recurrence, whereas neurons in visual cortex produce complex time-varying responses, even to static inputs. Towards addressing these inconsistencies with biology, here we study the emergent properties of a recurrent generative network that is trained to predict future video frames in a self-supervised manner. Remarkably, the resulting model is able to capture a wide variety of seemingly disparate phenomena observed in visual cortex, ranging from single-unit response dynamics to complex perceptual motion illusions, even when subjected to highly impoverished stimuli. These results suggest potentially deep connections between recurrent predictive neural network models and computations in the brain, providing new leads that can enrich both fields.

he fields of neuroscience and machine learning have long enjoyed productive dialogue, with neuroscience offering inspiration for how artificial systems can be constructed, and machine learning providing tools for modelling and understanding biological neural systems. Recently, as deep convolutional neural networks (CNNs) have emerged as leading systems for visual recognition tasks, they have also emerged—without any modification or tailoring to purpose—as leading models for explaining the population responses of neurons in primate visual cortex¹⁻⁴. These results suggest that the connections between artificial deep networks and brains may be more than skin deep.

However, while deep CNNs capture some important details of the responses of visual cortical neurons, they fail to explain other key properties of the brain. Notably, the level of strong supervision typically used to train CNNs is much greater than that available to our brain. To the extent that representations in the brain are similar to those in CNNs trained on, for example, ImageNet⁵, the brain must be arriving at these representations by different, largely unsupervised routes. Another key difference is that the majority of CNNs optimized for image recognition and subsequently used to predict neural responses are feedforward and thus fundamentally static, lacking recurrence and a notion of time (with notable recent exceptions^{4,6,7}). Neuronal systems, in contrast, are highly dynamic, producing responses that vary dramatically in time, even in response to static inputs.

Here, inspired by past success in using 'out-of-the-box' artificial deep neural networks as models of the visual cortex, we explore whether modern predictive recurrent neural networks built for unsupervised learning can also explain critical properties of neuronal responses and perception. In particular, we consider a deep predictive coding network ('PredNet's), a network that learns to perform next-frame prediction in video sequences. The PredNet is motivated by the principle of predictive coding⁹⁻¹²; the network

continually generates predictions of future sensory data via a topdown path, and it sends prediction errors in its feedforward path. At its lowest layer, the network predicts the input pixels at the next time step, and it has been shown to make successful predictions in real-world settings (for example, car-mounted camera datasets¹³). The internal representations learned from video prediction also proved to be useful for subsequent decoding of underlying latent parameters of the video sequence, consistent with the suggestion of prediction as a useful loss function for unsupervised/'self'-supervised learning¹⁴⁻²².

Self-supervised learning through video prediction has a rich history in machine learning literature and is a highly active area of current research²³⁻³³. Early implementations of spatiotemporal predictive learning include the work of Elman²⁰, Softky¹⁴ and Hawkins²¹. Recent approaches have incorporated adversarial^{15,16,23,25} and variational^{24,26,27} techniques, as well as novel recurrent units^{28,34}. With use cases including anomaly detection^{35,36} and robotic planning^{29,37}, state-of-the-art models are capable of successful predictions in datasets ranging from action recognition^{24,28} to robotic arm movement^{19,31}. In the neuroscience community, predictive coding also has a rich history³⁸⁻⁴⁶. Rao and Ballard helped popularize the notion of predictive coding in neuroscience in 1999, proposing that spatial predictive coding could explain several extra-classical receptive field effects in primary visual cortex (V1), such as end-stopping⁹. Predictive coding has been proposed as an explanatory framework for phenomena in a variety of sensory systems⁴⁷⁻⁴⁹. The PredNet formulates temporal and spatial predictive coding principles in a deep learning framework to work on natural sequences, providing an opportunity to test a wide range of neuroscience phenomena using a single model.

In the following, we show that despite being trained only to predict next frames in natural sequences, the PredNet captures a wide array of seemingly unrelated fundamental properties of neuronal

¹Harvard University, Cambridge, MA, USA. ²Boston Children's Hospital, Harvard Medical School, Boston, MA, USA. ³Center for Brains, Minds, and Machines (CBMM), Cambridge, MA, USA. ⁴MIT-IBM Watson AI Lab, Cambridge, MA, USA. ⁵IBM Research, Cambridge, MA, USA. ⁶Present address: DeepHealth, Inc., Cambridge, MA, USA. ^{III} Iotter.bill1@gmail.com

responses and perception, even when probed with synthetic stimuli. We begin by demonstrating that the PredNet can mimic neuronal response properties at multiple levels of the visual hierarchy, including both spatial properties (end-stopping/length suppression) and temporal properties (on/off firing rate dynamics, temporal sequence learning effects). We then demonstrate that the PredNet can also capture aspects of perception, even under conditions where subjective interpretation is dissociated from visual input, such as the spatial completion in illusory contours and the dynamic illusion of the flash-lag effect.

Results

The deep predictive coding network proposed in ref.⁸ (PredNet) consists of repeated, stacked modules, where each module generates a prediction of its own feedforward inputs, computes errors between these predictions and the observed inputs, and then forwards these error signals to subsequent layers (Fig. 1; also see Methods). The residual errors from a given layer thus become the prediction targets for the layer above. The model consists of four components: targets to be predicted (A_l) , predictions (\hat{A}_l) , errors between predictions and targets (E_i) , and a recurrent representation from which predictions are made (R_i) . On an initial time step, the feedforward pass can be viewed as a standard CNN consisting of alternating convolutional and pooling layers. Because of the pooling operation, the feedforward receptive field becomes twice as large at each successive layer. The pooling output of each layer is set as the prediction target for that layer, with the target at the lowest layer set to the actual input image itself, corresponding to the next frame in the input video sequence. Predictions are made in a top-down pass via convolutions over the representational units, which are first updated using the representational units from the layer above and errors from the previous time step as inputs. The error modules, E_{b} are calculated as a simple difference between the targets (A_l) and predictions (\hat{A}_l) , followed by splitting into positive and negative error populations. The network is trained to minimize the activations of the error units across the training set using (truncated) backpropagation through time, with the error units at each layer contributing to the total loss. Similar to the original work⁸, the results presented here use a model trained for next-frame prediction on a car-mounted camera dataset (KITTI¹³). Thus, the model is trained in an unsupervised or 'self'-supervised manner that does not require any external labels or other forms of supervision. The model consists of four layers and, with 0-indexing used here, layer 1 would be analogous to primary visual cortex (V1).

PredNet can capture spatial and temporal single unit response properties

We begin by comparing the response properties of units in the PredNet to established single unit response properties of neurons in the primate visual system, which have been studied extensively using microelectrode recordings. We investigate both spatial and temporal properties, first illustrating that the spatiotemporally trained PredNet can reproduce effects dependent on spatial statistics, akin to the spatially trained model of Rao and Ballard9. We then examine the temporal aspects of responses in the network, demonstrating both short-term ('inference mode') and long-term ('training mode') properties that are consistent with biology. Throughout, we primarily compare responses in the PredNet's error ('E') units, the output units of each layer, to neuronal recordings in the superficial layers of cortex. In each comparison, we show that the PredNet qualitatively exhibits the effect and then provide a quantitative comparison to biology, primarily using the metrics defined in the original biological studies. For completeness, response properties of other units in the PredNet (for example, the 'R' units) are included in the Extended Data, and would likely map onto other parts of the cortical circuit (Discussion).

ARTICLES



Fig. 1 | Deep predictive coding networks (PredNets). a, Each layer *I* consists of representation neurons (*R*_i), which output a layer-specific prediction at each time step (\hat{A}_i), which is compared against a target (A_i) to produce an error term (*E*_i), which is then propagated laterally and vertically in the network. **b**, Module operations for the case of video sequences. The target at the lowest layer of the network, A_{or} is set to the actual next image in the sequence. Conv, convolution; Conv LSTM, convolutional long short-term memory^{72,73}; Pool, 2 × 2 max-pooling; ReLU, rectified linear unit activation.

End-stopping and length suppression. One of the earliest nonlinear effects discovered in recordings of visual cortex is the property of end-stopping⁵⁰. End-stopping, or length suppression, is the phenomenon where a neuron tuned for a particular orientation becomes less responsive to a bar at this orientation when the bar extends beyond its classical receptive field. The predictive coding explanation is that lines/edges tend to be continuous in nature and thus the centre of a long bar can be predicted from its flanks^{9,45}. A short, discontinuous bar, however, deviates from natural statistics, and responding neurons signal this deviation. One potential source for conveying the long-range predictions in the case of an extended bar could be feedback from higher visual areas with larger receptive fields. This hypothesis was elegantly tested in ref. ⁵¹ using reversible inactivation of secondary visual cortex (V2) paired with V1 recordings in the macaque. As illustrated in Fig. 2a, cryoloop cooling of V2 led to a significant reduction in length suppression, indicating that feedback from V2 to V1 is essential for the effect.

Figure 2b demonstrates that length suppression and its mediation through top-down feedback are also present in the PredNet. The upper left panel contains the mean normalized response for units in the E_1 layer to bars of different lengths and the remaining panels contain example units. The red curves correspond to the original network (trained on the KITTI dataset¹³) and the blue curves correspond to zero-ing the feedback from R_2 to R_1 . Quantifying percent length suppression (%LS) as $100 \times \frac{r_{max} - T_{longest bar}}{r_{max}}$, with *r* indicating the response, the mean decrease in %LS upon removing top-down signalling was $16 \pm 7\%$ (mean \pm s.e.m.) for E_1 units (P=0.014, Wilcoxon signed rank test, one-sided, z=2.2), which is similar to the 20% decrease observed in the V1 population on cooling V2 in ref.⁵¹.

On/off temporal dynamics. Prediction in space and prediction in time are inextricably intertwined. A particular core temporal aspect of the visual cortical response that has a predictive quality is the on/off response trajectory to static stimuli. Figure 3a shows a typical response profile of a visual cortical neuron to a static input⁵². The neuron, recorded in the secondary visual cortex (V2) of a macaque monkey, produces a brief transient response to the onset of the visual stimulus, followed by near total suppression of that response. When the stimulus is removed, the neuron responds again with a transient



Fig. 2 | Length suppression. a, Responses of example macaque V1 units to bars of different lengths before (red), during (blue) and after (green) inactivation of V2 via cryoloop cooling (dashed lines indicate spontaneous activity). **b**, PredNet after training on the KITTI car-mounted camera dataset¹³. Top left: data are presented as mean over E_1 filter channels (\pm s.e.m.). The remaining three panels present examples. Red, original network; blue, feedback weights from R_2 to R_1 set to zero. We note that, in the PredNet, the 'after' inactivation response (green trace in neural data) would be equivalent to the 'before' inactivation response (blue). See Extended Data Fig. 1 for responses of A and R units. Figure reproduced with permission from ref. ⁵¹, Society for Neuroscience.

burst of activity (known as an 'off' response). The unpredictability of the stimulus and its precise onset correlate to the 'on' response, which decays as the image remains fixed, consistent with the common, slow-moving or static nature of real-world objects. Finally, the sudden and unexpected disappearance of the object drives the 'off' response. Figure 3b shows the average response of PredNet E units in different layers over a set of 25 naturalistic objects appearing on a grey background. The on/off dynamics are apparent on the population average level for all four layers of the network. One time step after image onset, the decay in average response ranges from 20% (E_1) to 49% (E_3) . As a point of reference, macaque inferior temporal cortex (IT) data from ref. 53 exhibit a 44% reduction in population response 100 ms after post-onset peak of a static image on a grey background. This decay rate is thus of the same magnitude observed in the PredNet, given that one time step in the model is loosely analogous to 100 ms, since the model was trained at a rate of 10 Hz. As the *E* units are the input drive to much of the rest of the network, it might be expected that the on/off dynamics are also present in the A and R layers, and indeed this is the case, as illustrated in Extended Data Fig. 2.

Sequence learning effects in visual cortex. Although the on/off dynamics of visual cortical responses may indeed reflect learned statistics of the natural world, there are perhaps even more striking examples of the sensitivity of neural responses to the long-range temporal structure in the visual world. For example, Meyer and Olson⁵⁴ demonstrated that neurons in IT could be strongly modulated by prior experience with sequences of presented images. After repeated presentations of arbitrary images with predictable transition statistics (for example, 'image B always follows image A'), neurons appeared to learn the sequence statistics, responding robustly only to sequence transitions that were unexpected. Figure 4a shows

the mean response of 81 IT neurons for predicted and unpredicted pairs. Figure 4b demonstrates a similar effect in the PredNet after an analogous experiment. Initialized with the weights after training on the KITTI car-mounted camera dataset, the model was then trained on five image pairs for 800 repetitions, matching the number of trials in the Meyer and Olson experiment. Figure 4c-e shows an example sequence and the corresponding next-frame predictions before and after training on the image pairs. The model, prior to exposure to the images in this experiment (trained only on KITTI¹³), settles into a noisy, copy-last-frame prediction mode. After exposure, the model is able to successfully make predictions for the expected image pair (row 2). Because the chosen image pair is unknown a priori, the initial prediction is the constant grey background when the first image appears. The model then rapidly copies this image for the ensuing three frames. Next, the model successfully predicts the transition to the second object (a stack of tomatoes in this case). In Fig. 4e, a sequence that differs from the training pair is presented. The model still makes the prediction of a transition to tomatoes, even though a chair is presented, but then copies the chair into subsequent predictions. Figure 4b shows that the unexpected transitions result in a significantly larger average response in the final *E* layer of the network (E_3 ; P = 0.002, paired *t*-test, one-sided, t(4) = 6.0). The PredNet E_3 units and the IT neurons in ref. ⁵⁴ both, in fact, exhibit over a 2× increase in response to unpredicted versus predicted stimuli (158% increase for IT, 108% increase for E_3). In the PredNet, there is indeed a larger response to the unpredicted images in all layers and all unit types (E, A, R; Extended Data Fig. 3).

PredNet can capture spatial and temporal illusory aspects of visual perception

Visual illusions can provide powerful insight into the underpinnings of perception. Building on the spatial and temporal single-unit



Fig. 3 | On/off temporal dynamics. a, Example macaque V2 neuron responding to a static image (Im). **b**, PredNet response to a set of naturalistic objects appearing on a grey background, after training on the KITTI car-mounted camera dataset¹³. Responses are grouped by layer for the *E* units and averaged across all units (all receptive fields and filter channels) and all stimuli, per layer. The average response trace is then normalized to have a range from 0 to 1. The average response peaks upon image appearance and disappearance. Given the large number of units in each layer, the s.e.m. is *O*(1%) of the mean. Responses of *A* and *R* units are contained in Extended Data Fig. 2. Panel **a** adapted with permission from ref. ⁵², American Physiological Society.

response properties reproduced by the PredNet, we ask if the model can also capture complex aspects of visual perception when probed with spatial and temporal visual illusions. We examine the PredNet's responses to illusory contours, a primarily spatially predictive phenomenon, and the flash-lag illusion, which has aspects of both spatial and temporal prediction. We note also that the PredNet has recently been shown to predict the illusory motion perceived in the rotating snakes illusion⁵⁵. In the following experiments, the network again has only been trained on the KITTI dataset and is evaluated in inference mode (that is, there is no additional training for the specific stimuli used in this section).

Illusory contours. Illusory contours, as in the Kanizsa figures⁵⁶, elicit perceptions of edges and shapes, despite the lack of enclosing lines. An example of such a figure is displayed at the bottom of Fig. 5. Importantly, the percept of the illusion is highly dependent on the spatial configuration of the components of the figure, as rotating these components lessens the effect (for example, the 'rotated J' figure in the bottom of Fig. 5). Neural correlates of these illusions have been discovered in the responses of visual neurons. Lee and Nguyen⁵⁷ found that neurons in monkey V1 are responsive to illusory contours, albeit at a reduced response and increased latency compared to physical contours. Figure 5a contains an example of such a neuron. The stimuli in the experiment consisted of sequences starting with an image of four circles, which then abruptly transitioned to one of numerous test images, including the illusion. Illustrated in Fig. 5b, the population average of 49 superficial V1 neurons responded more strongly to the illusion than similar, but non-illusory stimuli. This preference was also apparent in V2, with a response that was, interestingly, of a shorter latency compared to V1 (Fig. 5c).

Figure 5d–f demonstrates that the core effects discovered by Lee and Nguyen⁵⁷ are also largely present in the PredNet. In the population average of E_1 units, there is indeed a response to the illusory contour, with an onset at an increased latency compared to the physical contours (Fig. 5d). Additionally, Fig. 5e illustrates that the average E_1 response was moderately higher for the illusory contour than the response to the similar control images. This was also the case for E_2 units, with a peak response one time step before E_1 (Fig. 5f). Indeed, the size of the stimuli was chosen such that it was larger than the feedforward receptive field of the layer 1 neurons, but smaller than that of the layer 2 neurons (matching the protocol of ref. ⁵⁷). Using metrics proposed by Lee and Nguyen⁵⁷ to quantify the preference of the illusion to the amodal and rotated images for each individual unit, we find that the average is positive (higher response to the illusion) for both comparisons and all tested layers in the PredNet (E_1 , E_2 , A_1 , A_2 , R_1 , R_2), though not all statistically significant (Extended Data Fig. 5).

The flash-lag effect. Another illusion for which prediction has been proposed as having a role is the flash-lag effect. Fundamentally, the flash-lag effect describes illusions where an unpredictable or intermittent stimulus (for example, a line or dot) is perceived as 'lagging' behind the percept of a predictably moving stimulus nearby, even when the stimuli are, in fact, precisely aligned in space and time⁵⁸⁻⁶⁰. These illusions are sometimes interpreted as evidence that the brain is performing inference to predict the likely true current position of a stimulus, even in spite of substantial latency (up to hundreds of milliseconds) in the visual system^{61,62}. The version of the illusion tested here consists of an inner, continuously rotating bar and an outer bar that periodically flashes on. Figure 6 contains example next-frame predictions by the PredNet on a sample sequence within the flash-lag stimulus. The model was again only trained on the KITTI car-mounted camera dataset, and then evaluated on the flash-lag stimulus in inference mode. The rotation speed of the inner bar in the clip was set to 6° per time step. The first feature of note is that the PredNet is indeed able to make reasonable next-frame predictions for the inner rotating bar. If the model simply copied the last seen frame at every time step instead of making an actual prediction, the angle between the inner rotating bar in the outputted 'predicted' frame would be 6° behind the bar in the actual next frame. Instead, the inner bar in the PredNet predictions is on average only $1.4 \pm 1.2^{\circ}$ (s.d.) behind the actual bar (see Methods for quantification of the bar angle). As the model was trained on real-world videos, generalization to this impoverished stimulus is non-trivial. Second, the post-flash predictions made by the model tend to resemble the perceived illusion. In the PredNet next-frame predictions, the outer and inner bars are not co-linear, similar to the illusory percept (see additional post-flash predictions in Extended Data Fig. 6). As opposed to being aligned with a 0° difference in the actual image when the outer bar appears, the inner bar in the PredNet predictions lags the predicted outer bar by an average of $6.8 \pm 2.0^{\circ}$ (s.d.). For rotation speeds up to and including 25 rotations per minute, we find that the average angular difference between the predicted bars in the PredNet aligns well with perceptual estimates (Extended Data Fig. 8). Considering that the model was trained for next-frame prediction on a corpus of natural videos, this suggests that our percept matches the statistically predicted next frame



Fig. 4 | Image sequence learning effects. a,**b**, Population responses to predicted versus unpredicted image transitions: mean of 81 neurons recorded in macaque IT (**a**); mean (\pm s.e.m.) across all PredNet E_3 units (all spatial receptive fields and filter channels) (**b**). **c**-**e**, Next-frame predictions by the PredNet: predictions of a KITTI-trained PredNet model on an example sequence (**c**); PredNet predictions after repeated 'training' on the sequence (**d**); PredNet predictions for an unpredicted image transition (**e**). Panel **a** reproduced with permission from ref. ⁵⁴, PNAS.

(as estimated by the PredNet) more than the actual observed frame. These results thus support an empirical, natural statistics interpretation of the flash-lag illusion⁶³.

Discussion

We have shown that a recurrent neural network trained to predict future video frames can explain a wide variety of seemingly unrelated phenomena observed in visual cortex and visual perception. These phenomena range from core properties of the responses of individual neurons to complex visual illusions. Critically, while previous models have been used to explain subsets of the described phenomena, we illustrate that a single core PredNet model trained on natural videos can reproduce all the phenomena without being explicitly designed to do so (Extended Data Fig. 7). Our work adds to a growing body of literature showing that deep neural networks exclusively trained to perform relevant tasks can serve as surprisingly good models of biological neural networks, often even outperforming models exclusively designed to explain neuroscience phenomena.

A particular conceptual advantage of the PredNet training scheme, compared to typical supervised neural network training, is that it does not involve large amounts of supervision in the form of paired inputs and labels, which are neither required for biological learning nor are typically found in real life. Prediction is a learning signal that comes for 'free'; that is, it is a form of unsupervised or self-supervised learning. A prediction can be compared to the actual observed state of the world, and the errors in that prediction can drive learning directly. In addition, there is also intrinsic

ARTICLES



Fig. 5 | Illusory contours. First row: data from electrophysiological recordings in the rhesus monkey. Second row: PredNet responses. **a**, An exemplar V1 neuron, which exhibits a response to the illusory contour at an increased latency compared to stimuli with true contours. **b**, V1 population average, demonstrating a larger response to the illusory stimuli compared to similar, control stimuli. **c**, The V2 population average response is also larger for the illusory stimuli, and demonstrates an earlier latency than the V1 average. **d**, The PredNet E_1 average activity also demonstrates a response to the illusory contour. **e**, The E_1 average response is moderately larger for the illusory contour than the control stimuli. **f**, The PredNet E_2 response is also moderately larger for the illusory stimuli, with an earlier latency than E_1 . PredNet averages were computed across filter channels at the central receptive field. Error bars represent s.e.m. Bottom: illustration of the stimuli and the presentation paradigm, using the nomenclature proposed by Lee and Nguyen⁵⁷. For each trial in the monkey and PredNet experiments, the 'four circles' stimuli is first presented, followed by one of the test stimuli (in brackets). See Extended Data Fig. 4 for the responses of the *A* and *R* units. Panels **a**-**c** reproduced with permission from ref. ⁵⁷, PNAS.

behavioural value in the ability to predict—both in time and space. Temporal prediction enables more effective planning of actions and can also help mitigate lags found in the relatively slow processing pipeline of visual cortex, while spatial prediction can help fill in information lost due to occlusion, for example.

Analysis of model components in producing observed effects. Although the PredNet reproduces a diverse range of phenomena observed in the brain, we would not claim that the PredNet is a perfect or exact model of the brain, or that its precise architecture per se is required for the observed effects. Thus, it is useful to ask which features and components of the model are necessary to reproduce the biological phenomena described above. For example, one key feature of the PredNet is recurrent connectivity, both within layers (intrinsic recurrent connections) and between layers (feedback connections). It is straightforward to see that some form of recurrence is required to observe temporal dynamics (such as 'on' and 'off' responses), because a strictly feedforward version of the PredNet would lack temporal dynamics altogether. Similarly, recurrent connections are essential for the PredNet to demonstrate phenomena such as length suppression and end-stopping in early layers. Although it is possible that a strictly feedforward, nonlinear network could show end-stopping and surround suppression-like effects in higher layers, where receptive fields are large enough to include both the 'centre' and 'surround', such suppression is not possible in lower layers of the network without recurrence.

Related to recurrence, it is also clear that depth, with an associated increase in feedforward receptive field size over layers, is necessary to produce the observed phenomena. For example, the larger 'classical' receptive field of the E_2 layer versus the E_1 layer in the PredNet, combined with feedback, facilitates the effects observed in the illusory contour experiment, where both layers produce a response to the illusory figure, but the E_2 response is earlier.

Although recurrence, depth and nonlinearity can be seen to be essential from a first-principles analysis of the PredNet, the necessity of other specific features of the PredNet is less obvious. One notable feature of the PredNet, motivated by predictive coding principles⁹, is that it explicitly computes an error representation in a population of neurons, wherein predicted inputs are explicitly subtracted from actual inputs, and the activity of these 'error' units is



Fig. 6 | The flash-lag effect. Top row: a segment of the stimulus clip inputted to the PredNet. Bottom row: PredNet predictions after training on the KITTI car-mounted camera dataset¹³. Each column represents the actual next frame in the sequence (above) and the outputted next frame prediction from the model (\hat{A}_0 ; below). At the time step indicated as t_0 , the outer bar flashes on in the actual sequence and is co-linear with the inner bar. The PredNet's post-flash prediction (corresponding to t_{+1}) displays the two bars as not co-linear, similar to the perceptual illusion. Additional post-flash predictions are contained in Extended Data Fig. 6.

what is passed from layer to layer in a feedforward manner. One key feature that this kind of explicit representation and propagation of errors induces is a force that drives the activity of subpopulations of neurons towards zero. That is, with errors represented by the activity of explicit error-coding units, and a training objective based on reducing these errors, there is an explicit mechanism to encourage unit activity to go to zero. From a machine learning perspective, it is straightforward to design a version of the PredNet that is still trained to predict future inputs, but for which 'errors' are not passed (Extended Data Fig. 9). Biologically, this could still correspond to a loss/errors being instantiated by neurons, but where these neurons do not serve as a core drive for activity in the rest of the network. On training a version of the PredNet with removal of the error passing in the network, we find that it generally less faithfully reproduces the neural phenomena presented here (Extended Data Fig. 10). For example, it might be expected that the decay in unit activity after an 'on' response would be less dramatic in this control network than in the original PredNet and neural data, and indeed that is the case. Additionally, the control network actually exhibits enhanced length suppression upon the removal of top-down feedback and a decrease in response upon presentation of the illusory contours stimulus. However, some qualitative effects, such as a larger response to unexpected versus expected stimuli in the sequence learning experiment, are still present in this control network, suggesting that explicit error activity passing may or may not be essential to explain these phenomena. We note that, as the explicit error passing in the PredNet seems to improve overall biological faithfulness here, it was also demonstrated to improve next-frame prediction performance in the original work⁸. Specifically, compared to a network with a similar architecture to the PredNet except for lacking layer-wise error computations, the PredNet performed better in next-frame prediction for both synthetic and natural stimuli. The biologically inspired splitting of positive and negative errors at each layer was additionally illustrated to improve prediction performance.

Because the reduction of network activity induced by error propagation has some correlation with the observed effects in the PredNet, one might wonder whether other means of minimizing activity are sufficient to produce the effects, without necessarily requiring temporal prediction. For example, sparse coding-style networks also tend to minimize overall activity, and sparse coding has been invoked as a possible explanation for phenomena such as end-stopping⁶⁴. Sparse coding models are typically trained with a reconstruction loss, that is, a loss function based on representing the current stimulus, and, critically, they impose an L_1 penalty on activations. For static stimuli, it might be expected that reconstructive

and predictive models will behave similarly. However, the predictive 'what will appear next' nature of the sequence learning and flash-lag effect experiments described above lacks an obvious explanation in the context of a purely static reconstructive loss. Nonetheless, it is certainly conceivable that various timeframes of sensory input estimation, from past to present (reconstruction) and future, are utilized as a learning signal and encoding strategy in the brain^{65,66}.

Comparison of model components to biology. Many of the core features of the PredNet architecture-recurrent connectivity, depth with increasing receptive field size and activity minimization through explicit computation of errors- are central to reproducing the presented phenomena, but also correspond well with the known constraints of biological circuits. However, we note that the PredNet also contains elements that are not biologically plausible, or for which the mapping to biological implementation is not yet clear. Chief among these deviations from biology is the fact that the model uses scalar valued ('rate') activations rather than spiking, and that the model uses backpropagation for training. The extent to which these deviations matter is unclear. Some efforts have been made to show that rate-based models can be converted to spiking models⁶⁷, although there are numerous compelling computational proposals for ways that spike-based computation may be qualitatively different from rate-based computation68,69. The backpropagation algorithm used to train the PredNet requires updating neuronal connections with non-local information, and thus it is often cited as a key biologically implausible element of artificial neural networks. However, recent work has suggested several avenues by which backpropagation-like computations might be implemented with biological neurons70,71.

To the extent that the PredNet might mimic the architecture of cortex, it is interesting to consider how the elements of the model might map onto the elements of real cortical microcircuits. Indeed, it has been suggested that there is a tight correspondence between the canonical microcircuit and the connectivity pattern implied by predictive coding⁴⁰. The structure of a layer in the PredNet could also be seen as consistent with this mapping. In this view, the A_1 units in the PredNet would correspond to granular (L4) layer neurons in cortex, which largely serve as targets for feedforward inputs. The E_1 units would correspond to superficial (L1/2/3) layers, which receive input from the $A_1/L4$ neurons. The E_1 (superficial) neurons then serve as outputs of the circuit, passing information to subsequent, higher areas. These neurons also output to deep (L5/6) layers within the same microcircuit, which would correspond to the R_1 units in the PredNet. Finally, completing the circuit, the deep (R_1)

units input onto the granular (A_l/\hat{A}_l) layer. Interestingly, we find that there is some variation in the response characteristics among the unit types in the PredNet; specifically, the average R response does not exhibit length suppression (Extended Data Fig. 1) and also shows a smaller 'surprise' response compared to the A and E units (Extended Data Fig. 3). This raises an intriguing and testable question of whether such differences also exist between deep and superficial units in the brain. Finally, we note that in the PredNet at least, it is partly notation whether it is said that the 'activations' (A_l) or the 'errors' (E_l) are passed between layers, and the effects observed here in the *E* units are also present in the *A* units (Extended Data Figs. 1, 2, 3, 4 and 5). Overall, we do not intend to claim that there are precise classifications of 'error' versus 'activation/feature' neurons per se, rather that both of these types of computation are important and could map to the canonical microcircuit, with potentially even individual neurons providing a combination of both computations.

Conclusion

Neuroscience has been a long-standing inspiration for machine learning, where a core goal is to develop models with brain-like abilities. Conversely, developing computational models that reproduce and explain neural phenomena is a central aim in neuroscience. Here, we present an example of this cyclical dialogue, showing that a deep learning model inspired by theories of brain computation can reproduce a wide array of phenomena observed in visual cortex and visual perception. Importantly, the model was trained purely with a self-supervised, predictive loss. An especially salient motivation for pursuing such unsupervised/self-supervised methods is the ability of humans to excel in these regimes. Despite tremendous progress, artificial intelligence (AI) systems today still lag well behind humans in critical properties including extrapolation across domains, few shot learning and transfer learning. Thus, the ability of a self-supervised model to generalize from training on car-mounted camera videos to testing on impoverished, synthetic stimuli provides further inspiration for incorporating cognitive and neural constraints in designing AI models. In particular, that a simple objective-prediction-can produce such a wide variety of observed neural phenomena as demonstrated here underscores the idea that prediction may be a central organizing principle in the brain.

Methods

PredNet background. The original description of the PredNet can be found in ref.⁸. Briefly, the PredNet consists of a hierarchical stack of modules, where each module contains four different unit types: representational units (R_i) from which predictions are generated (\hat{A}_l) , targets to be predicted (A_l) and error units (E_l) , where l indicates the layer in the network. At each time step, the R_l units are first updated via a top-down pass, receiving input from both the error units at the same level (E_i) and the representational units from the layer above (R_{i+1}) , which are first spatially upsampled (nearest-neighbour) to match the spatial size of layer l. The R_l units are implemented as convolutional long short-term memory (LSTM) units7 After updating the R_l units, a bottom-up pass is made where the predicted next frame is first generated (\hat{A}_0) via a convolution of the R_0 units. The actual input frame, A_0 , is compared to \hat{A}_0 via unit-wise subtraction, followed by splitting into positive and negative error populations, forming E_0 . The splitting of the error populations is motivated by the existence of on-centre/off-surround and off-centre/ on-surround neurons in the early visual system. E_0 becomes the input into the next layer of the network, from which A_i is generated via a convolution over E_0 , followed by a 2×2 max-pooling operation. A prediction at this layer is generated via a convolution over R_1 and then this process is repeated forward in the network until errors are calculated at each level. A summary of the computations performed by each unit is contained in equations (1) to (4) in the following.

Given an input sequence of images, x_p the units at each layer l and time step t are updated according to

$$A_{l}^{t} = \begin{cases} x_{t} & \text{if } l = 0\\ \text{MaxPool}(\text{ReLU}(\text{Conv}(E_{l-1}^{t}))) & l > 0 \end{cases}$$
(1)

$$\hat{A}_{l}^{t} = \operatorname{ReLU}(\operatorname{Conv}(R_{l}^{t}))$$
(2)

ARTICLES

$$E_l^t = \left[\text{ReLU}(A_l^t - \hat{A}_l^t); \text{ ReLU}(\hat{A}_l^t - A_l^t) \right]$$
(3)

$$R_l^t = \text{ConvLSTM}(E_l^{t-1}, R_l^{t-1}, \text{Upsample}(R_{l+1}^t))$$
(4)

The only modification to the original PredNet that we make here, for the sake of biological interpretability, is replacing the tanh output activation function for the LSTMs with a ReLU activation (ReLU(x) = max(x, 0)), enforcing positive 'firing rates'. On the KITTI dataset this leads to a marginally (8%) worse prediction mean-squared error (MSE) than the standard formulation, but it is still 2.6 times better than the MSE that would be obtained by simply copying the last frame seen (compared to 2.8 for tanh).

The loss function of the PredNet is implemented as the weighted sum of the error unit activations at each layer and time step. The model is thus 'generative' in the sense that it generates predictions of future input data given previous input data, but not in the sense of an explicit probabilistic formulation, although future work could explore incorporating generative adversarial74 or variational75 components. We use the 'Lall' version of the model here, placing a non-zero loss weight on each layer in the network. For model training, weights are updated via backpropagation76 (through time) using the Adam optimizer77. The dataset used for training is the KITTI dataset13, a collection of videos obtained from a car-mounted camera while driving in Germany. The same training and pre-processing procedures were used as in the original PredNet paper, including training using sequences of 10 frames, with each frame centre-cropped and downsampled to 128×160 pixels. The numbers of filter channels (for example, convolutional kernels) per layer for both the A and R modules are 3, 48, 96 and 198, from layers 0 to 3, respectively. Given a 128×160 image, this means that there are $\frac{128}{2} \times \frac{160}{2} \times 48 = 245,760$ units in the A_1 and R_1 layers, for example, given the 2×2 max-pooling between each layer. There are thus 122,880 and 61,440 units in the A_2/R_2 and A_3/R_3 layers, respectively. For each layer of the hierarchy, there are twice as many E units given the splitting into positive and negative errors. Code for the PredNet, including training on the KITTI dataset, is available at https://github. com/coxlab/prednet.

End-stopping and length suppression. For each convolutional kernel in the PredNet, length suppression was evaluated at the central receptive field, with input images of size 128×128 pixels. We follow ref. ⁵¹ by first determining each unit's preferred orientation, implemented by measuring responses to Gabor filters at different orientations. Filters with a wavelength and envelope standard deviation of 5 pixels were used, and responses were summed over the presentation of ten time steps, after the presentation of a grey background for five time steps. Given the preferred orientation for each unit, bars of width 1 pixel and varying length were presented at this orientation. For each bar, a grey background was again presented for five time steps and then the bar was presented for ten time steps, with the total response quantified as the sum of the response over the ten time steps. A population average over all units was quantified by following the procedure above and then normalizing each unit to have a maximum response of 1, followed by averaging. Removal of feedback was implemented by setting the connection weights from R_2 to R_1 to zero. Statistical analysis comparing the original network to the removal of feedback was performed using units that had a non-constant response in both conditions, which amounted to 28 out of 96 units in E_1 , 21 out of 48 units in A_1 and 30 out of 48 units in R_1 (see Extended Data Fig. 1 for A_1 and R_1 results).

On/off temporal dynamics. The stimuli for the temporal dynamics experiment consisted of objects appearing on a grey background with images of size 128×128 pixels. A set of 25 objects was used, with examples displayed in Fig. 4. The input sequences consisted of a grey background for seven time steps, followed by an object on the background for six time steps. For comparing response decay rates in the PredNet to the macaque IT data from ref. ⁵³, the population average of single unit activity in the IT data was used.

Sequence learning effects in visual cortex. Stimuli for the sequence learning experiments in the PredNet consisted of five randomly chosen image pairs from a set of 25 images of objects appearing on a grey background of 128×128 pixels. Each image appeared in only one set of pairs. The training portion of the experiment (starting from the KITTI-trained weights) consisted of presenting each pair 800 times, matching the number of trials in the Meyer and Olson experiment⁴. Each trial consisted of a grey background for four time steps, followed by the first image for four time steps, then the second image for four time steps and finally the grey background again for four time steps. For model updates, the Adam⁷⁷ optimizer was used with default parameters. For testing, unpredicted pairs were created by randomly permuting the second images across the pairs. The population response in Fig. 4 was quantified by averaging across all units and image pairs (five predicted and unpredicted pairs) and normalizing this response to have a maximum of 1 across the duration of the trial. The difference between predicted and unpredicted responses was assessed at the peak of the response for the second image.

Illusory contours. PredNet responses in the illusory contours experiment were evaluated using units at the central receptive field for each convolutional kernel,

ARTICLES

using input images of size 128×128 pixels. Similar to the neural experiments by Lee and Nguven⁵⁷, the preferred orientation for each unit was first determined using a short bar stimulus, specifically a 1 pixel wide bar with a length of 8 pixels. Responses to the bar at different orientations were quantified as the sum of the response over a presentation of ten time steps, after the presentation of a grey background for five time steps. Responses to the test stimuli were then evaluated when presenting at the optimal orientation for each unit, meaning that, for example, one edge of the 'line square' (Fig. 5) was centred around the unit's receptive field and oriented at the unit's preferred orientation. The test sequences consisted of a grey background for five time steps, followed by the 'four circles' image for ten time steps and finally one of the test images for ten time steps. For the stimuli involving circles, the radius of the circles was set to 4 pixels with the distance between the centres of adjacent circles (or, equivalently, the length of a side in the square stimuli) set to 16 pixels. These sizes were chosen because 4 pixels is twice the size of the feedforward receptive field in the E_1 layer. The radius used in the Lee and Nguyen57 experiments was also approximately twice as large as the mapped receptive fields in V1. In Fig. 5d-f, the population response was calculated as an average over the responses of individual units, where the response of each unit was first normalized by division of the unit's max response over all stimuli. To be included in the population response, a unit had to have a non-zero response to the bar stimulus (at any orientation) and a non-zero response to at least one of the test sequences. The number of units meeting this criteria was 37 (out of 96) for E_1 , 32 (out of 192) for *E*₂, 27 (out of 48) for *A*₁, 32 (out of 96) for *A*₂, 31 (out of 48) for R₁ and 54 (out of 96) for R₂ (see Extended Data Figs. 4 and 5 for A and R results).

The flash-lag effect. The flash-lag stimulus was created with a rotation speed of 6° per time step, with a flash every six time steps for three full rotations and an input size of 160×160 pixels. Angles of the bars in the predictions were quantified over the last two rotations to allow a 'burn-in' period. The angles of the predicted bars were estimated by calculating the mean-squared error between the prediction and a probe bar generated at 0.1° increments and a range of centres, and taking the angle with the minimum mean-squared error.

Data availability

The primary dataset used in this work is the KITTI Dataset¹³, which can be obtained at http://www.cvlibs.net/datasets/kitti/raw_data.php. All other data may be obtained upon request to the authors.

Code availability

Code for the PredNet model is available at https://github.com/coxlab/prednet. All other code may be obtained upon request to the authors.

Received: 11 September 2019; Accepted: 13 March 2020; Published online: 20 April 2020

References

- Yamins, D. L. K. et al. Performance-optimized hierarchical models predict neural responses in higher visual cortex. *Proc. Natl Acad. Sci. USA* 111, 8619–8624 (2014).
- Yamins, D. L. K. & DiCarlo, J. J. Using goal-driven deep learning models to understand sensory cortex. *Nat. Neurosci.* 19, 356–365 (2016).
- Khaligh-Razavi, S.-M. & Kriegeskorte, N. Deep supervised, but not unsupervised, models may explain IT cortical representation. *PLoS Comput. Biol.* 10, 1–29 (2014).
- Nayebi, A. et al. Task-driven convolutional recurrent models of the visual system. In Advances in Neural Information Processing Systems 5290–5301 (NeurIPS, 2018).
- Deng, J. et al. ImageNet: a large-scale hierarchical image database. In IEEE Conference on Computer Vision and Pattern Recognition 248–255 (IEEE, 2009).
- Tang, H. et al. Recurrent computations for visual pattern completion. Proc. Natl Acad. Sci. USA 115, 8835–8840 (2018).
- 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. *Nat. Neurosci.* 22, 974–983 (2019).
- Lotter, W., Kreiman, G. & Cox, D. D. Deep predictive coding networks for video prediction and unsupervised learning. *International Conference on Learning Representations* (ICLR, 2017).
- Rao, R. P. N. & Ballard, D. H. Predictive coding in the visual cortex: a functional interpretation of some extra-classical receptive-field effects. *Nat. Neurosci.* 2, 79–87 (1999).
- Friston, K. A theory of cortical responses. *Philos. Trans. R Soc. Lond. B Biol.* Sci. 360, 815–836 (2005).
- 11. Spratling, M. W. Unsupervised learning of generative and discriminative weights encoding elementary image components in a predictive coding model of cortical function. *Neural Comput.* **24**, 60–103 (2012).
- Wen, H. et al. Deep predictive coding network for object recognition. Proc. 35th International Conference on Machine Learning 80, 5266–5275 (2018).

NATURE MACHINE INTELLIGENCE

- Geiger, A., Lenz, P., Stiller, C. & Urtasun, R. Vision meets robotics: the KITTI dataset. *Int. J. Robot. Res.* 32, 1231–1237 (2013).
- Softky, W. R. Unsupervised pixel-prediction. In Advances in Neural Information Processing Systems 809–815 (NeurIPS, 1996).
- Lotter, W., Kreiman, G. & Cox, D. Unsupervised learning of visual structure using predictive generative networks. *International Conference on Learning Representations* (ICLR, 2016).
- Mathieu, M., Couprie, C. & LeCun, Y. Deep multi-scale video prediction beyond mean square error. *International Conference on Learning Representations* (ICLR, 2016).
- Srivastava, N., Mansimov, E. & Salakhutdinov, R. Unsupervised learning of video representations using LSTMs. Proc. 32nd International Conference on Machine Learning 37, 843–852 (2015).
- Dosovitskiy, A. & Koltun, V. Learning to act by predicting the future. International Conference on Learning Representations (ICLR, 2017).
- Finn, C., Goodfellow, I. J. & Levine, S. Unsupervised learning for physical interaction through video prediction. In *Advances in Neural Information Processing Systems* 64–72 (NeurIPS, 2016).
- 20. Elman, J. L. Finding structure in time. Cogn. Sci. 14, 179–211 (1990).
- 21. Hawkins, J. & Blakeslee, S. *On Intelligence* (Times Books, 2004).
- Luo, Z., Peng, B., Huang, D.-A., Alahi, A. & Fei-Fei, L. Unsupervised learning of long-term motion dynamics for videos. In *The IEEE Conference on Computer Vision and Pattern Recognition* 7101–7110 (IEEE, 2017).
- Lee, A. X. et al. Stochastic adversarial video prediction. Preprint at https:// arxiv.org/pdf/1804.01523.pdf (2018).
- 24. Villegas, R. et al. High fidelity video prediction with large stochastic recurrent neural networks. In *Advances in Neural Information Processing Systems* 81–91 (NeurIPS, 2019).
- Villegas, R., Yang, J., Hong, S., Lin, X. & Lee, H. Learning to generate long-term future via hierarchical prediction. *International Conference on Learning Representations* (ICLR, 2017).
- Denton, E. & Fergus, R. Stochastic video generation with a learned prior. In Proceedings of 35th International Conference on Machine Learning 1174–1183 (ICML, 2018).
- Babaeizadeh, M., Finn, C., Erhan, D., Campbell, R. H. & Levine, S. Stochastic variational video prediction. *International Conference on Learning Representations* (ICLR, 2018).
- Wang, Y., Gao, Z., Long, M., Wang, J. & Yu, P. S. PredRNN++: towards a resolution of the deep-in-time dilemma in spatiotemporal predictive learning. *Proc. 35th International Conference on Machine Learning* 80, 5123–5132 (2018).
- Finn, C. & Levine, S. Deep visual foresight for planning robot motion. In International Conference on Robotics and Automation 2786–2793 (IEEE, 2017).
- Hsieh, J.-T., Liu, B., Huang, D.-A., Fei-Fei, L. F. & Niebles, J. C. Learning to decompose and disentangle representations for video prediction. In *Advances in Neural Information Processing Systems* 517–526 (NeurIPS, 2018).
- Kalchbrenner, N. et al. Video pixel networks. Proc. 34th International Conference on Machine Learning 70, 1771–1779 (2017).
- 32. Qiu, J., Huang, G. & Lee, T. S. Visual sequence learning in hierarchical prediction networks and primate visual cortex. In Advances in Neural Information Processing Systems 2662–2673 (NeurIPS, 2019).
- 33. Wang, Y. et al. Eidetic 3D LSTM: a model for video prediction and beyond. *International Conference on Learning Representations* (ICLR, 2019).
- Wang, Y., Long, M., Wang, J., Gao, Z. & Yu, P. S. PredRNN: recurrent neural networks for predictive learning using spatiotemporal LSTMs. In Advances in Neural Information Processing Systems 879–888 (NeurIPS, 2017).
- Liu, W., Luo, W., Lian, D. & Gao, S. Future frame prediction for anomaly detection - a new baseline. In *The IEEE Conference on Computer Vision and Pattern Recognition* 6536–6545 (IEEE, 2018).
- 36. Tandiya, N., Jauhar, A., Marojevic, V. & Reed, J. H. Deep predictive coding neural network for RF anomaly detection in wireless networks. In 2018 IEEE International Conference on Communications Workshops (IEEE, 2018).
- Ebert, F. et al. Visual Foresight: Model-based deep reinforcement learning for vision-based robotic control. Preprint at https://arxiv.org/pdf/1812.00568.pdf (2018).
- Rao, R. P. N. & Sejnowski, T. J. Predictive sequence learning in recurrent neocortical circuits. In Advances in Neural Information Processing Systems 164–170 (NeurIPS, 2000).
- 39. Summerfield, C. et al. Predictive codes for forthcoming perception in the frontal cortex. *Science* **314**, 1311–1314 (2006).
- Bastos, A. M. et al. Canonical microcircuits for predictive coding. *Neuron* 76, 695–711 (2012).
- Kanai, R., Komura, Y., Shipp, S. & Friston, K. Cerebral hierarchies: predictive processing, precision and the pulvinar. *Philos. Trans. R. Soc. Lond. BBiol. Sci* 370, 20140169 (2015).
- Srinivasan, M. V., Laughlin, S. B. & Dubs, A. Predictive coding: a fresh view of inhibition in the retina. *Proc. R. Soc. Lond. B Biol. Sci.* 216, 427–459 (1982).
- Atick, J. J. Could information theory provide an ecological theory of sensory processing. Network: Computation in neural systems 22, 4–44 (1992).

- 44. Murray, S. O., Kersten, D., Olshausen, B. A., Schrater, P. & Woods, D. L. Shape perception reduces activity in human primary visual cortex. *Proc. Natl Acad. Sci. USA* **99**, 15164–15169 (2002).
- Spratling, M. W. Predictive coding as a model of response properties in cortical area V1. J. Neurosci. 30, 3531–3543 (2010).
- Jehee, J. F. M. & Ballard, D. H. Predictive feedback can account for biphasic responses in the lateral geniculate nucleus. *PLoS Comput. Biol.* 5, 1–10 (2009).
- 47. Kumar, S. et al. Predictive coding and pitch processing in the auditory cortex. J. Cogn. Neurosci. 23, 3084–3094 (2011).
- Zelano, C., Mohanty, A. & Gottfried, J. A. Olfactory predictive codes and stimulus templates in piriform cortex. *Neuron* 72, 178–187 (2011).
- Mumford, D. On the computational architecture of the neocortex: II The role of cortico-cortical loops. *Biol. Cybern.* 66, 241–251 (1992).
- Hubel, D. H. & Wiesel, T. N. Receptive fields and functional architecture of monkey striate cortex. J. Physiol. 195, 215–243 (1968).
- Nassi, J. J., Lomber, S. G. & Born, R. T. Corticocortical feedback contributes to surround suppression in V1 of the alert primate. *J. Neurosci.* 33, 8504–8517 (2013).
- 52. Schmolesky, M. T. et al. Signal timing across the macaque visual system. J. Neurophysiol. **79**, 3272–3278 (1998).
- Hung, C. P., Kreiman, G., Poggio, T. & DiCarlo, J. J. Fast readout of object identity from macaque inferior temporal cortex. *Science* 310, 863–866 (2005).
- Meyer, T. & Olson, C. R. Statistical learning of visual transitions in monkey inferotemporal cortex. *Proc. Natl Acad. Sci. USA* 108, 19401–19406 (2011).
- Watanabe, E., Kitaoka, A., Sakamoto, K., Yasugi, M. & Tanaka, K. Illusory motion reproduced by deep neural networks trained for prediction. *Front. Psychol.* 9, 345 (2018).
- 56. Kanizsa, G. Organization in Vision: Essays on Gestalt Perception (Praeger, 1979). 57. Lee, T. S. & Nguyen, M. Dynamics of subjective contour formation in the
- early visual cortex. *Proc. Natl Acad. Sci. USA* **98**, 1907–1911 (2001). 58. Nijhawan, R. Motion extrapolation in catching. *Nature* **370**, 256–257 (1994).
- 59. Mackay, D. M. Perceptual stability of a stroboscopically lit visual field
- containing self-luminous objects. *Nature* 181, 507–508 (1958).
 60. Eagleman, D. M. & Sejnowski, T. J. Motion integration and postdiction in visual awareness. *Science* 287, 2036–2038 (2000).
- Khoei, M. A., Masson, G. S. & Perrinet, L. U. The flash-lag effect as a motion-based predictive shift. *PLoS Comput. Biol.* 13, 1–31 (2017).
- Hogendoorn, H. & Burkitt, A. N. Predictive coding with neural transmission delays: a real-time temporal alignment hypothesis. *eNeuro* 6, e0412–18.2019 (2019).
- Wojtach, W. T., Sung, K., Truong, S. & Purves, D. An empirical explanation of the flash-lag effect. *Proc. Natl Acad. Sci. USA* 105, 16338–16343 (2008).
- 64. Zhu, M. & Rozell, C. J. Visual nonclassical receptive field effects emerge from sparse coding in a dynamical system. *PLoS Comput. Biol.* 9, 1–15 (2013).
- Chalk, M., Marre, O. & Tkačik, G. Toward a unified theory of efficient, predictive and sparse coding. *Proc. Natl Acad. Sci. USA* 115, 186–191 (2018).
- 66. Singer, Y. et al. Sensory cortex is optimized for prediction of future input. *eLife* 7, e31557 (2018).

- Hunsberger, E. & Eliasmith, C. Training spiking deep networks for neuromorphic hardware. Preprint at https://arxiv.org/pdf/1611.05141.pdf (2016).
- Boerlin, M., Machens, C. K. & Denève, S. Predictive coding of dynamical variables in balanced spiking networks. *PLoS Comput. Biol.* 9, 1–16 (2013).
- Maass, W. in Pulsed Neural Networks (eds Maass, W. & Bishop, C. M.) 55–85 (MIT Press, 1999).
- Nøkland, A. Direct feedback alignment provides learning in deep neural networks, In Advances in Neural Information Processing Systems 1037–1045 (NeurIPS, 2016).
- Lillicrap, T. P., Cownden, D., Tweed, D. B. & Akerman, C. J. Random synaptic feedback weights support error backpropagation for deep learning. *Nat. Commun.* 7, 13276 (2016).
- Hochreiter, S. & Schmidhuber, J. Long short-term memory. *Neural Comput.* 9, 1735–1780 (1997).
- Shi, X. et al. Convolutional LSTM network: a machine learning approach for precipitation nowcasting. In Advances in Neural Information Processing Systems 802–810 (NeurIPS, 2015).
- Goodfellow, I. et al. Generative adversarial nets. In Advances in Neural Information Processing Systems 2672–2680 (NeurIPS, 2014).
- Kingma, D. P. & Welling, M. Auto-encoding variational bayes. International Conference on Learning Representations (ICLR, 2014).
- Rumelhart, D. E., Hinton, G. E. & Williams, R. J. Learning representations by back-propagating errors. *Nature* 323, 533–536 (1986).
- 77. Kingma, D. P. & Ba, J. Adam: a method for stochastic optimization. International Conference on Learning Representations (ICLR, 2015).

Acknowledgements

This work was supported by IARPA (contract no. D16PC00002), the National Science Foundation (NSF IIS 1409097) and the Center for Brains, Minds and Machines (CBMM, NSF STC award CCF-1231216).

Author contributions

W.L. and D.C. conceived the study. W.L. conceived the model and implemented the experiments and analysis. G.K. and D.C. supervised the study. All authors contributed to interpreting the results. All authors contributed to writing the manuscript.

Competing interests

The authors declare no competing interests.

Additional information

Extended data is available for this paper at https://doi.org/10.1038/s42256-020-0170-9.

Correspondence and requests for materials should be addressed to W.L.

Reprints and permissions information is available at www.nature.com/reprints.

Publisher's note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

© The Author(s), under exclusive licence to Springer Nature Limited 2020



Extended Data Fig. 1 [Length suppression analysis for A_1 and R_1 units. The average (\pm s.e.m) response of A_1 and R_1 units and exemplars are shown (expanding upon Fig. 2 in main text). Red: Original network. Blue: Feedback weights from R_2 to R_1 set to zero. The average A_1 response demonstrates length suppression, whereas the average R_1 response does not show a strong effect, with some units overall showing length suppression (for example, unit 15 - bottom right panel) and other units showing an opposite effect (for example, unit 33 - bottom middle panel). The removal of feedback led to a significant decrease in length suppression in A_1 , with a mean (\pm s.e.m) decrease in percent length suppression ($%LS = 100^* \frac{r_{max} - r_{longest bar}}{r_{max}}$) of $31\pm7\%$ (p=0.0004, Wilcoxon signed rank test, one-sided, z=3.3). The R_1 units exhibited a mean %LS decrease of $5\pm6\%$ upon removal of feedback, which was not statistically significant (p=0.18, z=0.93).



Extended Data Fig. 2 | Temporal dynamics in the *A* and *R* units in the PredNet. The average response of *A* and *R* units to a set of naturalistic objects on a gray background, after training on the KITTI car-mounted camera dataset¹³ is shown (expanding upon Fig. 3 in the main text). The *A* and *R* layers seem to generally exhibit on/off dynamics, similar to the *E* layers. R_1 also seems to have another mode in its response, specifically a ramp up between time steps 3 and 5 post image onset. The responses are grouped per layer and consist of an average across all the units (all filters and spatial locations) in a layer. The mean responses were then normalized between 0 and 1. Given the large number of units in each layer, the s.e.m. is *O*(1%) of the mean. Responses for layer 0, the pixel layer, are omitted because of their heavy dependence on the input pixels for the *A* and *R* layers. Note that, by notation in the network's update rules, the input image reaches the *R* layers at a time step after the *E* and *A* layers.

Unit Type	Layer 0	Layer 1	Layer 2	Layer 3
E	308*	90**	109**	108**
Α	N/A	78**	109**	108**
R	N/A	18	19	30*

Extended Data Fig. 3 | Response differential between predicted and unpredicted sequences in the sequence learning experiment. The percent increase of population peak response between predicted and unpredicted sequences is quantified for each PredNet layer. Positive values indicate a higher response for unpredicted sequences. *p < 0.05, **p < 0.005 (paired t-test, one-sided).



Extended Data Fig. 4 | Illusory contours responses for A and R units in the PredNet. The mean \pm s.e.m. is shown (expanding upon Fig. 5 in the main text). Averages are computed across filter channels at the central receptive field.

Source	Layer	IC_A	IC_R	Layer	IC_A	IC_R
Monkey A Monkey B PredNet PredNet	$V1S \\ V1S \\ E_1 \\ A_1$	$0.19*\ 0.10*\ 0.09\pm 0.06\ 0.03\pm 0.05$	$0.31^{*} \ 0.16^{*} \ 0.14 \pm 0.06^{*} \ 0.10 \pm 0.05^{*}$	$\begin{array}{c} \textbf{V2S}\\ \textbf{V2S}\\ E_2\\ A_2 \end{array}$	$0.21*\ 0.08*\ 0.15\pm 0.07*\ 0.15\pm 0.07*$	$0.11*\ 0.12*\ 0.09\pm 0.07\ 0.09\pm 0.07$
Monkey A Monkey B PredNet	V1D V1D R ₁	$0.09* \\ 0.04* \\ 0.11 \pm 0.05*$	$0.11*\ 0.13*\ 0.04\pm 0.03$	V2D V2D R ₂	$0.28* \\ 0.07* \\ 0.12 \pm 0.04*$	$0.24* \\ 0.20* \\ 0.03 \pm 0.05$

Extended Data Fig. 5 | Quantification of illusory responsiveness in the illusory contours experiment. Units in the monkey recordings of Lee and Nguyen⁵⁷ are compared to units in the PredNet. We follow Lee and Nguyen⁵⁷ in calculating the following two measures for each unit: $IC_a = \frac{R_i - R_a}{R_i + R_a}$ and $IC_r = \frac{R_i - R_a}{R_i + R_a}$, where R_i is the response to the illusory contour (sum over stimulus duration), R_a is the response to amodal stimuli, and R_i is the response to the rotated image. For the PredNet, these indices were calculated separately for each unit (at the central receptive field) with a non-uniform response. Positive values, indicating preferences to the illusion, were observed for all subgroups. Mean \pm s.e.m.; *p < 0.05 (t-test, one-sided).

ARTICLES



Extended Data Fig. 6 | Additional predictions by the PredNet model in the flash lag experiment. The images shown consist of next-frame predictions by the PredNet model after four consecutive appearances of the outer bar. The model was trained on the KITTI car-mounted camera dataset¹³.

ARTICLES

NATURE MACHINE INTELLIGENCE

	Model Name	Spatial Predictive Coding	Feedforward LN model with high pass temporal kernel	Retina Response CNN	Single Layer PC/BC	LGN-V1 PC Model	Bayesian HMAX	Traditional Deep CNN's	PredNet control w/o error penalization	PredNet
	Reference	Rao & Ballard, 1999	Adelson & Bergen, 1985	McIntosh et al., 2016	Spratling, 2010	Jehee & Ballard, 2009	Dura-Bernal et al, 2012	AlexNet, VGG, ResNet,	This work	This work
	Length Suppression	Y	N	?	Y	Ν	?	N	N	Y
Exhibits Phenemena?	On/Off Dynamics	N	Y	~Y	?	Y	?	N	~	Y
	Sequence Learning Effects	N	N	?	N	N	N	N	~	Y
	Illusory Contours	?	N	N	?	N	~	N	N	Y
	Flash-Lag Illusion	N	N	N	N	N	N	N	Y	Y
	Weights are learned	Y	Not proposed*	Y	N	Y	Y	Y	Y	Y
Learning Aspects	Trained on natural stimuli	Y	N/A	Y	N/A	Y	N	Y	Y	Y
	Trained w/ dynamic stimuli	N	N/A	~	N/A	N	N	N	Y	Y
	Trained w/ biol. plausible loss	Y	N/A	N/A	N/A	Y	~	N	Y	Y

Key Y

Y Yes N No

~ Partially

? Not tested, but possible

Extended Data Fig. 7 | Comparison of the PredNet to prior models. The models under comparison are a (non-exhaustive) list of prior models that have been used to probe the phenomena explored here. The top section indicates if a given model (column) exhibits each phenomenon (row). The bottom section considers various learning aspects of the models. From left to right, the models considered correspond to the works of Rao and Ballard (1999)⁹, Adelson and Bergen (1985)⁷⁸, McIntosh et al. (2016)⁷⁹, Spratling (2010)⁴⁵, Jehee and Ballard (2009)⁴⁶, and Dura-Bernal et al. (2012)⁸⁰. Additionally, traditional deep CNNs are considered (for example AlexNet⁸¹, VGGNet⁸², ResNet⁸³). The PredNet control (second column from right) refers to the model in Extended Data Figures 9 and 10.

*As this is a general model family, theoretically the kernels can be learned from data, though not proposed in the referenced work.

- 78. Adelson, E. H. & Bergen, J. R. Spatiotemporal energy models for the perception of motion. J. Opt. Soc. Am. A 2, 284-299 (1985).
- 79. McIntosh, L., Maheswaranathan, N., Nayebi, A., Ganguli, S. & Baccus, S. Deep learning models of the retinal response to natural scenes. In *Advances in Neural Information Processing Systems* 1369–1377 (NeurIPS, 2016).
- 80. Dura-Bernal, S., Wennekers, T. & Denham, S. L. Top-down feedback in an HMAX-like cortical model of object perception based on hierarchical bayesian networks and belief propagation. *PLoS ONE* 7, 1–25 (2012).
- 81. Krizhevsky, A., Sutskever, I. & Hinton, G. E. ImageNet classification with deep convolutional neural networks. In Advances in Neural Information Processing Systems, 1097–1105 (NeurIPS, 2012).
- 82. Simonyan, K. & Zisserman, A. Very deep convolutional networks for large-scale image recognition. *International Conference on Learning Representations* (ICLR, 2015).
- 83. He, K., Zhang, X., Ren, S. & Sun, J. Deep residual learning for image recognition. In *The IEEE Conference on Computer Vision and Pattern Recognition* 770–778 (IEEE, 2016).

RPM	PredNet	BK (Subject 1)	CL (Subject 2)
10	6.7 ± 1.6	-	-
15	8.8 ± 2.1	9.9 ± 1.8	7.8 ± 2.9
25	11 ± 2.8	12.3 ± 3.2	11.3 ± 1.3
35	9.9 ± 8.0	24.6 ± 4.9	12.2 ± 2.4

Extended Data Fig. 8 | Comparison of PredNet predictions in the flash-lag illusion experiment to psychophysical estimates. The psychophysical estimates come from Nijhawan, 1994⁵⁸. With the frame rate of 10 Hz used to train the PredNet as a reference, the average angular difference between the inner and outer bars in the PredNet predictions was quantified for various rotation speeds. The results are compared to the perceptual estimates obtained using two human subjects by Nijhawan⁵⁸. Mean and standard deviation are shown. For rotation speeds up to and including 25 rotations per minute (RPM), the PredNet estimates align well with the psychophysical results. At 35 RPM, the PredNet predictions become noisy and inconsistent, as evidenced by the high standard deviation.



Extended Data Fig. 9 | PredNet control model lacking explicit penalization of activity in "error units". An additional convolutional block (\hat{A}_0^{frame}) is added that generates the next-frame prediction given input from R_0 . The predicted frame is used in direct L_1 loss, with the removal of the activity of the *E* units from the training loss altogether. Thus, in this control model, the *E* units are unconstrained and there is no explicit encouragement of activity minimization in the network.



Extended Data Fig. 10 | **Results of control model with the removal of explicit minimization of "error" activity in the PredNet.** Overall, the control model less faithfully reproduces the neural phenomena presented here. a) The control network E_1 units exhibit *enhanced* length suppression when feedback is removed (opposite of the effect in biology and the original PredNet). b) The responses in the control network still peak upon image onset and offset, however the decay in activity after peak is non-monotonic in several layers and less dramatic overall than the results shown in Fig. 3. As opposed to the 20-49% decrease in response after image onset peak in the original PredNet and the 44% decrease in the Hung et al.⁵³ macaque IT data, the control network exhibited a 5% (E_1) to 30% (E_2) decrease. c) Response of the control network E_3 layer in the sequence pairing experiment. The unpredicted images actually elicit a higher response than the first image in the sequence and the predicted images hardly elicit any response, both effects which are qualitatively different than the macaque IT data from Meyer and Olson⁵⁴ and the original PredNet. d,e) The average E_1 response in the control network demonstrates a decrease in activity upon presentation of the illusory contour.