Bipartite invariance in mouse primary visual cortex

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Introduction

The key challenge of visual perception is inferring behaviorally relevant latent features in the world despite drastic variations in the environment. For example, to recognize food in a cluttered environment, the brain must extract relevant features from light patterns in a way that is robust and generalizable to variations such as viewing distance, 3D pose, scale, and illumination, while still consistently identifying the food. Contrary to often being labeled as “nuisance”, these variations in the input must also be represented by the brain since they play a crucial role in other tasks such as navigating the environment.

To understand how brains effectively disentangle high-dimensional sensory inputs and robustly extract latent variables, it is essential to identify the features that neurons exhibit selectivity and invariance toward. Invariance refers to the property of neurons to maintain consistent responses despite changes in the input.

However, characterizing neuronal tuning is difficult because identifying invariances in the enormous search space of visual stimuli is challenging, experimental time is limited, and information processing in the brain is non-linear. Because of this, it is infeasible to present all images at all possible variations to systematically map invariances. As a result, most previous studies have been limited to using parametric stimuli (e.g., gratings) or semantic categories (e.g., objects and faces) (1–6) with strong assumptions about the types of invariances encoded. The classic example of this approach is Hubel and Wiesel’s complex cells in the primary visual cortex (V1) (7), which are tuned to oriented gratings of a preferred orientation but are invariant to spatial phases. In contrast, simple cells, are selective to spatial phase. However, we know little about other types of invariances along the visual hierarchy beyond these classes of parametric stimuli. This is critical for understanding how the brain robustly disentangles latent variables from a variable visual input (8).

To systematically study invariances, we combined in-vivo large-scale calcium imaging of the primary visual cortex in mice with deep convolutional neural networks (CNNs) to learn accurate neural predictive models. Using the CNN models as a “digital twin” of the visual cortex, we synthesized diverse exciting inputs (DEIs) in the functional connectomes MICrONs data set, which opens the way towards a circuit-level mechanistic understanding of this novel type of invariance. Our study demonstrates the power of using a data-driven deep learning approach to systematically characterize neuronal invariances. By applying this method across the visual hierarchy, cell types, and sensory modalities, we can decipher how latent variables are robustly extracted from natural scenes, leading to a deeper understanding of generalization.

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A defining characteristic of intelligent systems, whether natural or artificial, is the ability to generalize and infer behaviorally relevant latent causes from high-dimensional sensory input, despite significant variations in the environment. To understand how brains achieve generalization, it is crucial to identify the features to which neurons respond selectively and invariantly. However, the high-dimensional nature of visual inputs, the non-linearity of information processing in the brain, and limited experimental time make it challenging to systematically characterize neuronal tuning and invariances, especially for natural stimuli. Here, we extended “inception loops” — a paradigm that iterates between large-scale recordings, neural predictive models, and in silico experiments followed by in vivo verification — to systematically characterize single neuron invariances in the mouse primary visual cortex. Using the predictive model we synthesized diverse exciting inputs (DEIs), a set of inputs that differ substantially from each other while each driving a target neuron strongly, and verified these DEIs’ efficacy in vivo. We discovered a novel bipartite invariance: one portion of the receptive field encoded phase-invariant texture-like patterns, while the other portion encoded a fixed spatial pattern. Our analysis revealed that the division between the fixed and invariant portions of the receptive fields aligns with object boundaries defined by spatial frequency differences present in highly activating natural images. These findings suggest that bipartite invariance might play a role in segmentation by detecting texture-defined object boundaries, independent of the phase of the texture. We also replicated these bipartite DEIs in the functional connectomes MICrONs data set, which opens the way towards a circuit-level mechanistic understanding of this novel type of invariance. Our study demonstrates the power of using a data-driven deep learning approach to systematically characterize neuronal invariances. By applying this method across the visual hierarchy, cell types, and sensory modalities, we can decipher how latent variables are robustly extracted from natural scenes, leading to a deeper understanding of generalization.

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To understand how brains effectively disentangle high-dimensional sensory inputs and robustly extract latent variables, it is essential to identify the features that neurons exhibit selectivity and invariance toward. Invariance refers to the property of neurons to maintain consistent responses despite changes in the input. However, characterizing neuronal tuning is difficult because identifying invariances in the enormous search space of visual stimuli is challenging, experimental time is limited, and information processing in the brain is non-linear. Because of this, it is infeasible to present all images at all possible variations to systematically map invariances. As a result, most previous studies have been limited to using parametric stimuli (e.g., gratings) or semantic categories (e.g., objects and faces) (1–6) with strong assumptions about the types of invariances encoded. The classic example of this approach is Hubel and Wiesel’s complex cells in the primary visual cortex (V1) (7), which are tuned to oriented gratings of a preferred orientation but are invariant to spatial phases. In contrast, simple cells, are selective to spatial phase. However, we know little about other types of invariances along the visual hierarchy beyond these classes of parametric stimuli. This is critical for understanding how the brain robustly disentangles latent variables from a variable visual input (8).

To systematically study invariances, we combined in-vivo large-scale calcium imaging of the primary visual cortex in mice with deep convolutional neural networks (CNNs) to learn accurate neural predictive models. Using the CNN models as a “digital twin” of the visual cortex, we synthesized a set of stimuli that highly activated neurons while being distinct from each other — “Diverse Exciting Inputs” (DEIs). We subsequently presented these stimuli back to the animal while recording the activity of the same neurons and verified in vivo that these neurons responded invariantly to
the stimulus set. Our experiments discovered a novel type of invariance where one part of the receptive field (RF) was highly activated by specific textures like checkerboards translated over a wide range of phases, creating a local phase invariance. In contrast, the other part of the RF was selective for a fixed spatial pattern. We found that this bipartite structure of the RF often matched with object boundaries defined by a frequency difference in highly activating natural stimuli. Taken together, these findings suggest that bipartite invariance may play a role in supporting object boundary detection based on texture cues. In addition, we adapted our methodology to compute invariances using natural movies, which we then applied to the functional connectomics MICrONS data set (9). Our analysis successfully replicated bipartite DEIs in the MICrONS data set, providing a pathway toward understanding the circuit-level mechanisms underlying this novel form of invariance.

Results

Diverse Exciting Inputs (DEIs) identify invariant stimulus manifold. In this study, we used inception loops (10, 11), a closed-loop experimental paradigm, to investigate the invariances of single neurons in mouse V1 (Fig. 1b). An inception loop encompasses the following steps:

1. Neuronal recordings: Large-scale recordings with high-entropy natural stimuli.
2. Fit digital twin: Train a deep CNN (10, 12, 13) to predict the activity of neurons to arbitrary natural images. The CNN uses a non-linear core composed of convolutional layers shared among all recorded neurons, followed by a neuron-specific linear readout (Fig. 1d).
3. In silico experiments: Use the CNN predictive model as a functional “digital twin” of the brain to perform in silico experiments to systematically study the computations of the modeled neurons and derive experimentally testable predictions.

We presented 5,100 unique natural images to awake, head-fixed mice while recording the activity of thousands of V1 layer 2/3 (L2/3) excitatory neurons using two-photon calcium imaging (Fig. 1e). We used the recorded neuronal activity to train CNNs to predict the responses of these neurons to arbitrary images. The predictive model included a shifter network to compensate for eye movements and a modulator network for predicting an adaptive gain based on behavioral variables like running velocity and pupil size (10, 12) (Fig. 1d). When tested on a held-out set of novel natural images, the model achieved a median normalized correlation coefficient of 0.71 (14) (Fig. 1e), comparable to state-of-the-art models of mouse V1 (13, 15, 16).

We adapted a previously introduced method (17) to map the invariances of individual neurons in silico. The method extends optimal stimulus synthesis described recently (10, 11):

instead of finding the Most Exciting Input (MEI) for each neuron, it generates a set of optimal images — Diverse Exciting Inputs (DEIs). The DEIs are generated such that each of them highly excites the target neuron while being maximally different from all other DEIs in the set. For every model neuron, we initialized the DEI optimization with its MEI perturbed with different seeds of white noise and then re-optimized the image so that each of them drives the target neuron strongly while at the same time achieving the greatest pixel-wise Euclidean distance to the other DEIs of that neuron (see Methods for optimization loss). Our DEI synthesis method correctly characterized the invariances of simulated Hubel & Wiesel simple and complex cells. Specifically, all the DEIs of a simulated simple cell exhibited the same orientation, spatial frequency and phase of a Gabor patch (Fig. 2a simulated), as expected from linear-nonlinear (LN) models (18–20). For a simulated complex cell, the DEIs exhibited different phases, as expected from the phase invariance of complex cells (Fig. 2a simulated).

The DEIs of the modeled mouse V1 neurons strongly resembled their corresponding MEIs but with specific image variations that defined different types of invariances (Fig. 2a model mouse V1 neurons, more examples in Fig. S1). Some neurons had DEIs that were almost identical, indicating that the cells were not invariant, like the simulated simple cells. Among the neurons that were strongly activated by stimuli deviating from Gabor-shaped RFs (10, 13), some appeared to be stimulated strongly by random crops from an underlying texture pattern, demonstrating global phase invariance. We refer to these neurons as texture cells, similar to those found in the hidden layers of Artificial Neural Networks (ANNs) trained on object recognition (17). Many neurons exhibited a novel type of invariance: a bipartite RF invariance. For these neurons, one portion of their RF responded to a fixed spatial pattern while the rest was invariant to phase shifts of the stimuli, similar to texture cells. We denote these neurons as bipartite cells. We verified that this bipartite RF structure is unlikely the result of signal contamination from neighboring neurons since the MEI and DEIs optimized for different imaging slices in depth from the same neurons are highly consistent (Fig. S2). To quantify the prevalence of this phenomenon, we computed a diversity index for each neuron using its DEIs. This index was defined as the normalized average pairwise Euclidean distance in pixel space across the DEIs (see Methods). In particular, an index of 0 and 1 corresponds to classical simple and complex cells, respectively, while bipartite cells will have an index in between. We observed that the diversity indexes of mouse V1 neurons spanned a continuous spectrum, with those of simulated simple and complex cells on the two extremes (Fig. 2b).

We validated the model’s predictions by confirming the responses of the synthesized DEIs in vivo. Note that the mean luminance and root mean square (RMS) contrast of all the DEIs and MEIs of all neurons were matched (see Methods). When presented back to the animal, both MEIs and DEIs were highly selective, consistently eliciting higher activity in their target neurons than in most other neurons (Fig. 2c, S3).

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Fig. 1. Deep neural network well captures mouse V1 responses to natural scenes. a, Illustration of the optimization of Most Exciting Inputs (MEI) and Diverse Exciting Inputs (DEIs). The vertical axes represent the activation of a model neuron with no obvious invariance (left) and another model neuron with phase invariance to its optimal stimulus (right) as a function of two example image dimensions. The black curves depict optimization trajectories of the same MEI (left) starting from different initializations and of the DEIs (right) as different perturbations starting from the MEI along the invariance ridge. b, Schematic of the inception loop paradigm. On day 1, we presented sequences of natural images and recorded in vivo neuronal activity using two-photon calcium imaging. Overnight, we trained an ensemble of CNNs to reproduce the measured neuronal responses and synthesized artificial stimuli for each target neuron in silico. On day 2, we showed these stimuli back to the same neurons in vivo to compare the measured and predicted responses. c, We presented 5,100 unique natural images to an awake mouse for 500 ms, interleaved with gray screen gaps of random length between 300 and 500 ms. A subset of 100 images was repeated 10 times each to estimate the reliability of neuronal responses. Neuronal activity was recorded at 8 Hz in V1 L2/3 using a wide-field two-photon microscope. Behavioral traces including pupil dilation and locomotion velocity were also recorded. d, Schematic of the CNN model architecture. Our network consists of a 3-layer convolutional core followed by a single-point readout predicting neuronal responses, a shifter network accounting for eye movements, and a behavior modulator predicting an adaptive gain for each neuron (10, 12). Traces on the right show average responses (gray) to test images of two example neurons and corresponding model predictions (black). e, Normalized correlation coefficient (\(CC_{\text{norm}}\), see Methods) (14) between measured and predicted responses to test images for all 22,083 unique neurons across 10 mice selected for MEI and DEIs generation, a metric measuring the fraction of variation in neuronal responses to identical stimuli accounted for by the model prediction (median = 0.71 as indicated by the dashed line). Neurons with \(CC_{\text{max}}\) smaller than 0.1 (0.31%) were excluded and \(CC_{\text{norm}}\) larger than 1 (1.19%) were clipped to 1 in the histogram.

Moreover, the closed-loop experiments also confirmed that our CNN model accurately predicted the responses to all synthesized MEIs and DEIs (Fig. 2d, e). Importantly, DEIs strongly activated their target neurons in vivo, achieving 75% of their corresponding MEI activation (Fig. 2f), close to the model prediction of 85%. One potential concern is that the difference across DEIs may be due to features that are indistinguishable to the mouse visual system. Therefore, we tested
if the mouse visual system can detect the changes across DEIs by presenting pairs of the most dissimilar DEIs in pixel space to V1 neurons. Using a logistic regression classifier, we decoded the DEI class using the V1 population responses on a trial-by-trial basis and found a median classification accuracy of 0.9, significantly higher than the chance level (Fig. 2g). This result confirmed that the mouse visual system is able to accurately detect the differences between DEIs.

Next, we wanted to test whether DEIs follow specific invariance trajectories from the MEI that preserve high activity whereas random directions are sharply tuned. Similar to the DEI generation, we perturbed the MEI with different additive Gaussian white noise to generate a set of 20 control images with maximal diversity (Fig. 3a synthesized controls). In contrast to the DEI generation, in this case, the loss function did not require the images to be highly activating, yet enforced all the synthesized control images to be closer to the MEI than DEIs as measured by Euclidean distance in pixel space (see Methods). When shown back to the animal during inception loop experiments, these control images elicited significantly lower responses to their target neurons compared to the DEIs (Fig. 3b).

In silico: $y = 0.85x$

In vivo: $y = 0.75x$

Normalized MEI response (a.u.)

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Fig. 3. DEIs evoked stronger responses than synthesized and natural control stimuli in target neurons and generalized across different synthesis conditions. a. Examples of non-parametric DEIs (top), synthesized controls (middle), and natural controls (bottom) for 2 neurons. Synthesized controls were generated by perturbing the MEI in random directions while natural controls were found by searching through random natural patches. Both natural and synthesized controls were restricted to be strictly closer to the MEI in pixel space than all DEIs. b–c. Each point corresponds to the normalized activity of a single neuron in response to its DEIs versus its synthesized controls (b) or natural (c) controls. Response to each stimulus type was averaged over 20 different images with single-repeat. DEIs activated their target neurons stronger than their corresponding synthesized (one-sided Wilcoxon signed-rank test, $W = 3258, P < 10^{-7}$) and natural image controls (one-sided Wilcoxon signed-rank test, $W = 6442, P < 10^{-7}$) with 47.2% and 41.0% of all neurons showing greater response to DEIs respectively ($P < 0.05$, one-tailed Welch’s t-test with 30.4 and 31.1 average d.f., respectively). Data were pooled over 318 neurons from 5 mice. d. Examples of non-parametric DEIs synthesized with diversity evaluated as Euclidean distance in pixel space (top), or as Pearson correlation in in silico population responses (middle), or using a predictive model of different architecture and trained on different stimulus domain (bottom).

Neurons in vivo (Fig. 3d, S4). Second, we wanted to determine if DEIs are robust across predictive models with distinct architecture trained on different stimulus domains. To do this, we developed a method to synthesize DEIs from the dynamic digital twin trained on natural movies in the MICrONS functional connectomics data set (9, 22). Specifically, we presented to the same population a static set of natural images as well as the identical set of natural movies used in the MICrONS data set. We then trained two different static image models: one directly on in vivo static responses, and another on in silico predicted responses to the identical static images from a recurrent neural network trained on movie stimuli (22). We found that the MEIs and DEIs generated from these two models were perceptually similar (Fig. 3d) and both strongly activated their corresponding neurons as evaluated on another independent model (Fig. S5b, c). The ability to reproduce these invariances in the MICrONS data set provides an avenue for exploring the synaptic-level mechanisms underlying bipartite invariances.

Bipartite parametrization of DEIs Up until this point, we only discussed the types of invariances we observed for the texture and bipartite invariant cells in qualitative terms. Here we provide a more interpretable and quantitative model to account for the invariances we observed. We first used a texture synthesis model that could characterize the invariances of neurons similar to classical complex cells which maintain high activation when presented with phase-shifted versions of their preferred texture. To do this, we followed a previous method (17) to synthesize a full-field texture for each neuron by maximizing the in silico average activation of uniformly sampled crops from the texture (Fig. 4a, middle rows). We then sampled random crops from the optimized texture which we denote as full-texture DEIs (DEIs$_{full}$). However, for many neurons this method failed to produce DEIs that resembled the original non-parametric DEIs (Fig. 4b middle vs. top rows, 4c middle vs. top rows). Indeed, when shown back to the animal, the responses of many neurons (52.9%) were significantly lower when presented with DEIs$_{full}$ compared to the non-parametric DEIs (Fig. 4d). We hypothesized that the difference in responses was due to the fact that texture phase invariance alone could not account for the bipartite cells that have heterogeneous subfields in the RF (Fig. 2a) and are thus not well characterized by a homogeneous full-field texture model.

Thus, we introduced a new model to explain bipartite DEIs. We parameterized DEIs as the summation of two non-overlapping subfields within the RF: (1) a phase invariant subfield, which is cropped from a synthesized full-field texture and (2) a fixed subfield, which is masked from the original MEI and shared across all DEIs (Fig. 4a, b bottom rows). The invariant subfield was determined by finding the region of the RF with high pixel-wise variance across the non-parametric DEIs (see Methods). We then optimized a full-field texture, using a similar approach as the DEIs$_{full}$ method, but with a key difference: we used crops masked with the invariant subfield instead of the whole RF. Finally, we combined randomly sampled crops from the optimized texture and the fixed subfield to achieve partial-texture DEIs (DEIs$_{partial}$) (for more details see Methods). This model has the flexibility to represent a spectrum of phase invariance properties ranging from simple cells (or other non-invariant neurons) to complex cells (or other texture phase invariant neurons), as well as the bipartite cells we observed. Under this model, for simple cells, the fixed subfield occupies the
Fig. 4. Partial-texture DEIs activated target neurons similarly as non-parametric DEIs. a–b, Schematic of non-parametric (DEIs), full-texture (DEIs
full), and partial-texture (DEIs
partial) DEIs synthesis for an example texture cell (left) and an example bipartite cell (right). DEIs
full were synthesized by optimizing an underlying texture canvas from which uniformly sampled cropped by the MEI mask maximally activate the target neuron. In contrast, DEIs
partial are composed of two non-overlapping subfields: a fixed one masked directly from the MEI, and a phase invariant one synthesized similarly to DEIs
full except that the mask used for texture optimization is only part of MEI mask. c, MEI, DEIs, DEIs
full, and DEIs
partial of 4 example neurons, with each type of DEIs indicated by the corresponding color in a. DEIs
partial visually resembles non-parametric DEIs for most neurons while DEIs
partial captures non-parametric DEIs only for texture-like neurons. d, e, Normalized responses to DEIs
full (d) and DEIs
partial (e) responses versus non-parametric DEIs. Each point corresponds to the normalized activity of a single neuron, averaged over 20 different images with single repeat. d, DEIs
full failed to stimulate their target neurons compared to non-parametric DEIs (one-sided Wilcoxon signed-rank test, \( W = 4389, P < 10^{-4} \)) with 52.9% of all neurons showing lower responses to DEIs
full (\( P < 0.05 \), one-tailed Welch’s t-test with 29.4 average d.f.). e, DEIs
partial activated their target neurons similarly to non-parametric DEIs (two-sided Wilcoxon signed-rank test, \( W = 32429, P = 7.01 \times 10^{-4} \)) with only 8.7% of all neurons showing different responses (\( P < 0.05 \), two-tailed Welch’s t-test with 33.5 average d.f.). Data were pooled over 8 mice, displaying a total of 401 neurons.

We next tested the necessity and specificity of the two sub-fields in DEIs
partial (Fig. S7 a). Both subfields were necessary — the removal of either the phase invariant or the fixed sub-field resulted in lower activation (Fig. S7 b, c). When we swapped the phase invariant subfields with texture crops optimized for other random neurons or swapped the fixed sub-fields with random natural patches, these control stimuli also failed to strongly activate the neurons (Fig. S7 d, e). This demonstrates the specificity of the combination of the fixed and phase invariant subfields.

All of the closed-loop verification results shown up until now...
came from neurons with high invariance levels (see Methods). We also verified that all results presented in this section generalized to randomly selected neuronal populations (Fig. S8). Together, these findings suggest that most mouse V1 neurons, regardless of their level of invariance, were better explained by a model that parameterizes heterogeneous properties by partitioning the RF into two non-overlapping subfields, one of which contains phase invariance.

Invariant and fixed subfields detect boundaries in natural images with spatial frequency differences. It has been shown that the complex spatial features exhibited by MEIs are frequently present in natural scenes (10). Here, we wondered what visual tasks in the real world would benefit from this particular RF property of subfield division with phase invariance. Intuitively, these neurons may be involved in segmentation, such as detecting boundaries based on luminance changes (“first-order” cues) or texture variations (“second-order” cues) (23–25). Since our partial-texture model partitions the DEIs into two subfields, often with distinct patterns, we hypothesized that these neurons may be preferentially activated by object boundaries defined by texture discontinuities.

To test this hypothesis, we selected a natural image data set with manual segmentation labels, namely Caltech-UCSD Birds-200-2011 (CUB) (26), that enables us to locate the corresponding boundary between the object (a bird) and the background in each image. We screened more than one million crops with standardized RMS contrast from the CUB data set to identify highly activating crops for each V1 neuron in silico (Fig. 5a). We found that across the population, highly activating crops were significantly more likely to contain object boundaries than a random set of crops (Fig. 5b). To assess how well the segmentation masks defined by the object boundary in these highly activating natural images were aligned with the RF subfields extracted from DEIs_partial, we computed a matching score (Fig. 5a). For each image, the matching score would equal to 1 or -1 if the object perfectly matched with the phase invariant or the fixed subfield, respectively. Across the population, the phase invariant subfield was significantly aligned with the object rather than the background of the image (Fig. 5e, for more examples see Fig. S9a). In contrast, random image crops were not significantly aligned to the bipartite masks of the DEIs showing that the effect we observed was not due to biases in the images.

Next, we asked what low-level visual statistics could contribute to such matching results. We noticed that the DEIs appeared to exhibit differences in the spatial frequency preference between the fixed and phase invariant subfields. To investigate this, we replaced the DEI content within the two subfields with synthesized textures of different spatial frequency content. The textures were generated using Perlin noise (27)—a pseudo-random procedural texture synthesis method with naturalistic properties (Fig. S10a). Our findings revealed that a significantly larger proportion of neurons (41.08%) were most activated when the phase invariant subfield contained higher frequency content than the fixed part while very few neurons (8.67%) preferred the opposite site (Fig. S10b). Therefore, we hypothesized that object boundaries defined by differences in spatial frequency would strongly activate V1 neurons. To explicitly address this, we replaced the original contents in the CUB data set with gratings of varying spatial frequencies while still keeping the naturalistic boundaries (CUB-grating images) and presented them to our modeled neurons (Fig. 5d). Our results confirmed that neurons preferred images with object boundaries defined by the different spatial frequencies of the gratings (71.08%) significantly more than boundaries defined by grating orientation alone (28.58%) and homogeneous gratings with no boundaries (0.34%) (Fig. 5e). Just like in the natural image patches that strongly activated neurons (Fig. 5c), segmentation masks based on object boundaries in highly activating CUB-grating images also aligned significantly with the bipartite mask extracted from DEIs_partial (Fig. 5f). Notably, the phase invariant and fixed portions aligned with high and low spatial frequency, respectively, rather than with the object or background of the CUB-grating images (Fig. 5f). Overall, we found that mouse V1 neurons preferred object boundaries constructed by frequency changes, with phase invariances biased toward the higher frequency subfields.

Discussion

Invariant object recognition, the ability to discriminate objects from each other despite tremendous variation in their appearances, is a hallmark of visual perception. From the perspective of the object manifold disentanglement theory, visual input is gradually transformed and re-represented by visual neurons along the ventral visual pathway and ultimately achieves linear decoding of object identity from the population representation at the top of the hierarchy (8). Experimental and theoretical evidence suggested that single neurons in higher visual areas extract and integrate information from simple feature detectors in lower areas to represent more complex features and also build up response invariance to feature transformations (28–31). Discovering neuronal invariances is critical to understand how the brain generalizes to novel stimuli, and helps define the latent variables that animals use to guide actions. However, single-cell invariance properties have not been characterized systematically despite a few classical examples discovered using parametric or semantically meaningful stimuli that are largely human-biased (6, 7, 32). A few recent studies have developed non-parametric image synthesis approaches to find preferred stimuli of visual neurons (10, 11, 33). Yet most efforts along this direction characterized purely neuronal selectivity instead of invariance, and also mostly focused on feature visualization without further interpreting the functional roles of neurons.

In this study, we extended the previous work of combining large-scale neuronal recording and deep neural networks to study neuronal selectivity (10) to the invariance problem. Particularly, we modified a diverse feature visualization approach previously developed in ANNs (17) to synthesize DEIs for individual neurons in mouse V1 layer 2/3 and verified these model-predicted near-optimal diverse im-
ages in vivo. Given that the changes across DEIs were decodable at the V1 population level and that DEIs preserved target neuron activation better than other diverse image sets that were strictly more similar to the MEI in pixel space, our DEIs successfully captured non-trivial and highly specific invariant tuning directions. Note that revealing such specific activation-preserving image perturbations is nearly impossible without the invention loop paradigm because of the extremely high-dimensional stimulus search space and the high sparsity of mouse visual neurons’ responses (10, 34). Importantly, we demonstrated that our DEI methodology yielded similar results across variations in diversity metric, stimulus domain, and model bias, so this paradigm should be adaptable to characterize neuronal invariances systematically across different cortical regions of interest, sensory modalities, and even animal species.

While the current study focused on invariances around the optimal stimulus, one could easily generalize our method to study the entire tuning landscape by varying the desired neural response. Ultimately, our work is a first attempt to characterize single-neuron invariances in biological systems in a systematic and unbiased fashion, paving the way for more holistic approaches that combine invariance and selectivity to understand the neuronal tuning landscape in the future.

Mouse V1 DEIs revealed a novel type of bipartite phase invariance that could be parsimoniously parameterized by a partial-texture generative model: one part prefers a fixed spatial pattern while the other prefers a random windowed crop from an underlying texture. These parametric DEIs$_{\text{partial}}$ are not only strikingly indistinguishable from the non-parametric version but also excited their target neurons similarly when shown back to the mice. In contrast, when homogeneous phase change throughout the entire RF was assumed, the optimized DEIs$_{\text{full}}$ failed to stimulate target neurons in vivo. Overall, the essence of such partial-texture parameterization is the introduction of heterogeneity in both selectivity and invariance between two subfields of the RF. This is a property that fundamentally deviates from the homogeneous represen-
tation assumed in traditional simple and complex cells. While we have focused primarily on phase invariance, we do not attempt to claim it is the only type of invariance existing in mouse V1. As an initial effort to parameterize novel empirical invariances, it is also worth acknowledging that our partial-texture model proposes a simple hypothesis of a binary division of the presence and absence of phase invariance in the RF without considering more complicated scenarios of multi-region invariances or nonlinear cross-subfield interactions. Nonetheless, we believe the novel bipartite phase invariance can be of great use as a computational principle for future designs of biologically-plausible or brain-inspired computer vision systems (35). On the other hand, it can also serve as an empirical test for theoretically-driven (36–38) or data-driven models (39) that aim to explain and predict neural responses in the visual system.

The observed heterogeneity between the two RF subfields corresponded not only in phase invariance but also in preferred spatial frequency. This property is remarkably similar to that observed in “high-low frequency detectors”, a term introduced previously to describe single units in ANNs that look for low-frequency patterns on one side of their RF and high-frequency ones on the other side (40). As these ANN units were speculated to encode natural boundaries defined by frequency variation, we indeed found supporting evidence that V1 layer 2/3 neurons preferred natural image patches containing object boundaries. This bipartite phase invariance with frequency bias might be common in both biological and artificial vision systems for boundary detection. Importantly, our results are consistent with previous findings showing that mice are behaviorally capable of utilizing texture-based cues to solve segmentation (25, 41). While these studies particularly focused on object boundaries constructed mostly by orientation or phase difference (41), our evidence suggests spatial frequency bias as yet another type of second-order (texture variation-based) visual cue (23) important for detecting object boundaries.

To investigate the neuronal wiring that allows the formation of bipartite invariance, one can take advantage of recent advances in functional connectomics, combining large-scale neuronal recordings with detailed anatomical information at the scale of single synapses. Here, we demonstrated that the DEIs of mouse V1 neurons were reproduced (Fig. S5e) in a recently published MICrONS functional connectomics data set spanning V1 and multiple higher areas of mouse visual cortex (9). This data set includes responses of >75K neurons to natural movies and the reconstructed sub-cellular connectivity of the same cells from electron microscopy data. The MICrONS data set provides unprecedentedly ample resources to link morphology and connectivity among neurons to examine the circuit-level mechanisms that give rise to bipartite phase invariance.

Methods

**Neurophysiological experiments** The following procedures were approved by the Institutional Animal Care and Use Committee of Baylor College of Medicine. Ten mice (*Mus musculus*: 6 male, 4 female) aged from 6 to 17 weeks, expressing GCaMP6s in excitatory neurons via Scl17a7-Cre and Ai162 transgenic lines (stock nos. 023527 and 031562, respectively; The Jackson Laboratory) were selected for experiments. The mice were anesthetized and a 4 mm craniotomy was made over the visual cortex of the right hemisphere as described previously (34, 42). For functional imaging, mice were head-mounted above a cylindrical treadmill and calcium imaging was performed using a Chameleon Ti-Sapphire laser (Coherent) tuned to 920 nm and a large field-of-view microscope equipped with a custom objective (0.6 numerical aperture, 21 mm focal length) (43). Laser power at the cortical surface was kept below 58 mW and maximum laser output of 61 mW was used at 245 ± 50 nm from the surface.

We also recorded the rostro-caudal treadmill movement as well as the pupil dilation and movement. The treadmill movement was measured via a rotary optical encoder with a resolution of 8,000 pulses per revolution and was recorded at approximately 100 Hz in order to extract locomotion velocity. The images of the left eye were reflected through a hot mirror and captured with a GigE CMOS camera (Genie Nano C1920M; Teledyne Dalsa) at 20 fps with a resolution of 246-384 pixels × 299-488 pixels.

A DeepLabCut model (44) was trained on 17 manually labeled samples from 11 animals to label each frame of the compressed eye video with 8 eyelid points and 8 pupil points at cardinal and intercardinal positions. Pupil points with likelihood >0.9 (all 8 in 93% ± 8% of frames) were fit with the smallest enclosing circle, and the radius and center of this circle was extracted. Frames with <3 pupil points with likelihood >0.9 (0.7% ± 3% frames per scan), or producing a circle fit with outlier >5.5 standard deviations from the mean in any of the three parameters (center x, center y, radius, <1.3% frames per scan) were discarded (total <3% frames per scan).

We delineated visual areas by manually annotating the retinotopic map generated by pixel-wise response to a drifting bar stimulus across a 4,000 × 3,600 µm² region of interest (0.2 px/µm⁻¹) at 200 µm depth from the cortical surface. The imaging site in V1 was chosen to minimize blood vessel occlusion and maximize stability. Imaging was performed using a remote objective to sequentially collect ten 630 × 630 µm² fields per frame at 0.4 px/µm⁻¹ xy resolution at approximately 8 Hz for all scans. We allowed only 5 µm spacing across depths to achieve dense imaging coverage of a 630 × 630 × 45 µm³ xyz volume. The most superficial circle positioned in L2/3 was around 200 µm from the surface of the cortex. Thanks to our dense sampling, cells in the imaged volume were heavily over-sampled, often appearing in at least 2 or more imaging planes. This allowed matching across days with 2.5 ± 2.6 µm vertical distance between masks (see details below). We performed raster and motion correction on the imaging data and then deployed CNMF algorithm (45) implemented by the CalMAn pipeline (46) to segment and deconvolve the raw fluorescence traces. Additionally, cells were selected by a classifier (46) trained to detect somata based on the segmented cell masks to result in 6,014–7,987 soma masks per scan. The full two-photon imaging process-
Visual stimuli were presented 15 cm away from the left eye with a 25° LCD monitor (31.8 x 56.5 cm, ASUS PB225Q) at a resolution 1080 x 1920 pixels and refresh rate of 60 Hz. We positioned the monitor so that it was centered on and perpendicular to the surface of the eye at the closest point, corresponding to a visual angle of 2.2°/cm on the monitor. In order to estimate the monitor luminance, we taped a photodiode at the top left corner of the monitor and recorded its voltage during stimulus presentation, which is approximately linearly correlated with the monitor luminance. The conversion between photodiode voltage and luminance was estimated from luminance measurements from a luminance meter (LS-100 Konica Minolta) for 16 equidistant pixel values ranging from 0 to 255 while simultaneously recording photodiode voltage. Since the relationship between photodiode voltage and luminance is usually stable, we only perform such measurement every a few months. In the beginning of every experimental session, we computed the gamma between pixel intensity and photodiode voltage by measuring photodiode voltage at 52 equidistant pixel values ranging from 0 to 255; then we further interpolated the corresponding luminance at each pixel intensity. For closed loop experiments, the pixel-luminance interpolation computed on day 1 was used throughout the loop. All stimuli used in the current study were presented at gamma value ranging from 1.60 to 1.74 and monitor luminance ranging from 0.07 ± 0.16 to 9.58 ± 0.65.

Monitor positioning across days In order to optimize the monitor position for centered visual cortex stimulation, we mapped the aggregate receptive field of the scan field region of interest (ROI) using a dot stimulus with a bright background (maximum pixel intensity) and a single dark dot (minimum pixel intensity). We tiled the center of the screen in a 10 x 10 grid with single dots in random locations, with 10 repetitions of 200 ms presentation at each location. The RF was then estimated by averaging the calcium trace of an approximately 15 x 15 µm² window in the ROI from 0.5 - 1.5 s after stimulus onset across all repetitions of the stimulus for each location. The resulted two-dimensional map was fitted with an elliptic two-dimensional Gaussian to find a center. To keep a consistent monitor placement across all imaging sessions, we positioned the monitor such that the aggregate RF of ROI in the first session was placed at the center of the monitor and then fixed the monitor position across the subsequent sessions within a closed loop experiment. An L-bracket on a six dimensional arm was fitted to the corner of the monitor at its location in the first session and locked in position such that the monitor could be returned to the same position between scans and across imaging sessions.

Cell matching across days In order to return to the same image site, the craniotomy window was leveled with regard to the objective with six d.f., five of which were locked between days. A structural 3D stack encompassing the volume was imaged at 0.8 x 0.8 x 1 µm³ resolution with 100 repeats. The stack contained two volumes each with 150 fields spanning from 50 µm above the most superficial scanning field to 50 µm below the deepest scanning field; each field was 500 x 800µm², together tiling a 800 x 800µm² field of view (300µm overlapped). This was used to register the scan average image into a shared xyz frame of reference between scans across days. To match cells across imaging scans, each two-dimensional scanning plane was registered to the three-dimensional stack through an affine transformation (with nine d.f.) to maximize the correlation between the average recorded plane and the extracted plan from the stack. Based on its estimated coordinates in the stack, each cell was matched to its closest cell across scans. To further evaluate the functional stability of neurons across scans, we included in every visual stimulus an identical set of 100 natural images with each repeated 10 times (referred as oracle images). For each pair of matched neurons from two different scans, we compute the correlation between their average-trial responses to the 100 oracle images. In order to be included for downstream analyses, the matched cell pair had to (1) have an inter-cellular distance smaller than 10µm; (2) achieve a functional correlation equal to or greater than the top 1 percentile of correlation distribution between all unmatched cell pairs (0.42); (3) survive manual curation of the matched pair’s physical appearance in the processed two-photon imaging average frame. On average, 58 ± 13% of closed-loop neurons survived all three criteria.

Presentation of natural stimuli To fit neurons’ responses, 5,100 natural images from ImageNet (ILSVRC2012) were cropped to fit a 16:9 monitor aspect ratio and converted to gray scale. To collect data for training a predictive model of the brain, we showed 5,000 unique images as well as 100 additional images repeated 10 times each. This set of 100 images were shown in every scan for evaluating cell response reliability within and between scans. Each image was presented on the monitor for 500 ms followed by a blank screen lasting between 300 and 500 ms, sampled uniformly.

Preprocessing of neural responses and behavioral data Neuronal responses were deconvolved using constrained nonnegative calcium deconvolution and then accumulated between 50 and 550ms after stimulus onset of each trial using a Hamming window. The corresponding pupil movement and treadmill velocity for each trial were also extracted and integrated using the same Hamming window. Each data set consists of 4500 and 500 unique images for training and validation, respectively; an additional set of 100 images presented with 10 repeats were used for model testing. The original stimuli presented to the animals were isotropically downsampled to 64 x 36 px for model training. Input images, neuronal responses, and behavioral traces were normalized (z-scored for input images and divided by standard deviation for the
rest) across either the training set or all data during model training.

**Predictive model architecture and training** We followed the same network architecture and training procedure as described previously (10, 12). The network is composed of four components: a common nonlinear core for all neurons, a linear readout dedicated for each neuron, a modulator network that scales the responses based on the animal’s pupil dilation and running velocity, and a shifter network predicting RF shifts from pupil position changes. The common core is a 3-layer CNN with full skip connections. Each layer contains a convolutional layer with no bias, followed by batch normalization, and an exponential linear unit (ELU) nonlinearity. The readout models the neural response as an affine function of the core outputs followed by ELU nonlinearity and an offset of 1 to guarantee positiveness. Additionally, we model the location of a neuron’s RF with a spatial transformer layer reading from a single grid point that extracts the feature vector from the same location at different scales of the downsampled feature outputs. The modulator computes a gain factor for each neuron that simply scales the output of the readout layer using a two-layer fully connected multiplayer perceptron with rectified linear unit (RELU) nonlinearity and a shifted exponential nonlinearity to ensure positive outputs. Finally, because training mice to fixate their gaze is impractical, we estimated the trial-by-trial RF displacement shared across all neurons using a shifter network composed of a three-layer MLP with a tanh nonlinearity. For model training, we followed the procedure detailed in (10). Four instances of the same network with different initialization were trained by minimizing the Poisson loss \[ \frac{1}{m} \sum_{i=1}^{m} (\hat{r}(i) - r(i) \log \hat{r}(i)) \]
where \( m \) denotes the number of neurons, \( \hat{r} \) the predicted neuronal response and \( r \) the observed response. We used all segmented neuronal masks for training, including the duplicate ones due to dense imaging. Predictions from these four models are then averaged at inference.

**Evaluation of neuronal reliability and model performance** To study how neurons encode visual stimuli using the predictive model, we restricted all analyses to neurons that exhibit reasonable level of response reliability as well as model predictive accuracy. We evaluated reliability using an oracle score for each neuron by correlating its leave-one-out mean response with that to the remaining trial across 100 images with each repeated 10 times in the test set. The model performance \( CC_{abs} \) for each neuron was computed on the test set as the correlation between the model predicted response \( \hat{\tau} \) and the recorded responses \( \tau \) averaged across 10 repetitions:

\[
CC_{abs} = \frac{Cov(\tau, \hat{\tau})}{\sqrt{Var(\tau)Var(\hat{\tau})}}.
\]

We hard thresholded on both of the evaluation metrics to select neurons for synthetic stimuli generation and downstream analyses (see further on). We also measured the fraction of variation in neuronal responses to identical stimuli accounted for by the model prediction as the normalized correlation coefficient (\( CC_{norm} \)) (14) estimated as

\[
CC_{norm} = \frac{CC_{abs}}{CC_{max}}.
\]

\( CC_{max} \) serves as the upper bound of the correlation coefficient and is computed as

\[
CC_{max} = \sqrt{\frac{NVar(y)}{(N-1)V ar(\tau)}},
\]

where \( y \) is the in vivo response, and \( N \) is the number of trials. We threshold neurons on \( CC_{max} \) to be larger than 0.1 (0.31% of neurons) to avoid unreliable estimate of \( CC_{norm} \). Then we clipped \( CC_{norm} \) values outside of 0 and 1 (1.19% of neurons).

**Neuron selection for synthetic stimuli generation** First excluded neuronal masks within 10 \( \mu m \) from the edge of the imaging volume, and then rank the remaining masks based on descending model predictive accuracy. To avoid duplicated neurons, we started from the lowest-ranked neuron and iteratively added neurons such that they are at least 25 \( \mu m \) apart and have functional correlation <0.4 with all neurons selected. Such filtering left us with 2,081–2,676 unique neurons for each of the ten mice. We then hard thresholded on oracle score and model test correlation to include approximately the top 22% of the population for mouse 1-2 for synthetic stimuli generation; we relaxed the thresholding to include approximately 79% of the population for mouse 3-10 and randomly sampled one-thirds of all unique neurons for synthetic stimuli generation.

**Generation of Most Exciting Input (MEI)** For each individual neuron, we adapted the activation maximization procedure described earlier (10) to find the stimulus that optimally drive each individual neuron. We started from a Gaussian white noise and iteratively added the gradient of the target neuron’s predicted response to the image. At every optimization step, we smoothed the gradient using a Gaussian filter with \( s.d = 1.0 \) to avoid high frequency artifact; after each gradient ascent step, we standardized the resultant image such that the mean and root mean square (RMS) contrast within the receptive field (estimated as the MEI mask, see details below) approximately match the corresponding values of the training set.

**Generation of MEI mask** We computed a weighted mask for each MEI to capture the region containing most of the variance in the image so that the resulting masked MEI activates the in silico neuron slightly less than the original unmasked MEI. We computed a pixel-wise z-score on the MEI and thresholded at z-score above 1.5 to identify the highly contributing pixels. Then we closed small holes/gaps using binary closing, searched for the largest connected region to create a binary mask \( M \) where \( M = 1 \) if the pixel is in the largest region identified. Then, a convex hull was calculated using the identified pixels. Lastly, to avoid edge artifacts, we smoothed out the mask using a Gaussian filter with \( \sigma = 1.5 \) to avoid potential edge effects.
**In silico image presentation**  In order to obtain the most trustworthy prediction from the predictive model, we would like to present all images in *silico* at statistics comparable to the training set images. On average, the training set images have approximately mean 0 and standard deviation 0.8 in the z-score space inside the MEI masks. To match the training set statistics, we synthesized MEIs with a range of full-field constraints and found the best match at approximately full-field mean 0 and RMS contrast 0.25. With this, we standardized all images before presenting them *in silico* to the predictive model following the same procedure: masking the image with the corresponding MEI mask and then normalizing the entire image to mean 0 and RMS contrast 0.25.

**Generation of Diverse Exciting Inputs (DEIs) and diversity evaluation** We modified procedures described previously (17) to optimize DEIs. For each individual neuron, we synthesized a set of images initiating from MEI that preserve high activation while differ as much as possible from each other. To optimize this set, we initiated from 20 instances of the target neuron’s MEI with different additive Gaussian white noises \( MEI + \sigma = I_i \) where \( 1 \leq i \leq 20 \) and iteratively minimize the loss:

\[
L = \frac{1}{n} \sum_{i=1}^{n} \max (c - \frac{r_i}{r_{MEI}} (0)) - \lambda \min_{i,j} d(I_i, I_j)
\]

where \( r_i \) and \( r_{MEI} \) are the model predicted response to DEI \( i \) and MEI, \( c \) is the minimum activation relative to \( r_{MEI} \) that we target each DEIs for, \( d(I_i, I_j) \) is the distance between DEI \( i \) and DEI \( j \) measured either in pixel Euclidean space or population representation space (see further on). The first term encourages all DEIs to achieve high activation, while the second term maximizes the minimum pair-wise distance among DEIs. Note that the minimum, instead of average distance, is used in the second term to avoid solutions that form the set of DEIs into clusters by pushing apart the most similar pair of DEIs at every iteration. Additionally, we employed two strategies to attenuate the prevalence of high-frequency artifact during CNN image synthesis - (1) we blurred the gradient for every image in the set at every step using a Gaussian filter; (2) we decayed the gradient for all images as a function of number of iterations. We performed the optimization for every target neuron with a series of diversity regularization hyper-parameter \( \lambda \), densely sampled from \( 1\times10^{-4} \) to \( 5\times10^{-2} \). For each neuron, the set optimized using the largest \( \lambda \) that preserved a minimal response greater than 85% of the MEI response was selected as the DEIs and used for downstream analyses.

**Selection of natural control and generation of synthesized control** To show that our predictive model indeed did a good job finding stimuli with both high activation and high diversity, we demonstrated that DEIs preserve activation better than control stimuli resulted from perturbation from the MEI in random directions. These stimuli need to be at comparable distance away from the MEI as the DEIs; to be extra conservative, we required each control stimulus to be strictly closer the MEI than all of the DEIs. We designed two types of such control stimuli through either natural image search or direct image synthesis. For each target neuron, we first computed the minimum distance from the DEIs to the MEI within the MEI mask as \( d_{target} \) and use it as the distance budget for searching for or synthesizing control images. For natural control, we augmented a pool of 124,647 ImageNet images of 256x144 px to 41,881,392 image patches of 64x36 px and searched for patches that have distance from the MEI within the MEI mask smaller than \( d_{target} \). Since it would be pointless to include controls too close to the MEI (i.e. no perturbation), we additionally excluded images that have distance to MEI smaller 80% of \( d_{target} \). For the synthesized controls, we synthesized a diverse set of random perturbations from MEI following our DEIs synthesis objective except that the first term is changed to match a targeted distance from the MEI rather than a desired *in-silico* activation as follows:

\[
L = \frac{1}{n} \sum_{i=1}^{n} (c - d(I_i, MEI)) - \lambda \min_{i,j} d(I_i, I_j)
\]

For both types of controls, we created 20 different images for each neuron and presented them *in vivo* with single repetition each.

**Diversity index** To calculate neurons’ diversity indexes on a meaningful spectrum with interpretable reference points, we estimated theoretical references for diversity indexes of idealized simple and complex cells. Particularly, we estimated the lower/upper bounds \( d_{lower} \) and \( d_{upper} \) as the median average pairwise Euclidean distance of DEIs from a population of noiseless simple/complex cells, respectively. To get a precise estimation, we identified DEIs via performing an exhaustive search through the Gabor filter parameter spaces. Since DEIs are standardized with a fixed mean and standard deviation, idealized simple cells have the same average pairwise Euclidean distance regardless of the underlying Gabor parameters. Similarly, idealized complex cells with different Gabor parameters have identical but higher average pairwise Euclidean distance among themselves. Then for each real neuron, a diversity index \( D \) is calculated for each mouse V1 neuron \( i \) based on the average Euclidean pairwise distance of its DEIs \( d_i \) as

\[
D(i) = \frac{d(i) - d_{lower}}{d_{upper} - d_{lower}}
\]

**Neuron selection for closed-loop verification** For 9 out of 10 mice, we randomly selected neurons with relatively high level of invariance (detailed below); for 1 mouse, we randomly selected neurons from all candidates that survived our oracle score and model performance criteria (see above). To remove the confounding effect of RF size on neurons’ invariance level, we fit a least squares regression from the MEI mask size to the diversity index computed from DEIs (see above) using 1000 random neurons compiled across 8 pilot data sets. For each neuron, the residual between the actual average DEIs pair-wise Euclidean distance and the predicted distance from the MEI mask size was calculated as its diversity residual. This diversity residual served as an size-independent evaluation of neuron’s invariance level. For each of the 9 mice, we randomly selected 80 neurons from the top
50 percentile among all neurons with positive diversity residuals.

**Presentation of synthetic stimuli** For all MEIs, DEIs, DEIs controls, and texture DEIs (see further on), we masked the stimuli with the MEI mask and standardized all masked stimuli such that they have fixed value of mean and RMS contrast in the luminance space within the MEI mask with very small amount of variation due to clipping within the 8-bit range. The fixed mean and contrast valued were chosen to be close to the corresponding values of the training set while minimizing the amount of clipping. All pixels outside of the MEI mask are at 128, the same intensity as the blank screen in between consecutive trials. For each neuron, its MEI was presented for 20 repeats and each of its 20 DEIs or 20 DEIs controls were presented once.

**Decoding DEIs from population responses** To confirm whether there exists perceivable difference across DEIs, for each target neuron, we selected the most different pair of DEIs (based on the corresponding diversity measurement space) from 20 synthesized DEIs. We presented each DEI (DEI1 and DEI2) for 20 repeats per neuron for the whole V1 population. We performed a 5-fold cross-validation logistic classification with L2 regularization on the population responses to classify whether each single trial response originated from DEI1 or DEI2. The regularization strength was optimized empirically from fitting logistic regression to decode DEIs across 1 pilot data set.

**Simple and complex cell models** The response of an idealized simple cell was modeled as convolution with a 2D Gabor filter followed by a rectified linear activation function (ReLU). An idealized complex cell was formulated by the classical energy model, where the response was modeled as the square root of the summation of squared outputs to a quadrature pair of 2D Gabor filters. In order to obtain a noisy predictive model for these idealized simulated neurons, we followed the same protocol of data presentation and model training for these idealized neurons as that for the real brain neurons. Specifically, we collected the idealized responses to 5000 random ImageNet images and used each response as the mean of a Poisson distribution from which a noisy response was sampled. Then we used the noisy input-response data set to train a predictive model with an identical architecture described above on the resultant simple and complex cell predictive models.

**Generation of DEIs in population representational space** To test whether DEIs could generalize across diversity representational spaces, we optimized DEIs using the same loss function as in 5 but now measuring pair-wise DEI diversity $d(DEI_i, DEI_j)$ as the negative Pearson correlation between model predicted population neuronal response vectors $r_i$ and $r_j$ to DEI$_i$ and DEI$_j$:

$$d(\text{DEI}_i, \text{DEI}_j) = -\frac{\sum (r_i - \mu_i)(r_j - \mu_j)}{\sqrt{\sum (r_i - \mu_i)^2 \sum (r_j - \mu_j)^2}}$$  \hspace{1cm} (4)

where $\mu_i, \mu_j$ are the mean population responses. We obtained the population responses by shifting all neurons’ RF center to that of the target neuron that the DEIs were optimized for. Therefore, each population response vector served as a proxy of the "perceptual representation" of the corresponding image. We denote these DEIs as population-based DEIs.

**Replication of DEIs in functional connectomics data set** Recently, a large-scale functional connectomics data set of mouse visual cortex ("MICrONS data set"), including responses of >75k neurons to full-field natural movies and the reconstructed sub-cellular connectivity of the same cells from electron microscopy data (9). A dynamic recurrent neural network (RNN) model (cite eric) of this mouse visual cortex—digital twin—exhibits not only a high predictive performance for natural movies, but also accurate out-of-domain performance on other stimulus classes such as drifting Gabor filters, directional pink noise, and random dot kinematograms. Here, we took advantage of the model’s ability to generalize to other visual stimulus domains and presented MEIs and DEIs to this digital twin model in order to relate specific functional properties to the neurons’ connectivity and anatomical properties. Specifically, we recorded the visual activity of the same neuronal population to static natural images as well as to the identical natural movies that were used in the MICrONS data set. Neurons were matched anatomically as described for the closed loop experiments. Based on the responses to static natural images we trained a static model as described above, and from the responses to natural movies we trained a dynamic model using a RNN architecture described in REF. We then presented the same static natural image set that we showed to the mice also to their dynamic model counterparts and trained a second static model using these predicted in silico responses. This enabled us to compare the MEIs and DEIs for the same neurons generated from two different static models: one trained directly on responses from real neurons, and another trained on synthetic responses to static images from dynamic models (D-MEI and D-DEIs). To quantify similarity, we presented both versions of MEIs and DEIs to an independent static model trained on the same natural images and responses but initialized with a different random seed, thereby avoiding model-specific biases.

**Generation of full-texture and partial-texture DEIs** A neuron with perfect partial phase invariance should maintain high activation to phase-shifted version of its preferred texture pattern within the spatial region that tolerates phase changes. Thus, we proposed a texture generative model to produce texture-based DEIs composed of two complementary sub-fields as follows:

$$\text{DEIs} = m_1 \cdot T + (1 - m_1) \cdot \text{MEI}$$  \hspace{1cm} (5)

where the first term is the phase invariant subfield randomly cropped from an optimized texture canvas $T$ using a mask $m_1$. The second term is a fixed subfield masked directly from the original MEI. This model could be simplified to a full-texture model to describe global phase invariance if the
entire receptive field (i.e. the MEI mask \( m_{MEI} \)) was used as \( m_I \). We generated the texture \( T \) following (17) by maximizing the average activation of randomly sampled crops from \( T \) using \( m_I \). We followed the same loss as in non-parametric DEIs generation 1 to jointly maximize the activation and diversity of DEIs with the same regularizations (i.e. Gaussian blurring on the gradient and learning rate decay) but in this case, DEIs were parameterized as in 5. The parametric DEIs were standardized to the same mean and RMS contrast as the MEI after each gradient step.

To ensure that \( m_I \) captures the region of the non-parametric DEIs from which we observed the most diversity, we pre-computed a series of \( m_I \)'s with increasing sizes by varying the threshold on the pixel-wise variance across the DEIs. Specifically, starting from the pixels with the largest variance across DEIs, we kept expanding the \( m_I \) by requesting increasing fraction of the total variance from 0.2 to 0.6 within the phase invariant subfield. In general, the average predicted activation of the texture-based DEIs decreased as the size of \( m_I \) increased. While an ideal complex cell should exhibit an almost flat line, an ideal simple cell should exhibit rapid activation decay. Most V1 neurons behaved as if in between the classical cell types, demonstrating an initial slow decrease in activation before steeper decay when the proposed phase invariant region expanded beyond the desirable target (Fig. S6a).

For each set of texture-based DEIs resultant from a phase invariant mask \( M_{I,j} \), we computed the harmonic mean between normalized activation and diversity index.

\[
H_j = \frac{2 \tau_j d_{DEIs,j}}{\tau_j + d_{DEIs,j}}
\]

where \( \tau_j \) is the average normalized activation, and \( d_{DEIs,j} \) is the average pair-wise Euclidean distance normalized by the maximal value across all different sets of stimulus corresponding to the series of \( M_{I,j} \). We denoted the set of texture-based DEIs resultant with the maximum score as the partial-texture DEIs. On the other hand, the set of texture DEIs resultant from the full-texture model were denoted as full-texture DEIs.

Necessity and specificity of the two subfields in partial-texture DEIs To test whether both the phase invariant and the fixed subfields are necessary for the high activation of the partial texture DEIs, we presented the content within each subfield alone and see if the neuronal activation decreased compared to the MEI activation. To ensure pixels within each subfield stay at their entirety, we computed a binary mask for each subfield and smoothed the edge of the mask only outside of the binary mask. To test the specificity of the content within each subfield, we swapped the subfield content to random patterns and see if the activation is lower than that of the original partial-texture DEIs. Particularly, we swapped each neuron’s phase invariant content to samples cropped from textures optimized for different random neurons and each neuron’s fixed content to random natural image crops. When presented to the animals, these content-swapped stimuli were standardized to match the mean and RMS contrast in the luminescence space as the original partial texture DEIs.

The CUB and CUB-grating data sets To study the relationship between invariance and natural stimuli, we created a data set from the Caltech-UCSD Birds-200-2011 (CUB) image data set (26). The original data set contains 11,788 natural images belonging to 200 bird categories. Each image contains a single bird appearing in its natural habitat resulting in ideal and ethnologically relevant stimuli for mice. Each original image is then resized to \( 64 \times 64 \) pixels and then sampled using a window of size \( 36 \times 36 \) pixels with a stride of size 2. In addition, each image has a semantic segmentation label identifying the region of the image containing the bird by hand. This segmentation label is then thresholded at 0.5 to get the final binarized segmentation. To test if object boundaries defined by differences in spatial frequency would strongly activate V1 neurons, we replaced textures in the CUB data set with gratings (CUB-grating images) with the frequency of 5.83/cycle to 15.55/cycle (high-frequency range) and 15.55/cycle to 58.3/cycle (low-frequency range). These frequency bounds were derived empirically from fitting gratings to 1000 random neurons compiled across 8 pilot data sets.

Analyses on highly activating crops in the CUB and CUB-grating data set To test whether the division of subfields identified from neurons’ bipartite parameterization corresponds to object boundary in natural images, we searched for highly activating natural crops in silico for each target neuron. We masked these crops with the MEI mask and standardized all masked stimuli such that they have the same mean and RMS contrast within the MEI mask as the target neuron’s MEI. We screened through 1 million of these standardized crops for each target neuron and identified 100 most highly activating natural crops as well as 100 random ones. A natural crop was classified as containing both object and background if it contained at least 20% of both object and background within the RF of the target neuron. For the CUB-grating images, we followed the same protocol but reported on the most exciting crop for simplicity (Fig. S5e). Note that our reported results were stable regardless of the specific choice of the number of stimuli searched and the definition of natural crops containing both object and background.

To measure the alignment of the image crop segmentation label and neurons’ bipartite mask in both the CUB and the CUB-grating data sets, we devised a matching score as follows. For each neuron, we first defined a bipartite mask \( m_{bipartite} \) from the bipartite parameterization of DEIs as \( m_{MEIb} - m_{Ib} \), where \( m_{MEIb} \) and \( m_{Ib} \) denotes the binarized version of the MEI mask \( m_{MEI} \) and the phase invariant subfield mask \( m_I \), respectively, when we hard-thresholded on the masks at 0.3. In \( m_{bipartite} \), a value of 1 and -1 denotes pixels within the phase invariant subfield and within the fixed subfield, respectively, and 0 elsewhere. On the other hand, for each crop from the CUB and the CUB-grating data sets, we first assigned pixels within the bird to 1 and pixels belonging to the background to -1 and then masked it with \( m_{MEIb} \).
to achieve the segmentation mask $m_{seg}$ of the crop. For each target neuron $i$ and each of its highly activating crops $j$, a matching score $S^{(i)}_{m}$ is calculated as follow:

$$S^{(i)} = \frac{m_{bi}^{(i)} * m_{seg}^{(i)}}{\sum m_{bi}^{(i)} * E_{i}}$$

A highly activating crop would yield a matching score of 1 if the phase invariant subfield perfectly matched with the bird and the fixed subfield perfectly matched with the background and a score of -1 if the reverse was true. For random crops, however, a score of 0 is expected.

**Perlin noise stimuli**

To quantify if the phase invariant and fixed subfields extracted from DEIs partially systematically preferred different frequencies, we replaced the DEI content in each subfield with synthesized textures of Perlin noise (27) with different spatial frequency biases. Specifically, the texture images with size of $64 \times 64$ pixels were generated by combining 8 octaves of Perlin noises with lacunarity factor of 2 and a persistence of either 1.0 (high frequency texture images) or 0.1 (low frequency texture images).

**In vivo response comparisons and statistical analysis**

Recorded responses were normalized across all presented images per scan. The normalized responses of the matched neurons were then averaged across either 20 repetitions of a single image for MEI or single repetitions of 20 different images for all other image types including non-parametric DEIs (both pixel-based and population-based), synthesized & natural image controls, partial- & full-texture DEIs, fixed & phase invariant subfields alone, partial-texture DEIs with swapped subfields. For single neurons, the statistical significance of the difference in responses was assessed using either one-tailed or two-tailed Welch’s t-tests. The overall difference in average responses pooled across all neurons was assessed using either one-sample t-test, although this was not explicitly tested.

**Statistics**

All statistical tests performed with the corresponding sample sizes, statistical values, and p-values are reported in the figure caption. All permutation tests and bootstrappings were performed with 10,000 permutations/resamplings. Normality was assumed for Welch’s t-test and one-sample t-test, although this was not explicitly tested. When p-values were less than $10^{-9}$, they were reported as $P < 10^{-9}$; otherwise the exact p-values were reported.

**Data and code availability.** All data and the analysis code will be made publicly available in an online repository latest upon journal publication. Please contact us if you would like to get access before that time.

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**AUTHOR CONTRIBUTIONS**


**References**

Supplementary Information

Supplementary Fig. 1 — Example stimuli presented in closed-loop experiments
Supplementary Fig. 2 — MEI, DEIs, and bipartite mask of partial texture DEIs generated from signals at different depths of the same neuron
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Supplementary Fig. 9 — DEI Bipartite masks align with object boundaries in highly activating natural crops
Supplementary Fig. 10 — Phase invariant subfields prefer higher frequency content than fixed subfields
Fig. S1. Example stimuli presented in closed-loop experiments. MEI, non-parametric DEIs, natural & synthesized controls, and partial- & full-texture DEIs that were presented back to the animals in closed-loop experiments for 9 example neurons.
Fig. S2. MEI, DEIs, and bipartite mask of partial-texture DEIs generated from signals at different depths of the same neuron. Most neurons were scanned multiple times due to the dense calcium imaging with field 5 µm apart in depths. MEI, DEIs, and partial-texture DEIs were generated for neuronal slices at different depths.
Fig. S3. MEI and DEIs activated neurons with high specificity in all mice. The confusion matrices show the responses of each neuron to the MEI (left) and DEIs (right) of all target neurons in individual scans where we presented the stimuli back to the mouse on day 2 and beyond in closed-loop experiments. MEI responses were averaged across 20 repeats of the same image while DEIs responses were averaged across 20 different images with single repeat. The responses of each neuron were normalized, and each row was scaled so the maximum response across all images equals 1. Responses of neurons to their own MEI and DEIs (along the diagonal) were larger than to other MEIs and DEIs, respectively (one-sided permutation test, $P < 10^{-3}$ for both cases across all mice).
Fig. S4. Population-based DEIs evoked strong and selective responses in target neurons while containing perceivable differences. a. Examples of MEI and non-parametric DEIs generated with pixel-based (odd rows) or in silico population-based (even rows) diversity for 4 example neurons. b. Population-based DEIs activated neurons with high specificity. The confusion matrices show the responses of each neuron to MEI (left) and DEIs (right) of 44 neurons. MEI responses were averaged across 20 repeats of the same image while DEIs responses were averaged across 20 different images with single repeat. The responses of each neuron were normalized, and each row was scaled so the maximum response across all images equals 1. Responses of neurons to their own MEI and population-based DEIs (along the diagonal) were larger than to other MEIs and DEIs respectively (one-sided permutation test, $P < 10^{-10}$ for both cases). c. In vivo response to population-based DEIs versus MEI. Each point corresponds to the normalized response of a single neuron. The linear relationship between DEIs and MEI responses was estimated by averaging over 1000 repeats of robust linear regression using the RANSAC algorithm (21). DEIs stimulated in vivo closely to the level predicted in silico with respect to MEI (76% versus 85%) (two-sided Wilcoxon signed-rank test, $W = 8464, P = 7.69 \times 10^{-3}$), with only 12.1% of all neurons showing different responses between DEIs and 85% of MEI ($P < 0.05$, two-tailed Welch’s $t$-test with 32.6 average d.f.). Data were pooled over 207 neurons from 5 mice. d, e. Each point corresponds to the normalized activity of a single neuron in response to its population-based DEIs versus its synthesized controls (d) or natural image (e) controls. Response to each stimulus type was averaged over 20 different images with single repeat. DEIs activated their target neurons stronger than their corresponding synthesized and natural image controls (one-sided Wilcoxon signed-rank test, $W = 3441, P < 10^{-9}$ and $W = 3466, P < 10^{-13}$ respectively) with 26.6% and 28.5% of all neurons showing greater responses to non-parametric DEIs ($P < 0.05$, one-tailed Welch’s $t$-test with 31.0 and 30.6 average d.f., respectively). f. Differences between the most different pair of DEIs in pixel space are distinguishable by the mouse V1 population. Logistic regression classifiers were used to decode DEI identity of individual trials from V1 population responses. Decoding accuracies across neurons (median=0.93) were higher than chance level (0.5 as indicated by the dashed line) (one sample $t$-test, $t = 138.5, P < 10^{-9}$). Data were pooled over 235 neurons from 3 mice.
Fig. S5. Replication of DEIs in the MICrONS functional connectomics data set. a, Illustration of how we applied our DEI methodology in the “MICrONS” functional connectomics data set. We used neuronal responses to natural images to train a static model and optimize MEIs and DEIs (“static”, S). Then, we used the responses of the same neurons to natural movie stimuli used in the MICrONS data set to first train a dynamic model and then predict responses to static images. Based on these predictions, we trained a separate static model and optimized MEI and DEIs for the same neurons (“Dynamic”, D). b, Neuron responses to natural image crops were sparse and smaller than those to MEI and DEIs synthesized from both static and dynamic static models. The gray curves show the fraction of natural image crops that elicit no less than certain normalized activation (normalized by S-MEI activation) for 150 random model neurons; the black curve indicates the average. Arrows denote the average activation of each stimulus type on the x axis. On average, only 1.0% of all 5000 natural crops produced in silico responses above D-DEIs. There existed only 0.1% of natural crops in between D-MEI and S-MEI activation and only 1.0% in between D-DEIs and S-DEIs activation. For a representative cell (red), example images at different activation levels were shown, along with its S-MEI, D-MEI, S-DEIs, and D-DEIs. Data were pooled over 3 mice for a total of 935 neurons. c, Cumulative distribution of neurons with an equal or larger fraction of natural images crops eliciting activation in between D-MEI and S-MEI activation (gray) and in between D-DEIs and S-DEIs activation (black) (90% and 70% of all neurons have only 1% of natural patches in between D-MEI and S-MEI activation and in between D-DEIs and S-DEIs activation, respectively (red dashed line)). d, Diversity index of D-DEIs versus S-DEIs. Data was pooled over 3 mice for a total of 4268 neurons for MEI and 679 for DEI analysis. e, We used the above pipeline to optimize MEIs and DEIs for neurons in the MICrONS data set itself. This allows future circuit-level dissections towards understanding the mechanism underlying the novel bipartite invariances in mouse visual cortex.
Fig. S6. Selection and quantification of partial-texture DEIs. 

**a**, Each curve shows how the normalized average in silico activation of texture-based DEIs decreases as the normalized size of phase invariant subfield increases. Texture DEIs with no phase invariant subfield (i.e., fixed subfield only) are equivalent to the MEI. Texture DEIs with phase invariant subfield spanning the entire RF is equivalent to full-texture DEIs. The set of partial-texture DEIs is selected for each neuron to jointly maximize both the average response and diversity (see Method for more details). 

**b**, Diversity index of partial-texture DEIs versus non-parametric DEIs. Data were shown for a total of 401 neurons pooled over 8 mice that have been tested in closed-loop experiments.

Fig. S7. Both subfields of the partial-texture DEIs are necessary and specific for evoking high in vivo responses. 

**a**, Examples of the MEI, fixed or phase invariant subfield of the MEI only, partial-texture DEIs, and partial-texture images with either fixed or phase invariant subfield swapped to random content for two example neurons. The fixed subfield was swapped to 20 different random natural patches whereas the phase invariant subfield was swapped to patches cropped from 20 different random non-self neurons’ preferred textures for the phase invariant subfield. 

**b-e**, Each point corresponds to the normalized response of a single neuron averaged over 20 repeats (MEI) or averaged over 20 single-trial different stimuli of the same type. 

**b-c**, Fixed or phase invariant subfield of the MEI only activated the target neurons weaker than the MEI (one-sided Wilcoxon signed-rank test, \( W = 21650, P < 10^{-3} \), and \( W = 17873, P < 10^{-9} \), respectively) with 50.2% and 21.4% of all neurons showing weaker responses than their MEIs (\( P < 0.05 \), one-tailed Welch’s t-test with 29.9 and 33.1 average d.f., respectively). 

**d-e**, Both fixed and phase invariant subfield swapped stimuli activated the target neurons weaker than partial-texture DEIs (one-sided Wilcoxon signed-rank test, \( W = 31710, P < 10^{-3} \), and \( W = 27850, P < 10^{-6} \), respectively) with 40.9% and 22.0% of all neurons showing weaker responses than their partial-texture DEIs (\( P < 0.05 \), one-tailed Welch’s t-test with 30.0 and 32.5 average d.f., respectively). Data were pooled over 5 mice, displaying a total of 259 neurons.
Fig. S8. DEI closed-loop verification for randomly selected neurons. a Confusion matrices for MEI and non-parametric DEIs show the responses of each neuron to the MEI (left) and DEIs (right) of all target neurons. The responses of each neuron were normalized, and each row was scaled so the maximum response across all images equals 1. Responses of neurons to their own MEI and DEIs (along the diagonal) were larger than to other MEIs and DEIs respectively (one-sided permutation test, $P < 10^{-9}$ for both cases). b-h. Each point corresponds to the normalized response of a single neuron averaged over 20 repeats (MEI) or averaged over 20 single-trial different stimuli of the same type. b. In vivo response to DEIs versus MEI. Each point corresponds to the normalized response of a single neuron. The linear relationship between DEIs and MEI responses was estimated by averaging over 1000 repeats of robust linear regression using the RANSAC algorithm (21). DEIs stimulated neurons in vivo closely to the level predicted in silico with respect to MEI (78% versus 85%) (two-sided Wilcoxon signed-rank test, $W = 982, P = 0.008$), with only 14% neurons showing different responses between DEIs and 85% of MEI ($P < 10^{-10}$, two-tailed Welch’s t-test with 33.0 average d.f.). c. DEIs failed to stimulate their target neurons compared to non-parametric DEIs (one-sided Wilcoxon signed-rank test, $W = 212, P = 1.53 \times 10^{-8}$) with 48.4% of all neurons showing weaker responses than non-parametric DEIs ($P < 0.05$, one-tailed Welch’s t-test with 30.5 average d.f.) d. DEIs activated their target neurons similarly to non-parametric DEIs (two-sided Wilcoxon signed-rank test, $W = 851, P = 0.21$) with only 9.4% of neurons showing different responses from corresponding non-parametric DEIs ($P < 0.05$, two-tailed Welch’s t-test with 34.4 average d.f.). e-f. Fixed or phase invariant subfield of the MEI only activated the target neurons weaker than the MEI (one-sided Wilcoxon signed-rank test, $W = 1942, P < 10^{-9}$, and $W = 1824, P = 7.9 \times 10^{-8}$, respectively) with 48.4% and 20.3% neurons showing weaker responses than their partial-texture DEIs ($P < 0.05$, one-tailed Welch’s t-test with 28.1 and 31.7 average d.f., respectively). g-h. Both fixed and phase invariant subfield swapped stimuli activated the target neurons weaker than partial-texture DEIs (one-sided Wilcoxon signed-rank test, $W = 1942, P < 10^{-9}$, and $W = 1824, P = 7.9 \times 10^{-8}$, respectively) with 48.4% and 20.3% neurons showing weaker responses than their partial-texture DEIs ($P < 0.05$, one-tailed Welch’s t-test with 28.1 and 31.7 average d.f., respectively). Data were collected from 1 mouse, displaying a total of 64 neurons.
Fig. S9. DEI bipartite masks align with object boundaries in highly activating natural crops. a, For each of the 4 example neurons, we show the MEI, the bipartite mask (black denotes the fixed subfield and white denotes the phase invariant subfield) extracted from partial-texture DEIs, 5 of the 100 highly activating natural crops, the corresponding original full-field natural images from which the crops were taken, and the corresponding segmentation labels of the natural crops. For each highly activating crop, a matching score is computed based on its segmentation label and the neuron’s bipartite mask. The value on the bottom right of each natural image crop shows the normalized model predicted activation of the image. Red dashed boxes on the full-field natural images indicate the natural image crops where the MEI masks were applied. The value on the bottom right of each segmentation label image shows its matching score. b, Histogram of averaged in silico response of top 100 highly activating natural crop sampled from CUB data set after normalized by their corresponding MEI (median is 0.695). c, When the bipartite mask was flipped with regard to the center of the RF, the matching score for the highly activating natural crops decreased (one-sided Wilcoxon signed-rank test $W = 204486$, $P < 10^{-7}$) with 41.3% of all neurons showing weaker matching scores than original bipartite mask ($P < 0.05$, one-tailed Welch’s t-test with 186.4 average d.f.). Example original bipartite mask and flipped bipartite mask were shown for one neuron (top left and bottom right respectively). The result was pooled over 6 mice for a total of 1200 neurons.
Fig. S10. Phase invariant subfields prefer higher frequency content than fixed subfields. a, Schematic illustrating the synthesis of Perlin noise stimuli ([27]) with different frequency biases. The MEI, bipartite mask, and synthesized stimuli for one example neuron were shown for demonstration. Two populations of full-field textures containing Perlin noise with different persistence coefficients (1.0 and 0.25) were synthesized (1 million each) to capture the diversity of natural textures at high and low frequencies (top and bottom). For each target neuron, the MEI mask was applied onto either high-frequency or low-frequency textures to create homogeneous stimuli (1 million high f-high f and 1 million low f-low f, respectively). In addition, the phase invariant subfield mask and the fixed subfield mask were applied onto different texture populations to create heterogeneous stimuli (1 million high f-high f and 1 million low f-low f). b, For each neuron, we screened through all 4 million synthesized stimuli to identify the most exciting one. The percentage of neurons preferring each stimulus type is reported in the 4 quadrants of the stimulus space. Each quadrant also demonstrates one example stimulus created for one example neuron, with the value on the bottom right of each image showing its normalized model predicted activation. Neurons preferred stimuli with higher frequency in the phase invariant subfield more than stimuli with low or high frequency in both subfields, as well as stimuli with lower frequency in the phase invariant subfield (one-way chi-squared test, $\chi^2 = 314$, $P < 10^{-9}$ and one-sided test using bootstrapping, $P < 10^{-9}$ for all three comparisons). The percentages of neurons for each quadrant with bootstrapped standard deviation were reported. Data was pooled over 6 mice for a total of 1200 neurons.