

# MIT Open Access Articles

## *Unsupervised Natural Visual Experience Rapidly Reshapes Size-Invariant Object Representation in Inferior Temporal Cortex*

The MIT Faculty has made this article openly available. *[Please](https://libraries.mit.edu/forms/dspace-oa-articles.html) share* how this access benefits you. Your story matters.

**Citation:** Li, Nuo, and James J. DiCarlo. "Unsupervised Natural Visual Experience Rapidly Reshapes Size-Invariant Object Representation in Inferior Temporal Cortex." Neuron 67, no. 6 (September 23, 2010): 1062–1075. © 2010 Elsevier Inc.

**As Published:** http://dx.doi.org/10.1016/j.neuron.2010.08.029

**Publisher:** Elsevier

**Persistent URL:** <http://hdl.handle.net/1721.1/96054>

**Version:** Final published version: final published article, as it appeared in a journal, conference proceedings, or other formally published context

**Terms of Use:** Article is made available in accordance with the publisher's policy and may be subject to US copyright law. Please refer to the publisher's site for terms of use.





## <span id="page-1-0"></span>Unsupervised Natural Visual Experience Rapidly Reshapes Size-Invariant Object Representation in Inferior Temporal Cortex

## Nuo Li<sup>1</sup> and James J. DiCarlo<sup>1,\*</sup>

1McGovern Institute for Brain Research, Department of Brain and Cognitive Sciences, Massachusetts Institute of Technology, Cambridge, MA 02139, USA \*Correspondence: [dicarlo@mit.edu](mailto:dicarlo@mit.edu)

DOI 10.1016/j.neuron.2010.08.029

#### **SUMMARY**

We easily recognize objects and faces across a myriad of retinal images produced by each object. One hypothesis is that this tolerance (a.k.a. ''invariance'') is learned by relying on the fact that object identities are temporally stable. While we previously found neuronal evidence supporting this idea at the top of the nonhuman primate ventral visual stream (inferior temporal cortex, or IT), we here test if this is a general tolerance learning mechanism. First, we found that the same type of unsupervised experience that reshaped IT position tolerance also predictably reshaped IT size tolerance, and the magnitude of reshaping was quantitatively similar. Second, this tolerance reshaping can be induced under naturally occurring dynamic visual experience, even without eye movements. Third, unsupervised temporal contiguous experience can build new neuronal tolerance. These results suggest that the ventral visual stream uses a general unsupervised tolerance learning algorithm to build its invariant object representation.

## INTRODUCTION

Our ability to recognize objects and faces is remarkably tolerant to variation in the retinal images produced by each object. That is, we can easily recognize each object even though it can appear in different positions, sizes, poses, etc. In the primate brain, the solution to this ''invariance'' problem is thought to be achieved through a series of transformations along the ventral visual stream. At the highest stage of this stream, the inferior temporal cortex (IT), a tolerant object representation is obtained in which individual IT neurons have a preference for some objects (''selectivity'') over others, and this rank-order preference is largely maintained across identity-preserving image transformations ([Ito et al., 1995; Logothetis and Sheinberg,](#page-13-0) [1996; Tanaka, 1996; Vogels and Orban, 1996\)](#page-13-0). Though most IT neurons are not strictly ''invariant'' [\(DiCarlo and Maunsell,](#page-13-0) [2003; Ito et al., 1995; Logothetis and Sheinberg, 1996; Vogels](#page-13-0) [and Orban, 1996](#page-13-0)), reasonably sized populations of these socalled ''tolerant'' neurons can support object recognition tasks [\(Afraz et al., 2006; Hung et al., 2005; Li et al., 2009](#page-13-0)). However, we do not yet understand how IT neurons construct this tolerant response phenomenology.

One potentially powerful idea is that time can act as an implicit teacher, in that the temporal contiguity of object features during natural visual experience can instruct the learning of tolerance, potentially in an unsupervised manner ([Foldiak, 1991; Masque](#page-13-0)[lier et al., 2007; Masquelier and Thorpe, 2007; Sprekeler et al.,](#page-13-0) [2007; Stryker, 1991; Wiskott and Sejnowski, 2002; Wyss et al.,](#page-13-0) [2006\)](#page-13-0). The overarching logic is as follows: during natural visual experience, objects tend to remain present for seconds or more, while object motion or viewer motion (e.g., eye movements) tend to cause rapid changes in the retinal image cast by each object over shorter time intervals (hundreds of ms). In theory, the ventral stream could construct a tolerant object representation by taking advantage of this natural tendency for temporally contiguous retinal images to belong to the same object, thus yielding tolerant object selectivity in IT cortex. A recent experimental result in adult nonhuman primate IT has provided some neuronal support for this temporal contiguity hypothesis ([Li and DiCarlo, 2008](#page-13-0)). Specifically, we found that alterations of unsupervised experience of temporally contiguous object image changes across saccadic eye movements can induce rapid reshaping (within hours) of IT neuronal position tolerance (i.e., a reshaping of each IT neuron's ability to respond with consistent object selectivity across the retina). This IT neuronal learning likely has perceptual consequences because similar temporal contiguity manipulations of eye-movementdriven position experience can produce qualitatively similar changes in the position tolerance of human object perception [\(Cox et al., 2005\)](#page-13-0).

However, these previous studies have two key limitations. First, they only uncovered evidence for temporal contiguity learning under a very restricted set of conditions: they showed learning effects only in the context of eye movements, and they only tested one type of tolerance—position tolerance. Because eye movements drive a great deal of the image statistics relevant only to position tolerance (temporally contiguous image translations), the previous results could reflect only a special case of tolerance learning. Second, the previous studies did not directly show that temporally contiguous image statistics can *build* new tolerance, but only showed that alterations of

<span id="page-2-0"></span>

#### Figure 1. Experimental Design and Prediction

(A) IT selectivity was tested in the *Test Phases* whereas animals received experience in the altered visual world in the *Exposure Phases*. (B) The chart shows the full exposure design for a single IT site in Experiment I. Arrows show the temporal contiguity experience of retinal images (arrow heads point to the retinal images occurring later in time, e.g., A). Each arrow shows a particular exposure event type (i.e., temporally linked images shown to the animal), and all eight exposure event types were shown equally often (randomly interleaved) in each *Exposure Phase*.

(C) Prediction for IT responses collected in the *Test Phase*: if the visual system builds size tolerance using temporal contiguity, the swap exposure should cause incorrect grouping of two different object images (P and N). The qualitative prediction is a decrease in object selectivity at the swap size (images and data points outlined in red) that grows stronger with increasing exposure (in the limit, reversing object preference as illustrated schematically here), and little or no change in object selectivity at the non-swap size. The experiment makes no quantitative prediction for the selectivity at the medium size (gray oval, see text).

those statistics can disrupt normal tolerance. Because of these limitations, we do not know if the naive ventral stream uses a general, temporal contiguity-driven learning mechanism to construct its tolerance to all types of image variation.

Here, we set out to test the temporal contiguity hypothesis in three ways. First, we reasoned that, if the ventral stream is using temporal contiguity to drive a general tolerance-building mechanism, alterations in that temporal contiguity should reshape other types of tolerance (e.g., size tolerance, pose tolerance, illumination tolerance), and the magnitude of that reshaping should be similar to that found for position tolerance. We decided to test size tolerance, because normal size tolerance in IT is much better described [\(Brincat and Connor, 2004; Ito et al., 1995; Logothetis](#page-13-0) [and Sheinberg, 1996; Vogels and Orban, 1996](#page-13-0)) than pose or illumination tolerance. Our experimental logic follows our previous work on position tolerance ([Cox et al., 2005; Li and Di-](#page-13-0)[Carlo, 2008](#page-13-0)). Specifically, when an adult animal with a mature (e.g., size-tolerant) object representation is exposed to an altered visual world in which object identity is consistently swapped across object size change, its visual system should learn from those image statistics such that it predictably ''breaks'' the size tolerance of that mature object representation. Assuming IT conveys this object representation [\(Afraz et al.,](#page-13-0) [2006; Hung et al., 2005; Logothetis and Sheinberg, 1996;](#page-13-0) [Tanaka, 1996\)](#page-13-0), that learning should result in a specific change in the size tolerance of mature IT neurons (Figure 1).

Second, many types of identity-preserving image transformations in natural vision do not involve intervening eye movements (e.g., object motion producing a change in object image size). If the ventral stream is using a general tolerance-building mechanism, we should be able to find size tolerance reshaping even without intervening eye movements, and we should also be able to find size tolerance reshaping when the dynamics of the image statistics mimic naturally occurring image dynamics.

Third, our previous studies ([Cox et al., 2005; Li and DiCarlo,](#page-13-0) [2008\)](#page-13-0) and our first two aims above use the breaking of naturally occurring image statistics to try to break the normal tolerance observed in IT (i.e., to weaken existing IT object selectivity in a position- or size-specific manner; Figure 1). Such results support the inference that naturally occurring image statistics instruct the ''building'' of that tolerance in the naive ventral stream. However, we also sought to test that inference more directly by looking for evidence that temporally contiguous image statistics can build new tolerance in IT neurons with immature tolerance (i.e., can produce an increase in existing IT object selectivity in a positionor size-specific manner).

Our results showed that targeted alterations in the temporal contiguity of visual experience robustly and predictably reshaped IT neuronal size tolerance over a period of hours. This change in size tolerance grew gradually stronger with increasing visual experience, and the rate of reshaping was very similar to previously reported position tolerance reshaping [\(Li and DiCarlo,](#page-13-0) [2008\)](#page-13-0). Second, we found that the size tolerance reshaping occurred without eye movements, and it occurred when the dynamics of the image statistics mimicked naturally occurring dynamics. Third, we found that exposure to ''broken'' temporal contiguity image statistics could weaken and even reverse the previously normal IT object selectivity at a specific position or

size (i.e., exposure could break old correct tolerance and build new ''incorrect'' tolerance), and that naturally occurring temporal contiguity image statistics could build new, correct position or size tolerance. Taken together with previous work, these results argue that the ventral stream uses unsupervised, natural visual experience and a common learning mechanism (a.k.a. unsupervised temporal tolerance learning, or UTL) to build and maintain its tolerant (invariant) object representation.

#### RESULTS

In three separate experiments (Experiments I, II, III), two unsupervised nonhuman primates (Rhesus monkeys, *Macaca mulatta*) were exposed to altered visual worlds in which we manipulated the temporal contiguity statistics of the animals' visual experience with object size [\(Figure 1A](#page-2-0), *Exposure Phases*). In each experiment, we recorded multiunit activity (MUA) in an unbiased sample of recording sites in the anterior region of IT to monitor any experience-induced change ([Figure 1](#page-2-0)A, *Test Phases*). Specifically, for each IT site, a preferred object (P) and a less-preferred object (N) were chosen based on testing of a set of 96 objects ([Figure 1B](#page-2-0)). We then measured the baseline IT neuronal selectivity for P and N at three retinal sizes  $(1.5^{\circ}, 4.5^{\circ},$ and  $9^{\circ}$ ) in a *Test Phase* ( $\sim$ 10 min) by presenting the object images in a rapid but naturally paced sequence (5 images/s) on the animals' center of gaze. For all the results below, we report selectivity values determined from these *Test Phases*, which we conducted both before and after experience manipulations. Thus, all response data shown in the results below were collected during orthogonal behavioral tasks in which object identity and size were irrelevant ([Supplemental Experi](#page-13-0)[mental Procedures](#page-13-0) available online).

Consistent with previous reports ([Kreiman et al., 2006\)](#page-13-0), the initial *Test Phase* data showed that each IT site tended to maintain its preference for object P over object N at each size tested here ([Figures 3](#page-6-0) and [S3](#page-13-0) available online). That is, most IT sites showed good, baseline size tolerance. Following the logic outlined in the [Introduction,](#page-1-0) the goal of Experiments I–III was to determine if consistently applied, unsupervised experience manipulations would predictably reshape that baseline size tolerance of each IT site (see [Figure 1](#page-2-0) for the basic prediction). In particular, we monitored changes in each IT site's preference for object P over N at each of the three objects sizes, and any change in that selectivity following experience that was not seen in control conditions was taken as evidence for an experience-induced reshaping of IT size tolerance.

In each experiment, the key experience manipulation was deployed in one or more *Exposure Phases* that were all under precise, automated computer-display control to implement spatiotemporally reliable experience manipulations (see [Experi](#page-12-0)[mental Procedures](#page-12-0)). Specifically, during each *Exposure Phase* the animals freely viewed a gray display monitor on which images of object P or N intermittently appeared at randomly chosen retinal positions away from the center of gaze (object size: 1.5°, 4.5°, or 9°). The animals almost always looked to foveate each object (>95% of object appearances) within  $\sim$ 124 ms (mean; median, 109 ms), placing the object image on the center of gaze. Following that object acquisition saccade,

we reliably manipulated the visual experience of the animals over the next 200–300 ms. The details of the experience manipulation (i.e., which object sizes where shown and the timing of those object images) were different in the three experiments, but all three experiments used the same basic logic outlined in the [Introduction](#page-1-0) and in [Figure 1](#page-2-0).

#### Experiment I: Does Unsupervised Visual Experience Reshape IT Size Tolerance?

In Experiment I, following the object acquisition saccade, we left the newly foveated object image unchanged for 100 ms, and then we changed the size of the object image (while its retinal position remained on the animal's center of gaze) for the next 100 ms ([Figure 1](#page-2-0)A). We reasoned that this creates a temporal experience linkage (''exposure event'') between one object image at one size and another object image at another size. Importantly, on half of the exposure events, one object was swapped out for the other object: for example, a medium-sized  $(4.5^{\circ})$  object P would become a big  $(9^{\circ})$  object N [\(Figure 1](#page-2-0)A, ''swap exposure event''). As one key control, we also exposed the animal to more normal exposure events in which object identity did not change during the size change [\(Figure 1](#page-2-0)A, ''non-swap exposure event''). The full exposure design for one IT site is shown in [Figure 1](#page-2-0)B; the animal received 800–1600 swap exposures within the time period of 2–3 hr. Each day, we made continuous recordings from a single IT site, and we always deployed the swap exposure at a particular object size (either 1.5 $^{\circ}$  or 9 $^{\circ}$ , i.e., swap size) while keeping the other size as a control (i.e., non-swap size). Across different IT sites (i.e., different recording days), we strictly alternated the object size at which swap manipulation took place so that object size was counterbalanced across our recorded IT population ( $n = 27$ ).

UTL theory makes the qualitative prediction that the altered experience will induce a size-specific confusion of object identity in the IT response as the ventral stream learns to associate the temporally linked images. In particular, our exposure design should cause the IT site to reduce its original selectivity for images of object P and N at the swap size (perhaps even reversing that selectivity in the limit of large amounts of experience; [Figure 1](#page-2-0)C, red). UTL is not currently specific enough to make a quantitative prediction of what this altered experience should do for selectivity among the medium object size images because those images were temporally paired in two ways: with images at the swap size (altered visual experience) and with the images at the non-swap size (normal visual experience). Thus, our key experimental prediction and planned comparison is between the selectivity (P versus N) at the swap and non-swap size: we predict a selectivity decrease at the swap size that should be much larger than any selectivity change at the nonswap object size ([Figure 1](#page-2-0)C, blue).

This key prediction was born out by the data: as the animals received experience in the altered visual world, IT selectivity among objects P and N began to decrease at the swap size, but not at the control size. This change in selectivity grew stronger with increasing experience over the time course of 2–3 hr ([Figure 2](#page-4-0)A). To quantify the selectivity change, for each IT site, we took the difference between the selectivity ( $P - N$ , response difference in units of spikes/s, see [Experimental](#page-12-0)

<span id="page-4-0"></span>



[Procedures\)](#page-12-0) in the first (pre-exposure) and last *Test Phase*. This  $\Delta(P - N)$  sought to quantify the total amount of selectivity change for each IT site induced by our experience manipulation. On average, there was a significant decrease in selectivity at the swap size (Figure 2B, p < 0.0001, two-tailed t test against 0) and no significant change at the non-swap control size (Figure  $2B$ ,  $p = 0.89$ ). Incidentally, we also observed a significant decrease in selectivity at the medium size ( $p = 0.002$ ). This is not surprising given that the images at the medium object size were exposed to the altered statistics half of the time when they were temporally paired with the images at the swap size. Because no prediction was made about the selectivity change at the medium size, we concentrate below on the planned comparison between the swap and non-swap size. We statistically confirmed the size

## Figure 2. Experimental I and II Key Results

(A) Mean  $\pm$  SEM. IT object selectivity change,  $\Delta(P - N)$ , from the first *Test Phase* as a function of the number of exposure events is shown. Each data point shows the average across all the sites tested for that particular amount of experience  $(n = 27, 800$  exposure events;  $n = 22, 1600$  exposure events). (B) Mean  $\pm$  SEM selectivity change at the swap, non-swap, and medium size (4.5°). For each IT site (n = 27), total  $\Delta(P - N)$ was computed using the data from the first and last *Test Phase*, excluding any middle *Test Phase* data. Hence, not all data from (A) were included.  $p < 0.05$  by two-tailed t test;  $*$ <sup>\*</sup> $p$  < 0.01; n.s.  $p > 0.05$ .

(C) For each IT site (n = 27), we fit a line (linear regression) to the  $(P - N)$  data as a function of the number of exposure events (insert). We used the slope of the line fit,  $\Delta s(P - N)$ , to quantify the selectivity change. The  $\Delta s(P - N)$  is a measure that leverages all our data while normalizing out the variable of exposure amount [for sites with only two *Test Phases*,  $\Delta s(P - N)$  equals  $\Delta(P - N)$ ].  $\Delta s(P - N)$  was normalized to show selectivity change per 800 exposure events. Error bars indicate the standard error of the procedure to compute selectivity ([Supplemental Experimental Procedures\)](#page-13-0). M1, monkey 1; M2, monkey 2.

(D) Mean  $\Delta s(P - N)$  at the swap and non-swap size (n = 27 IT sites; M1: 7, M2: 20). Error bars indicate SEM over neuronal sites.

(E) Change in selectivity,  $\Delta s(P - N)$ , of all IT sites from Experiment II at the swap and non-swap size.

(F) Mean  $\pm$  SEM  $\Delta s(P - N)$  at the swap and non-swap size.

specificity of the experience-induced decrease in selectivity by two different approaches: (1) a direct t test on the  $\Delta(P - N)$  between the swap and nonswap size ( $p < 0.001$ , two-tailed), and (2) a significant interaction of "exposure  $\times$  object size" on the raw selectivity measurements  $(P - N)$ —that is, IT selectivity was decreased by exposure only at the swap size ( $p = 0.0018$ , repeated-measures ANOVA;  $p = 0.006$ , bootstrap, see [Supplemental](#page-13-0) [Experimental Procedures\)](#page-13-0).

To ask if the experience-induced selectivity change was specific to the manipulated objects or the features contained in those objects, we also tested each IT site's responses to a second pair of objects (P' and N', control objects; see [Experimental Procedures](#page-12-0)). Images of these control

objects at three sizes were tested together with the swap objects during all *Test Phases* (randomly interleaved), but they were not shown during the *Exposure Phase*. On average, we observed no change in IT selectivity among these unexposed control objects [\(Figure S4\)](#page-13-0). This shows that that the experience-induced reshaping of IT size tolerance has at least some specificity for the experienced objects or the features contained in those objects.

We next set out to quantify the amount of IT size tolerance reshaping induced by the altered visual experience. Because each IT site was tested for different amounts of exposure time (due to experimental time constraints), we wanted to control for this and still leverage all the data for each site to gain maximal power. To do so, we fit linear regressions to the  $(P - N)$  selectivity of individual sites at each object size (Figure 2C, insert). The

slope of the line fit, which we will refer to as  $\Delta s(P - N)$ , provided us with a sensitive, unbiased measure of the amount of selectivity change that normalizes the amount of exposure experience. The  $\Delta s(P - N)$  for the swap size and non-swap size is shown in [Figures 2](#page-4-0)C and 2D, which qualitatively confirmed the result obtained in [Figure 2](#page-4-0)B (using the simple measure of selectivity change), and showed a mean selectivity change of  $-9.2$  spikes/s for every 800 swap exposure events.

Importantly, we note that this reshaping of IT tolerance was induced by unsupervised exposure to temporally linked images that did not include a saccadic eye movement to make that link ([Figure 1](#page-2-0)A). We also considered the possibility that small intervening microsaccades might still have been present, but found that they cannot account for the reshaping ([Figure S7\)](#page-13-0). The size specificity of the selectivity change also rules out alternative explanations such as adaptation, which would not predict this specificity (because our exposure design equated the amount of exposure for both the swap and non-swap size). We also found the same amount of tolerance reshaping when the sites were grouped by the physical object size at which we deployed the swap (1.5 $\degree$  versus 9 $\degree$ , p = 0.26, t test). Thus the learning is independent of low-level factors like the total luminance of the swapped objects. In sum, we found that unsupervised, temporally linked experience with object images across object size change can reshape IT size tolerance.

## Experiment II: Does Size Tolerance Learning Generalize to the ''Natural'' Visual World?

In the natural world, objects tend to undergo size change smoothly on our retinas as a result of object motion or viewer motion, but, in Experiment I (above), the object size changes we deployed were discontinuous: one image of an object was immediately replaced by an image of another object with no smooth transition ([Figure 2,](#page-4-0) top). Therefore, although those results show that unsupervised experience with object images at different sizes linked in time could induce the predicted IT selectivity change, we wanted to know if that learning was also found during exposure to more natural (i.e., temporally smooth) image dynamics.

To answer this question, we carried out a second experiment (Experiment II) in which we deployed essentially the same manipulation as Experiment I (object identity changes during object size changes, no intervening eye movement), but with natural (i.e., smooth-varying) stimulus sequences. The dynamics in these movie stimuli were closely modeled after the kind of dynamics that our visual system encounters daily in the natural environment ([Figure S2](#page-13-0)). To create smooth-varying object identity changes over object size changes, we created morph lines between pairs of objects we swapped in Experiment I (P and N). This allowed us to parametrically transform the shape of the objects [\(Figure 2](#page-4-0), bottom). All other experimental procedures were identical to Experiment I except, in the *Exposure Phases*, objects underwent size change smoothly while changing identity (swap exposure) or preserving identity (nonswap exposure, [Figure S2\)](#page-13-0).

When we carried out this temporally smooth experience manipulation on a new population of IT sites ( $n = 15$ ), we replicated the Experiment I results [\(Figures 2](#page-4-0)E and 2F): there was a

predicted decrease in IT selectivity at the swap size and not at the non-swap control size. This size specificity of the effect was, again, confirmed statistically by (1) direct t test on the total selectivity change,  $\Delta(P - N)$ , between the swap and non-swap size  $[\Delta(P - N) = -10.3$  spikes/s at swap size, +2.8 at nonswap size;  $p < 0.0001$ , two-tailed t test]; and (2) a significant interaction of "exposure  $\times$  object size" on the raw selectivity measurements  $(P - N)(p < 0.001$ , repeated-measures ANOVA;  $p = 0.001$ , bootstrap). This result suggests that image linking across time is sufficient to induce tolerance learning in IT and is robust to the temporal details of that image linking (at least over the  $\sim$ 200 ms time windows of linking used here). More importantly, Experiment II shows that unsupervised size tolerance learning occurs in a spatiotemporal image regime encountered in real-world vision.

## Size Tolerance Learning: Observations and Effect Size **Comparison**

Despite a wide diversity in the initial tuning of the recorded IT multiunit sites, our experience manipulation induced a predictable selectivity change that was large enough to be observed in individual IT sites: 40% (17/42 sites, Experiment I and II data combined) of the individual IT sites showed a significant selectivity decrease at the swap size within a single recording session (only 7% of sites showed significant selectivity decrease at the non-swap size, which is essentially the fraction expected by chance; 3/42 sites, p < 0.05, permutation test, see [Supplemental](#page-13-0) [Experimental Procedures](#page-13-0)). Eight example sites are shown in [Figure 3.](#page-6-0)

We found that the magnitude of size-tolerance reshaping depended on the initial selectivity at the medium object size, 4.5 $\degree$  (Pearson correlation, r = 0.54, p < 0.01). That is, on average, IT sites that we initially encountered with greater object selectivity at the medium size underwent greater exposure-induced selectivity change at the swap size. This correlation is not simply explained by the hypothesis that it is easier to break highly selective neurons (e.g., due to factors that might have nothing to do with neuronal learning, such as loss of isolation), because the correlation was not seen for changes in selectivity at the non-swapped size ( $r = -0.16$ ,  $p = 0.35$ ) and we found no average change in selectivity at the non-swapped size [\(Figure 2](#page-4-0) and statistics above). Instead, this observation is consistent with the overarching hypothesis of this study: the initial image selectivity at the medium object size provides (at least part of) the driving force for selectivity learning because those images are temporally linked with the swapped images at the swap size.

The change in selectivity produced by the experience manipulation was found throughout the entire time period of the IT response, including the earliest part of that period where IT neurons are just beginning to respond above baseline  $(\sim$ 100 ms from stimulus onset, [Figure S5\)](#page-13-0). This shows that the experience-induced change in IT selectivity cannot be explained by changes in long lag feedback alone (>100 ms; also see [Discussion](#page-9-0)). On average, the selectivity change at the swap size resulted from both a decrease in the response to the image of the preferred object (P) and an increase in the response to the less preferred object (N). Consistent with this, we found that the

<span id="page-6-0"></span>

experience manipulation produced no average change in the IT sites' mean response rate ([Figure S5\)](#page-13-0).

In this study, we concentrated on multiunit response data because it had a clear advantage as a direct test of our hypothesis—it allowed us to longitudinally track IT selectivity during altered visual experience across the entirety of each experimental session. We also examined the underlying single-unit data and found results that were consistent with the multiunit data. [Figure 4](#page-7-0)A shows an example of a rare single-unit IT neuronal recording that we were able to track across an entire recording session  $(\sim 3$  hr). The confidence that we were recording from the same unit comes from the consistency of the unit's waveform and its consistent pattern of response among the nonexposed control object images [\(Figure 4B](#page-7-0)). During this stable recording, the  $(P - N)$  selectivity at the swap size gradually decreased while the selectivity at the non-swap size remained stable, perfectly mirroring the multiunit results described above. However these  $\sim$ 3 hr single-unit recordings were very rare because single units have limited hold-time in the awake primate physiology preparation. Thus we took a more standard population approach to analyze the single-unit data [\(Baker et al., 2002; Kobatake et al., 1998; Sakai and Miya](#page-13-0)[shita, 1991; Sigala et al., 2002\)](#page-13-0). Specifically, we performed spike-sorting analyses to obtain clear single units from each *Test Phase* ([Experimental Procedures](#page-12-0)). We considered each single unit obtained from each *Test Phase* as a sample of the IT population, taken either before or after the experience in the altered visual world. This analysis does not require that the

#### Figure 3. Example Single IT Sites

Mean  $\pm$  SEM. IT response to P (solid square) and N (open circle) as a function of object size for eight example IT sites (from both Experiment I and II). The data shown are from the first (''before exposure'') and last (''after exposure'') *Test Phase.* (A) Swap size, 1.5°; (B) swap size, 9° (highlighted by red boxes and arrows). Gray dotted lines show the baseline response to a blank image (interleaved with the test images).

sampled units were the same neurons. The prediction is that IT single units sampled after exposure (i.e., at the last *Test Phase* of each day) would be less size tolerant at the swap size than at the nonswap size. This prediction was clearly observed in our single-unit data ([Figure 4C](#page-7-0), after exposure, p < 0.05; for reference, the size tolerance before the exposure is also shown and we observed no difference between the swap and non-swap size). The result was robust to the choice of the criteria to define ''single units'' [\(Figure S6](#page-13-0)). Similarly, we found that each single-unit population sampled after successively more exposure showed a successively larger change in size tolerance ([Figure 4](#page-7-0)D).

We next aimed to quantify the absolute magnitude of this size tolerance learning effect across the different experience manipulations deployed here, and to compare that magnitude with our previous results on position-tolerance learning ([Li](#page-13-0)

[and DiCarlo, 2008](#page-13-0)). To do this, we plotted the mean selectivity change at the swap size from each experiment as a function of number of swap exposures ([Figure 5\)](#page-8-0). We found that Experiments I and II produced a very similar magnitude of learning:  $\sim$ 5 spikes/s per 400 swap exposures (also see [Discussion](#page-9-0) for comparison to previous work). This effect grew larger at this approximately constant rate for as long as we could run each experiment, and the magnitude of the size tolerance learning was remarkably similar to that seen in our previous study of position tolerance [\(Li and DiCarlo, 2008](#page-13-0)).

### Size and Position Tolerance Learning: Reversing Old IT Object Selectivity and Building New IT Object Selectivity

The results on size tolerance presented above and our previous study of position tolerance [\(Li and DiCarlo, 2008](#page-13-0)) both used the breaking of naturally occurring temporal contiguity experience to discover that we can break normal position tolerance and size tolerance (i.e., we can cause a decrease in adult IT object selectivity in a size- or position-specific manner). While these results are consistent with the inference that naturally occurring image statistics instruct the original building of that normal tolerance (see [Introduction\)](#page-1-0), we next sought to test that inference more directly. Specifically, we asked if the temporal contiguity statistics of visual experience can instruct the creation of new IT tolerance (i.e., if they can cause an increase in IT object selectivity in a size- or position-specific manner). Our experimental data offered two ways to test this idea (below), and both ways revealed that unsupervised temporal contiguity learning could

<span id="page-7-0"></span>

#### Figure 4. Single-Unit Results

(A) P versus N selectivity of a rare single-unit IT neuron that was isolated across an entire recording session  $(\sim]3$  hr).

(B) The example single-unit's response to the six control object images during each *Test Phase* and its waveforms (gray: all traces from a *Test Phase*; red: mean). (C) Mean ± SEM size tolerance at the swap (red) and non-swap (blue) size for single units obtained before and after exposure. Size tolerance for the control objects is also shown at these two sizes (black). Each neuron's size tolerance was computed as  $(P - N)/(P - N)_{\text{medium}}$ , where  $(P - N)$  is the selectivity at the tested size and  $(P - N)$ <sub>medium</sub> is the selectivity at the medium object size. Only units that showed selectivity at the medium size were included  $[(P - N)_{\text{medium}} > 1 \text{ spikes/s}]$ . The top and bottom panels include neurons that had selectivity for the swap objects, the control objects, or both. Thus they show different but overlapping populations of neurons. The result is unchanged if we only examine populations for which each neuron has selectivity for both the swap and control objects (i.e., the intersections of the neuronal populations in top and bottom panels; [Figure S6](#page-13-0)).

(D) Mean ± SEM size tolerance at the swap size further broken out by the amount of exposure to the altered visual statistics. To quantify the change in IT size tolerance, we performed linear regression of the size tolerance as a function of the amount of experience. Consistent with the multiunit results, we found a significant negative slope ( $\Delta$  size tolerance = -0.84 per 800 exposure; p = 0.002, bootstrap; c.f. -0.42 for multiunit, [Figure S6\)](#page-13-0). No decrease in size tolerance was observed at the non-swap control size ( $\Delta$  size tolerance = 0.30; c.f. 0.12 for multiunit).

indeed build new IT tolerance. To do these analyses, we took advantage of the fact that we found very similar effects for both size tolerance and position tolerance ([Li and DiCarlo,](#page-13-0) [2008](#page-13-0)), and we maximized our power by pooling the data across this experiment ([Figure 5](#page-8-0): size experiment I, II;  $n = 42$  MUA sites) and our previous position experiment ( $n = 10$  MUA sites). This pooling did not qualitatively change the result—the effects shown in [Figures 5 and 6](#page-8-0) below were seen in the size tolerance data alone [\(Figure S9](#page-13-0)).

First, as outlined in [Figure 1C](#page-2-0), a strong form of the UTL hypothesis predicts that our experience manipulation should not only degrade existing IT selectivity for P over N at the swap size/position, but should eventually reverse that selectivity and then build new incorrect selectivity for N over P ([Figure 1](#page-2-0)C; note that we refer to this as incorrect selectivity because the full IT response pattern is inappropriate for the veridical world in which objects maintain their identity across changes in position and size). Though the plasticity we discovered is remarkably strong  $(\sim5$  spikes/s per hour), it did not produce a selectivity reversal for the ''mean'' IT site within the 2 hr recording session ([Figure S5D](#page-13-0)). Instead, it only produced a  $\sim$  50% decrease in selectivity for that mean site, which is entirely consistent with

the fact that our mean IT site had reasonably strong initial selectivity for P over N (mean P  $-$  N =  $\sim$  20 spikes/s). To look more deeply at this issue, we made use of the well-known observation that not all adult IT neurons are identical— some have a large amount of size or position tolerance, whereas others show a small amount of tolerance [\(DiCarlo and Maunsell, 2003; Ito](#page-13-0) [et al., 1995; Logothetis and Sheinberg, 1996; Op De Beeck](#page-13-0) [and Vogels, 2000](#page-13-0)). Specifically, some IT sites strongly prefer object P to N at some sizes/positions, but show only weak  $(P - N)$  selectivity at the swap sizes/positions (this neuronal response pattern is illustrated schematically at the top of [Figure 6](#page-9-0)). We reasoned that examination of these sites should reveal whether our experience manipulation is capable of causing a reversal in selectivity and building of new selectivity. Thus, we used independent data to select neuronal subpopulations from our data pool with varying amounts of initial selectivity at the swap size/position [\(Supplemental Experimental Proce](#page-13-0)[dures](#page-13-0)). Note that all of these neuronal sites had robust selectivity for P over N at the medium sizes/positions (as schematically illustrated in [Figure 6](#page-9-0)A). This analysis revealed that our manipulation caused neuronal sites with weak initial selectivity at the swap size/position to reverse their selectivity, and to build new

<span id="page-8-0"></span>

#### Figure 5. Effect Size Comparisons across Different Experience **Manipulations**

Mean object selectivity change as a function of the number of swap exposure events for different experiments. For comparison, the data from a position tolerance learning experiment [\(Li and DiCarlo, 2008](#page-13-0)) are also shown. Plot format is the same as [Figure 2](#page-4-0)A without the error bars. Mean  $\pm$  SEM  $\Delta$  $(P - N)$  at the non-swap size/position is shown in blue (all experiments pooled). SUA, single-unit activity; MUA, multiunit activity.

selectivity (building incorrect selectivity for N over P), exactly as predicted by the UTL hypothesis [\(Figure 6\)](#page-9-0).

A second way in which our data might reveal whether UTL can build tolerance is to carefully look for any changes in selectivity at the non-swap (control) size/position. Our experiment was designed to present a large number of normal temporal contiguity exposures at that control size/position so that we would perfectly equate its amount of retinal exposure with that provided at the swap size/position. Although some forms of unsupervised temporal contiguity theory might predict that these normal temporal contiguity exposures should increase the  $(P - N)$  selectivity at the control size/position, we did not initially make that prediction ([Figure 1](#page-2-0)C, blue) because we reasoned that most IT sites would already have strong, adultlike selectivity for object P versus N at that size/position, such that further supporting statistics would have little to teach those IT sites ([Figure 7](#page-10-0)A, top right). Consistent with this, we found little mean change in  $(P - N)$  selectivity for the control condition in either our position tolerance experiment [\(Li and DiCarlo, 2008\)](#page-13-0) or our size tolerance experiment ([Figure 2,](#page-4-0) blue). However, examination of all of our IT sites revealed that some sites happened to have initially weak  $(P - N)$  selectivity at the control size/position while still having strong selectivity at the medium size/position ([Figure 7A](#page-10-0), top left). This suggested that these sites might be in a more naive state with respect to the particular objects being tested such that our temporal contiguity statistics might expand their tolerance for these objects (i.e., increase their  $P - N$  selectivity at the control size/position). Indeed, examination of these sites reveals that our exposure experiment caused a clear, significant building of new, correct selectivity among these sites ([Figure 7B](#page-10-0)), again directly demonstrating that unsupervised temporal contiguity experience can build IT tolerance.

## Experiment III: Does the Learning Depend on the Temporal Direction of the Experience?

Our results show that targeted alteration of unsupervised natural visual experience rapidly reshapes IT size tolerance—as predicted by the hypothesis that the ventral stream uses a temporal contiguity learning strategy to build that tolerance in the first place. Several instantiated computational models show how this conceptual strategy can build tolerance ([Foldiak, 1991; Mas](#page-13-0)[quelier et al., 2007; Masquelier and Thorpe, 2007; Wallis and](#page-13-0) [Rolls, 1997; Wiskott and Sejnowski, 2002; Wyss et al., 2006\)](#page-13-0), and such models can be implemented using variants of Hebbian-like learning rules that are dependent on the timing of spikes [\(Gerstner et al., 1996; Sprekeler et al., 2007; Wallis and Rolls,](#page-13-0) [1997; Morrison et al., 2008; Sprekeler and Gerstner, 2009](#page-13-0)). The time course and task independence of the observed learning are consistent with synaptic plasticity [\(Markram et al., 1997; Me](#page-14-0)[liza and Dan, 2006](#page-14-0)), but our data do not constrain the underlying mechanism. One can imagine ventral stream neurons using almost temporally coincident activity to learn which sets of its afferents correspond to features of the same object across size changes. If tolerance learning is spike timing dependent, any experience-induced change in IT selectivity might reflect any temporal asymmetries at the level of the underlying synaptic learning mechanism. For example, one hypothesis is that lingering postsynaptic activity caused by temporally leading images drives synaptic plasticity in afferents activated by temporally lagging images. Alternatively, afferents activated by temporally leading images might be modified by the later arrival of postsynaptic activity caused by temporally lagging images. Or a combination of both hypotheses might be the case. To look for reflections of any such underlying temporal asymmetry, we carried out a third experiment (Experiment III) centered on the question, ''Do temporally leading images teach temporally lagging ones, or vice-versa?''

We deployed the same experience manipulation as before (linking of different object images across size changes, the same as Experiment I), but this time only in one direction (compare single-headed arrows in [Figure 8A](#page-11-0) with doubleheaded arrows in [Figure 1](#page-2-0)B). For example, during the recording of a particular IT site, the animal only received experience seeing objects temporally transition from a small size (arrow ''tail'' in [Fig](#page-11-0)[ure 8A](#page-11-0)) to a large size (arrow ''head'' in [Figure 8](#page-11-0)A) while swapping identity. We strictly alternated the temporal direction of the experience across different IT sites. That is, for the next IT site we recorded, the animal experienced objects transitioning from a large size to a small size while swapping identity. Thus, object size was counterbalanced across our recorded population, so that we could isolate changes in selectivity among the temporally leading stimuli (i.e., arrow tail stimuli) from changes in selectivity among the temporally lagging stimuli (i.e., arrow head stimuli). As in Experiments I and II, we measured the expression of any experience-induced learning by looking for any change in  $(P - N)$ selectivity at each object size measured in a neutral task with all images randomly interleaved (*Test Phase*). We replicated the results in Experiments I and II in that a decrease in  $(P - N)$ selectivity was found following swapped experience (red bars are negative in [Figure 8](#page-11-0)B). When we sorted our data based on the temporal direction of the animals' experience, we found greater selectivity change (i.e., learning) for the temporally lagging images ([Figure 8B](#page-11-0)). This difference was statistically significant ( $p = 0.038$ ,  $n = 31$ , two-tailed t test) and cannot be explained by any differences in the IT sites' initial selectivity [\(Figure S4](#page-13-0)C; also see [Figure S4](#page-13-0)B for results with all sites

## Neuron Natural Experience Reshapes IT Size Tolerance

<span id="page-9-0"></span>

### Figure 6. Altered Statistics in Visual Experience Builds Incorrect Selectivity

(A) Prediction: *top*, most adult IT neurons start with fully position/size tolerant selectivity (left). In the limit of a large amount of altered visual experience, temporal contiguity learning predicts that each neuron will acquire fully altered tolerance (right). *Bottom*, at the swap position/size (red), the selectivity for P over N is predicted to reverse in the limit (prefer N over P). Because we could only record longitudinally from a multiunit site for less than 3 hr, we do not expect our experience manipulation within a session to produce the full selectivity reversal (pre versus post) among neuronal sites with strong initial selectivity. However, because different IT sites differ in their degrees of initial selectivity, they start at different distances from selectivity reversal. Thus, our manipulation should produce selectivity reversal among the initially weakly selective sites and build new (''incorrect'') selectivity.

(B) Mean ± SEM normalized response to object P and N at the swap position/size among subpopulations of IT multiunit sites. Sites are grouped by their initial selectivity at the swap position/size using independent data. Data from the size and position tolerance experiments ([Li and DiCarlo,](#page-13-0) [2008](#page-13-0)) were combined to gain maximal power (size experiment I, II; position experiment, see [Supplemental Experimental Procedures](#page-13-0)). These sites show strong selectivity at the non-swap (control) position/size, and no negative change in that selectivity was observed (not shown). \*\*p  $<$ 0.01; \*p < 0.05, one-tailed t test against no change. (Size experiment data only, group  $1-6$ :  $p < 0.01$ ;  $p < 0.01$ ;  $p < 0.01$ ;  $p = 0.02$ ;  $p = 0.07$ ; n.s.).

I and II). We observed significant selectivity change for the medium size image pairs ([Figure 8B](#page-11-0), middle bar;  $p = 0.01$ , two-tailed t test against zero), which suggests that the tolerance learning has some degree of spread (but not to very different objects; [Figure S4\)](#page-13-0). Finally, the effect size observed in Experiment III was consistent with, and can explain the effect sizes observed in Experiments I and II. That is, based on the Experiment III effect sizes for the temporally lagging and leading images, a first-order prediction of the net effect in Experiments I and II is the average of these two effects

included). This result is consistent with an underlying learning mechanism that favors experience-induced plasticity of the afferents corresponding to temporally lagging images.

To test if the tolerance learning spread beyond the specifically experienced images, here, we also tested object images at an intermediate size  $(3^{\circ})$  between the two exposed sizes ([Figure 8](#page-11-0)). Unlike as in Experiments I and II, this medium size was not exposed to the animals during the *Exposure Phase* (it was also at a different physical size from the medium size in Experiments

(because Experiments I and II employed a 50-50 mix of the experience manipulations considered separately in Experiment III). That prediction is very close to what we found ([Figure 5](#page-8-0)).

## **DISCUSSION**

The overarching goal of this work is to ask whether the primate ventral visual stream uses a general, temporal contiguity driven learning mechanism to construct its tolerance to



<span id="page-10-0"></span>

#### Figure 7. Normal ("Correct") Statistics in Visual Experience Builds Tolerant Selectivity

(A) Prediction follows the same logic as in [Figure 6](#page-9-0)A, but here for the control conditions in which normal temporal contiguity statistics were provided [\(Figure 1\)](#page-2-0). *Top*, temporal contiguity learning predicts that neurons will be taught to build new "correct" selectivity (i.e., normal tolerance), and neurons starting with initially weak position/size tolerant selectivity (left) have the highest potential to reveal that effect. *Bottom*, at the non-swap position/size (blue), our manipulation should build new correct selectivity for P over N among IT sites with weak initial selectivity. (B) Mean  $\pm$  SEM normalized response to object P and N at the non-swap position/size among subpopulations of IT multiunit sites. Sites are grouped by their initial selectivity at the nonswap position/size using independent data. Other details are the same as those in [Figure 6](#page-9-0)B. (Size experiment data only, group 1-5:  $p = 0.06$ ;  $p <$ 0.01;  $p = 0.05$ ; n.s.; n.s.).

building statistics in the natural world: temporally contiguous image changes without intervening eye movements, and temporally smooth dynamics. Our results confirmed this prediction: we found that size tolerance was robustly reshaped in both of these conditions [\(Figure 2](#page-4-0)), and the magnitude of reshaping was similar to that seen with eye-movement-contingent reshaping of IT position tolerance [\(Li and DiCarlo, 2008,](#page-13-0) [Figure 5\)](#page-8-0). Third, we asked if experience with temporal contiguous image statistics could not only break existing IT tolerance, but could also build new tolerance. Again, our results confirmed this prediction: we found that experience with incorrect statistics can build incorrect tolerance [\(Figure 6\)](#page-9-0) and that experience with correct statistics can build correct tolerance (Figure 7). Finally, we found that this tolerance learning is temporally

object-identity-preserving image transformations. Our strategy was to use experience manipulations of temporally contiguous image statistics to look for changes in IT neuronal tolerance that are predicted by this hypothetical learning mechanism. Here we tested three key predictions that were not answered by previous work [\(Li and DiCarlo, 2008\)](#page-13-0). First, we asked if these experience manipulations predictably reshaped the size tolerance of IT neurons. Our results strongly confirmed this prediction: we found that the change in size tolerance was large  $(\sim 5$ spikes/s,  $\sim$ 25% IT selectivity change per hour of exposure) and grew gradually stronger with increasing visual experience. Second, we asked if this tolerance reshaping was induced under visual experience that mimics the common size-toleranceasymmetric and spreads beyond the specifically experienced images ([Figure 8,](#page-11-0) medium size), results that have implications for underlying mechanisms (see below).

Given these results, it is now highly likely that our previously reported results on eye-movement-contingent tolerance learning ([Li and DiCarlo, 2008](#page-13-0)) were only one instance of a general tolerance learning mechanism. Taken together, our two studies show that unsupervised, temporally contiguous experience can reshape and build at least two types of IT tolerance, and that they can do so under a wide range of spatiotemporal regimes encountered during natural visual exploration. In sum, we speculate that these studies are both pointing to the same general learning mechanism that builds adult IT tolerance,

<span id="page-11-0"></span>

Figure 8. Experiment III Exposure Design and Key Results (A) *Exposure Phase* design (*top*, same format as in [Figure 1B](#page-2-0)) and example object images used (*bottom*).

(B) Mean  $\pm$  SEM selectivity change,  $\Delta s(P - N)$ , among the temporally leading images, the nonexposed images at the medium object size  $(3^{\circ})$ , and the temporally lagging images.  $\Delta s(P - N)$  was normalized to show selectivity change per 800 exposure events.  $p = 0.038$ , two-tailed t test.

and we have previously termed this mechanism ''unsupervised temporal slowness learning'' ([Li and DiCarlo, 2008](#page-13-0)).

Our suggestion that UTL is a general tolerance learning mechanism is supported by a number of empirical commonalities between the size tolerance learning here and our previously reported position tolerance learning [\(Li and DiCarlo, 2008\)](#page-13-0). (1) Object specificity: the experience-induced changes in IT size tolerance and position tolerance have at least some specificity for the exposed object. (2) Learning induction (driving force): in both studies, the magnitude of learning depended on the initial selectivity of the temporally adjacent images (medium object size here, foveal position in the position tolerance study), which is consistent with the idea that the initial selectivity may provide at least part of the driving force for the learning. (3) Time course of learning expression: learning increased with increasing amount of experience and changed the initial part of IT response (100 ms after stimulus onset). (4) Response change of learning expression: in both studies, the IT selectivity change arose from a response

decrease to the preferred object (P) and a response increase to the less preferred object (N). (5) Effect size: our different experience manipulations here as well as our previous position manipulation revealed a similar effect magnitude  $(\sim 5$  spikes/s per 400 swap exposures). More specifically, when measured as learning magnitude per exposure event, size tolerance learning was slightly smaller than that found for position tolerance learning [\(Figure 5\)](#page-8-0), and when considered as learning magnitude per unit time, the results of all three experiments were nearly identical [\(Figure S8\)](#page-13-0). However, we note that our data cannot cleanly deconfound exposure amount from exposure time.

#### Relation to Previous Literature

Previous psychophysical studies have shown that human object perception depends on the statistics of visual experience (e.g., [Brady and Oliva, 2008; Fiser and Aslin, 2001; Turk-Browne](#page-13-0) [et al., 2005\)](#page-13-0). Several studies have also shown that manipulating the spatiotemporal contiguity statistics of visual experience can alter the tolerance of human object perception ([Cox et al., 2005;](#page-13-0) Wallis et al., 2009; Wallis and Bülthoff, 2001). In particular, an earlier study ([Cox et al., 2005\)](#page-13-0) showed that the same type of experience manipulation deployed here (experience of different object images across position change) produces increased confusion of object identities across position—a result that qualitatively mirrors the neuronal results reported here and in our previous neuronal study [\(Li and DiCarlo, 2008\)](#page-13-0). Thus, the available psychophysical data suggest that UTL has perceptual consequences. However, this remains an open empirical question (see ''Limitations and Future Direction'' subsection).

Previous neurophysiological investigations in the monkey ventral visual stream showed that IT and perirhinal neurons could learn to give similar responses to temporally nearby stimuli when instructed by reward (i.e., so-called ''paired associate'' learning; [Messinger et al., 2001; Miyashita, 1988; Sakai and Miyashita,](#page-14-0) [1991\)](#page-14-0), or sometimes, even in the absence of reward [\(Erickson](#page-13-0) [and Desimone, 1999\)](#page-13-0). Though these studies were motivated in the context of visual memory ([Miyashita, 1993\)](#page-14-0) and used visual presentation rates of seconds or more, it was recognized that the same associational learning across time might also be used to learn invariant visual features for object recognition (e.g., [Foldiak, 1991; Stryker, 1991; Wallis, 1998; Wiskott and Sej](#page-13-0)[nowski, 2002](#page-13-0)). Our studies provide a direct test of these ideas by showing that temporally contiguous experience with object images can specifically reshape the size and position tolerance of IT neurons' selectivity among visual objects. This is consistent with the hypothesis that the ventral visual stream relies on a temporal contiguity strategy to learn its tolerant object representations in the first place. Our results also demonstrate that UTL is somewhat specific to the experienced objects' images (i.e., object, size, position specificity) and operates over natural, very fast time scales (hundreds of ms, faster than those previously reported) in a largely unsupervised manner. This suggests that, during natural visual exploration, the visual system can leverage an enormous amount of visual experience to construct its object invariance.

Computational models of the ventral visual stream have put forms of the temporal contiguity hypothesis to test, and have shown that learning to extract slowly varying features across

<span id="page-12-0"></span>time can produce tolerant feature representations with units that mimic the basic response properties of ventral stream neurons [\(Masquelier et al., 2007; Masquelier and Thorpe, 2007; Sprekeler](#page-14-0) [et al., 2007; Wallis and Rolls, 1997; Wiskott and Sejnowski, 2002;](#page-14-0) [Wyss et al., 2006\)](#page-14-0). These models can be implemented using variants of Hebbian-like learning rules ([Masquelier and Thorpe, 2007;](#page-14-0) [Sprekeler and Gerstner, 2009; Sprekeler et al., 2007; Wallis and](#page-14-0) [Rolls, 1997\)](#page-14-0). The time course and task independence of UTL reported here is consistent with synaptic plasticity [\(Markram](#page-14-0) [et al., 1997; Rolls et al., 1989](#page-14-0)), and the temporal asymmetry in learning magnitude [\(Figure 8\)](#page-11-0) constrains the possible underlying mechanisms. While the experimental approach used here may seem to imply that experience with all possible images of each object is necessary for UTL to build an invariant IT object representation, this is not believed to be true in a full computational model of the ventral stream. For example, V1 complex cells that encode edges may learn position tolerance that ultimately supports the invariant encoding of many objects. Our observation of partial spread of tolerance learning to nonexperienced images [\(Figure 8](#page-11-0)) is consistent with this idea. In particular, at each level of the ventral stream, afferent input likely reflects tolerance already constructed for simpler features at the previous level (e.g., in the context of this study, some IT afferents may respond to an object's image at both the medium size and the swap size). Thus any modification of the swap-size-image-afferents would result in a partial generalization of the learning beyond the specifically experienced images.

#### Limitations and Future Direction

Because the change in object selectivity was *expressed* in the earliest part of the IT response after learning [\(Figure S5A](#page-13-0)), even while the animal was performing tasks unrelated to the object identity, this rules out any simple attentional account of the effect. However, our data do not rule out the possibility that attention or other top down signals may be required to mediate the learning during the *Exposure Phase*. These potential topdown signals could include nonspecific reward, attentional, and arousal signals. Indeed, psychophysical evidence ([Seitz](#page-14-0) [et al., 2009; Shibata et al., 2009](#page-14-0)) and physiological evidence [\(Baker et al., 2002; Freedman and Assad, 2006; Froemke](#page-13-0) [et al., 2007; Goard and Dan, 2009; Law and Gold, 2008\)](#page-13-0) both suggest that reward is an important factor that can modulate or gate learning. We also cannot rule out the possibility that the attentional or the arousal system may be required for the learning to occur. In our work, we sought to engage the subjects in natural exploration during the *Exposure Phases* under the assumption that visual arousal may be important for ongoing learning, even though we deployed the manipulation during the brief periods of fixation during that exploration. Future experiments in which we systematically control these variables will shed light on these questions, and will help expose the circuits that underlie UTL.

Although the UTL phenomenology induced by our experiments was a very specific change in IT neuronal selectivity, the magnitude of this learning effect was quite large when expressed in units of spikes per second [\(Figure 5](#page-8-0):  $\sim$ 5 spikes/s,  $\sim$ 25% change in IT selectivity per hour of exposure). This is comparable to or larger than other important neuronal phenomenology (e.g., attention, [Maunsell and Cook, 2002\)](#page-14-0). However, because this effect size was evaluated from the multiunit signal, without knowledge of how many neurons we are recording from, this effect size should be interpreted with caution. Furthermore, connecting this neuronal phenomenology (i.e., change in IT image selectivity) to the larger problem of size or position tolerance at the level of the IT population or the animal's behavior is not straightforward. Quantitatively linking a neuronal effect size to behavioral effect size requires a more complete understanding of how that neuronal representation is read out to support behavior, and large effects in confusion of object identities in individual IT neurons may or may not correspond to large confusions of object identities in perception. Such questions are the target of our ongoing and future monkey studies in which one has simultaneous measures of the neuronal learning and the animal's behaviors (modeled after those such as [Britten et al.,](#page-13-0) [1992; Cook and Maunsell, 2002](#page-13-0)).

The rapid and unsupervised nature of UTL gives us new experimental access to understand how cortical object representations are actively maintained by the sensory environment. However, it also calls for further characterization of the time course of this learning to inform our understanding of the stability of ventral stream object representations in the face of constantly available, natural visual experience. This sets the stage for future studies on how the ventral visual stream assembles its neuronal representations at multiple cortical processing levels, particularly during early postnatal visual development, so as to achieve remarkably powerful adult object representation.

#### EXPERIMENTAL PROCEDURES

#### Animals and Surgery

Aseptic surgery was performed on two male Rhesus monkeys (8 and 6 kg) to implant a head post and a scleral search coil. After brief behavioral training (1–3 months), a second surgery was performed to place a recording chamber to reach the anterior half of the temporal lobe. All animal procedures were performed in accordance with National Institute of Health guidelines and the Massachusetts Institute of Technology Committee on Animal Care.

#### General Design

On each experimental day, we recorded from a single IT multiunit site for 2–3 hr. During that time, the animal was provided with altered visual experience in *Exposure Phases* and we made repeated measurements of the IT site's selectivity during *Test Phases* ([Figure 1\)](#page-2-0). The study consisted of three separate experiments (Experiments I, II, and III), which differed from each other only in the *Exposure Phase* design (described below). We focused on one pair of objects (swap objects) that the IT site was selective for (preferred object P, and nonpreferred object N, chosen using a prescreening procedure; see [Supplemental Experimental Procedures\)](#page-13-0).

## Experiment I

Objects (P and N at 1.5°, 4.5°, or 9°) appeared at random positions on a gray computer screen and animals naturally looked to the objects. The image of the just-foveated object was replaced by an image of the other object at a different size (swap exposure event, [Figure 1](#page-2-0)A) or an image of the same object at a different size (non-swap exposure event, [Figure 1A](#page-2-0)). The image change was initiated 100 ms after foveation and was instantaneous [\(Figure 2](#page-4-0), top). We used a fully symmetric design illustrated graphically in [Figure 1](#page-2-0)B. This experience manipulation temporally linked pairs of object images [\(Figure 1A](#page-2-0) shows one such link) and each link could go in both directions [\(Figure 1](#page-2-0)B shows full design example). For each IT site, we always deployed the swap manipulation at one particular size (referred to as the swap size:  $1.5^{\circ}$  or  $9^{\circ}$ , prechosen, strictly alternated between sites), keeping the other size as the exposureequalized control (referred to as the non-swap size).

#### <span id="page-13-0"></span>Experiment II

All design parameters were identical to *Experiment I* except that the image changes were smooth across time [\(Figure 2,](#page-4-0) bottom). The image change sequence started immediately after the animal had foveated the image and the entire sequence lasted for 200 ms (Figure S2). Identity-changing morph lines were only achievable on the silhouette shapes. Only Monkey 2 was tested in Experiment II (given the stimulus class assignment).

#### Experiment III

We used an asymmetric design that is illustrated graphically in [Figure 8](#page-11-0)A: for each IT site, we only gave the animals experience of image changes in one direction  $(1.5^{\circ} \rightarrow 4.5^{\circ}$  or vice versa, prechosen, strictly alternated between sites). The timing of the image change was identical to that in Experiment I.

Another pair of control objects (P' and N', not shown in the *Exposure Phase*) was also used to probe the IT site's responses in the *Test Phase*. The selectivity among the control objects served as a measure of recording stability (below). In each *Test Phase,* the swap and control objects were tested at three sizes (Experiments I and II: 1.5°, 4.5°, 9°; Experiment III: 1.5°, 3°, 4.5°) by presenting them briefly (100 ms) on the animals' center of gaze (50–60 repetitions, randomized) during orthogonal behavioral tasks in which object identity and size were irrelevant. See Supplemental Experimental Procedures for details of the task design and behavioral monitoring.

#### Neuronal Assays

We recorded MUA from the anterior region of IT using standard single microelectrode methods. Our previous study on IT position tolerance learning showed that we could uncover the same learning in both single-unit activity and MUA with comparable effect size (Li and DiCarlo, 2008), so here, we only recorded MUA to maximize recording time. Over a series of recording days, we sampled across IT and sites selected for all our primary analyses were required to be selective among object P and N (ANOVA, object  $\times$  sizes, p < 0.05 for ''object'' main effect or interaction) and pass a stability criterion (n = 27 for Experiment I; 15 for Experiment II; 31 for Experiment III). We verified that the key result is robust to the choice of the stability criteria (Figure S4). See Supplemental Experimental Procedures for details of the recording procedures and site selections.

#### Data Analyses

All the analyses and statistical tests were done in MATLAB (Mathworks, Natick, MA) with either custom-written scripts or standard statistical packages. The IT response to each image was computed from the spike count in a 150 ms time window (100–250 ms poststimulus onset, data from *Test Phases* only). Neuronal selectivity was computed as the response difference in units of spikes/s between images of object P and N at different object sizes. To avoid any bias in this estimate of selectivity, for each IT site we define the labels P (preferred) and N by using a portion of the pre-exposure data to determine these labels, and the remaining data to compute the selectivity values reported in the text (Supplemental Experimental Procedures). In cases where neuronal response data was normalized and combined ([Figures 6 and 7\)](#page-9-0), each site's response from each *Test Phase* was normalized to its mean response to all object images in that *Test Phase*. The key results were evaluated statistically using a combination of t tests and interaction tests (Supplemental Experimental Procedures). For analyses presented in [Figure 4,](#page-7-0) we extracted clear single units from the waveform data of each *Test Phase* using a PCA-based spike sorting algorithm (Supplemental Experimental Procedures).

#### SUPPLEMENTAL INFORMATION

Supplemental Information for this article includes nine figures and Supplemental Experimental Procedures and can be found with this article online at [doi:10.1016/j.neuron.2010.08.029.](http://dx.doi.org/doi:10.1016/j.neuron.2010.08.029)

#### ACKNOWLEDGMENTS

We thank Professors T. Poggio, N. Kanwisher, and E. Miller and the members of our laboratory for valuable discussion and comment on this work. We also thank J. Deutsch, B. Andken, and Dr. R. Marini for technical support. This work

was supported by the NIH (grant R01-EY014970 and its ARRA supplement to J.J.D., NRSA 1F31EY020057 to N.L.) and The McKnight Endowment Fund for Neuroscience.

Accepted: August 5, 2010 Published: September 22, 2010

#### REFERENCES

Afraz, S.R., Kiani, R., and Esteky, H. (2006). Microstimulation of inferotemporal cortex influences face categorization. Nature *442*, 692–695.

Baker, C.I., Behrmann, M., and Olson, C.R. (2002). Impact of learning on representation of parts and wholes in monkey inferotemporal cortex. Nat. Neurosci. *5*, 1210–1216.

Brady, T.F., and Oliva, A. (2008). Statistical learning using real-world scenes: extracting categorical regularities without conscious intent. Psychol. Sci. *19*, 678–685.

Brincat, S.L., and Connor, C.E. (2004). Underlying principles of visual shape selectivity in posterior inferotemporal cortex. Nat. Neurosci. *7*, 880–886.

Britten, K.H., Shadlen, M.N., Newsome, W.T., and Movshon, J.A. (1992). The analysis of visual motion: a comparison of neuronal and psychophysical performance. J. Neurosci. *12*, 4745–4765.

Cook, E.P., and Maunsell, J.H.R. (2002). Attentional modulation of behavioral performance and neuronal responses in middle temporal and ventral intraparietal areas of macaque monkey. J. Neurosci. *22*, 1994–2004.

Cox, D.D., Meier, P., Oertelt, N., and DiCarlo, J.J. (2005). 'Breaking' positioninvariant object recognition. Nat. Neurosci. *8*, 1145–1147.

DiCarlo, J.J., and Maunsell, J.H.R. (2003). Anterior inferotemporal neurons of monkeys engaged in object recognition can be highly sensitive to object retinal position. J. Neurophysiol. *89*, 3264–3278.

Erickson, C.A., and Desimone, R. (1999). Responses of macaque perirhinal neurons during and after visual stimulus association learning. J. Neurosci. *19*, 10404–10416.

Fiser, J., and Aslin, R.N. (2001). Unsupervised statistical learning of higherorder spatial structures from visual scenes. Psychol. Sci. *12*, 499–504.

Foldiak, P. (1991). Learning invariance from transformation sequences. Neural Comput. *3*, 194–200.

Freedman, D.J., and Assad, J.A. (2006). Experience-dependent representation of visual categories in parietal cortex. Nature *443*, 85–88.

Froemke, R.C., Merzenich, M.M., and Schreiner, C.E. (2007). A synaptic memory trace for cortical receptive field plasticity. Nature *450*, 425–429.

Gerstner, W., Kempter, R., van Hemmen, J.L., and Wagner, H. (1996). A neuronal learning rule for sub-millisecond temporal coding. Nature *383*, 76–81.

Goard, M., and Dan, Y. (2009). Basal forebrain activation enhances cortical coding of natural scenes. Nat. Neurosci. *12*, 1444–1449.

Hung, C.P., Kreiman, G., Poggio, T., and DiCarlo, J.J. (2005). Fast readout of object identity from macaque inferior temporal cortex. Science *310*, 863–866.

Ito, M., Tamura, H., Fujita, I., and Tanaka, K. (1995). Size and position invariance of neuronal responses in monkey inferotemporal cortex. J. Neurophysiol. *73*, 218–226.

Kobatake, E., Wang, G., and Tanaka, K. (1998). Effects of shape-discrimination training on the selectivity of inferotemporal cells in adult monkeys. J. Neurophysiol. *80*, 324–330.

Kreiman, G., Hung, C.P., Kraskov, A., Quiroga, R.Q., Poggio, T., and DiCarlo, J.J. (2006). Object selectivity of local field potentials and spikes in the macaque inferior temporal cortex. Neuron *49*, 433–445.

Law, C.T., and Gold, J.I. (2008). Neural correlates of perceptual learning in a sensory-motor, but not a sensory, cortical area. Nat. Neurosci. *11*, 505–513. Li, N., and DiCarlo, J.J. (2008). Unsupervised natural experience rapidly alters invariant object representation in visual cortex. Science *321*, 1502–1507.

<span id="page-14-0"></span>Li, N., Cox, D.D., Zoccolan, D., and DiCarlo, J.J. (2009). What response properties do individual neurons need to underlie position and clutter ''invariant'' object recognition? J. Neurophysiol. *102*, 360–376.

Logothetis, N.K., and Sheinberg, D.L. (1996). Visual object recognition. Annu. Rev. Neurosci. *19*, 577–621.

Markram, H., Lübke, J., Frotscher, M., and Sakmann, B. (1997). Regulation of synaptic efficacy by coincidence of postsynaptic APs and EPSPs. Science *275*, 213–215.

Masquelier, T., and Thorpe, S.J. (2007). Unsupervised learning of visual features through spike timing dependent plasticity. PLoS Comput. Biol. *3*, e31.

Masquelier, T., Serre, T., Thorpe, S.J., and Poggio, T. (2007). Learning complex cell invariance from natural video: a plausibility proof. CBCL Paper (Cambridge, MA: Massachusetts Institute of Technology).

Maunsell, J.H.R., and Cook, E.P. (2002). The role of attention in visual processing. Philos. Trans. R. Soc. Lond. B Biol. Sci. *357*, 1063–1072.

Meliza, C.D., and Dan, Y. (2006). Receptive-field modification in rat visual cortex induced by paired visual stimulation and single-cell spiking. Neuron *49*, 183–189.

Messinger, A., Squire, L.R., Zola, S.M., and Albright, T.D. (2001). Neuronal representations of stimulus associations develop in the temporal lobe during learning. Proc. Natl. Acad. Sci. USA *98*, 12239–12244.

Miyashita, Y. (1988). Neuronal correlate of visual associative long-term memory in the primate temporal cortex. Nature *335*, 817–820.

Miyashita, Y. (1993). Inferior temporal cortex: where visual perception meets memory. Annu. Rev. Neurosci. *16*, 245–263.

Morrison, A., Diesmann, M., and Gerstner, W. (2008). Phenomenological models of synaptic plasticity based on spike timing. Biol. Cybern. *98*, 459–478.

Op De Beeck, H., and Vogels, R. (2000). Spatial sensitivity of macaque inferior temporal neurons. J. Comp. Neurol. *426*, 505–518.

Rolls, E.T., Baylis, G.C., Hasselmo, M.E., and Nalwa, V. (1989). The effect of learning on the face selective responses of neurons in the cortex in the superior temporal sulcus of the monkey. Exp. Brain Res. *76*, 153–164.

Sakai, K., and Miyashita, Y. (1991). Neural organization for the long-term memory of paired associates. Nature *354*, 152–155.

Seitz, A.R., Kim, D., and Watanabe, T. (2009). Rewards evoke learning of unconsciously processed visual stimuli in adult humans. Neuron *61*, 700–707.

Shibata, K., Yamagishi, N., Ishii, S., and Kawato, M. (2009). Boosting perceptual learning by fake feedback. Vision Res. *49*, 2574–2585.

Sigala, N., Gabbiani, F., and Logothetis, N.K. (2002). Visual categorization and object representation in monkeys and humans. J. Cogn. Neurosci. *14*, 187–198.

Sprekeler, H., and Gerstner, W. (2009). Robust learning of position invariant visual representations with OFF responses (Salt Lake City: In COSYNE).

Sprekeler, H., Michaelis, C., and Wiskott, L. (2007). Slowness: an objective for spike-timing-dependent plasticity? PLoS Comput. Biol. *3*, e112.

Stryker, M.P. (1991). Neurobiology. Temporal associations. Nature *354*, 108–109.

Tanaka, K. (1996). Inferotemporal cortex and object vision. Annu. Rev. Neurosci. *19*, 109–139.

Turk-Browne, N.B., Jungé, J., and Scholl, B.J. (2005). The automaticity of visual statistical learning. J. Exp. Psychol. Gen. *134*, 552–564.

Vogels, R., and Orban, G.A. (1996). Coding of stimulus invariances by inferior temporal neurons. Prog. Brain Res. *112*, 195–211.

Wallis, G. (1998). Spatio-temporal influences at the neural level of object recognition. Network *9*, 265–278.

Wallis, G., and Bülthoff, H.H. (2001). Effects of temporal association on recognition memory. Proc. Natl. Acad. Sci. USA *98*, 4800–4804.

Wallis, G., and Rolls, E.T. (1997). Invariant face and object recognition in the visual system. Prog. Neurobiol. *51*, 167–194.

Wallis, G., Backus, B.T., Langer, M., Huebner, G., and Bülthoff, H. (2009). Learning illumination- and orientation-invariant representations of objects through temporal association. J. Vis. *9*, 6.

Wiskott, L., and Sejnowski, T.J. (2002). Slow feature analysis: unsupervised learning of invariances. Neural Comput. *14*, 715–770.

Wyss, R., König, P., and Verschure, P.F. (2006). A model of the ventral visual system based on temporal stability and local memory. PLoS Biol. *4*, e120.