

# Perceptual learning modifies the functional specializations of visual cortical areas

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Training can improve performance of perceptual tasks. This phenomenon, known as perceptual learning, is strongest for the trained task and stimulus, leading to a widely accepted assumption that the associated neuronal plasticity is restricted to brain circuits that mediate performance of the trained task. Nevertheless, learning does transfer to other tasks and stimuli, implying the presence of more widespread plasticity. Here, we trained human subjects to discriminate the direction of coherent motion stimuli. The behavioral learning effect substantially transferred to noisy motion stimuli. We used transcranial magnetic stimulation (TMS) and functional magnetic resonance imaging (fMRI) to investigate the neural mechanisms underlying the transfer of learning. The TMS experiment revealed dissociable, causal contributions of V3A (one of the visual areas in the extrastriate visual cortex) and MT+ (middle temporal/medial superior temporal cortex) to coherent and noisy motion processing. Surprisingly, the contribution of MT+ to noisy motion processing was replaced by V3A after perceptual training. The fMRI experiment complemented and corroborated the TMS finding. Multivariate pattern analysis showed that, before training, among visual cortical areas, coherent and noisy motion was decoded most accurately in V3A and MT+, respectively. After training, both kinds of motion were decoded most accurately in V3A. Our findings demonstrate that the effects of perceptual learning extend far beyond the retuning of specific neural populations for the trained stimuli. Learning could dramatically modify the inherent functional specializations of visual cortical areas and dynamically reweight their contributions to perceptual decisions based on their representational qualities. These neural changes might serve as the neural substrate for the transfer of perceptual learning.

perceptual learning | motion | ps choph sics | transcranial magnetic stimulation | functional magnetic resonance imaging

Perceptual learning, an enduring improvement in the performance of a sensory task resulting from practice, has been widely used as a model to study experience-dependent cortical plasticity in adults (1). However, at present, there is no consensus on the nature of the neural mechanisms underlying this type of learning. Perceptual learning is often specific to the physical properties of the trained stimulus, leading to the hypothesis that the underlying neural changes occur in sensory coding areas (2). Electrophysiological and brain imaging studies have shown that visual perceptual learning alters neural response properties in primary visual cortex (3, 4) and extrastriate areas including V4 (5) and MT+ (middle temporal/medial superior temporal cortex) (6), as well as object selective areas in the inferior temporal cortex (7, 8). An alternative hypothesis proposes that perceptual learning is mediated by downstream cortical areas that are responsible for attentional allocation and/or decision-making, such as the intraparietal sulcus (IPS) and anterior cingulate cortex (9, 10).

Learning is most beneficial when it enables generalized improvements in performance with other tasks and stimuli. Although specificity is one of the hallmarks of perceptual learning, transfer of learning to untrained stimuli and tasks does occur, to a greater or lesser extent (2). For example, visual perceptual learning of an orientation task involving clear displays (a Gabor patch) also improved performance of an orientation task involving noisy displays (a Gabor patch embedded in a random-noise mask) (11). Transfer of perceptual learning to untrained tasks indicates that neuronal plasticity accompanying perceptual learning is not restricted to brain circuits that mediate performance of the trained task, and perceptual training may lead to more widespread and profound plasticity than we previously believed. However, this issue has rarely been investigated. Almost all studies concerned with the neural basis of perceptual learning have used the same task and stimuli for training and testing. One exception is a study conducted by Chowdhury and DeAngelis (12). It is known that learning of fine depth discrimination in a clear display can transfer to coarse depth discrimination in a noisy display (13). Chowdhury and DeAngelis (12) examined the effect of fine depth discrimination training on the causal contribution of macaque MT to coarse depth discrimination. MT activity was essential for coarse depth discrimination before training. However, after training, inactivation of MT had no effect on coarse depth discrimination. This result is striking, but the neural substrate of learning transfer was not revealed.

Here, we performed a transcranial magnetic stimulation (TMS) experiment and a functional magnetic resonance imaging (fMRI) experiment, seeking to identify the neural mechanisms involved in the transfer of learning from coherent motion (i.e., a motion stimulus containing 100% signal) to a task involving noisy motion (i.e., a motion stimulus containing only 40% signal and 60% noise:40% coherent motion). By testing with stimuli other than the

## **Significance**

Using transcranial magnetic stimulation and functional magnetic resonance imaging techniques, we demonstrate here that the transfer of perceptual learning from a task involving coherent motion to a task involving noisy motion can induce a functional substitution of V3A (one of the visual areas in the extrastriate visual cortex) for MT+ (middle temporal/medial superior temporal cortex) to process noisy motion. This finding suggests that perceptual learning in visually normal adults shapes the functional architecture of the brain in a much more pronounced way than previously believed. The effects of perceptual learning extend far beyond the retuning of specific neural populations that mediate performance of the trained task. Learning could dramatically modify the inherent functional specializations of visual cortical areas and dynamically reweight their contributions to perceptual decisions based on their representational qualities.

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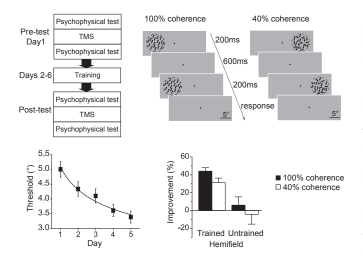
trained stimulus, we uncovered much more profound functional changes in the brain than expected. Before training, V3A and MT+ were the dominant areas for the processing of coherent and noisy motion, respectively. Learning modified their inherent functional specializations, whereby V3A superseded MT+ as the dominant area for the processing of noisy motion after training. This change in functional specialization involving key areas within the cortical motion processing network served as the neural substrate for the transfer of motion perceptual learning.

## Results

**Perceptual Learning of Motion Direction Discrimination.** In our first experiment, we used TMS to identify the causal contributions of V3A and MT+ to coherent and noisy motion processing before and after training. We focused on V3A and MT+ because they are both pivotal areas in the cortical network that supports motion perception (14). Furthermore, both V3A and MT+ are bilateral, which allowed us to train one visual hemifield and left the other hemifield untrained. The experiment consisted of three phases: pretraining test (Pre), motion direction discrimination training, and posttraining test (Post) (Fig. 14).

Psychophysical tests and TMS were performed on the days before (Pre) and after (Post) training. Motion direction discrimination thresholds were measured for each combination of stimulus type (100% coherent: the trained stimulus; 40% coherent: the untrained stimulus) and hemifield (trained and untrained) before and after TMS. TMS was delivered using an offline continuous theta burst stimulation (cTBS) protocol. cTBS induces cortical suppression for up to 60 min (15), which was enough time for all subjects to complete the motion direction discrimination threshold measures. Subjects were randomly assigned to receive TMS of V3A (n = 10) or MT+ (n = 10). Only the hemisphere that was contralateral to the trained hemifield was stimulated.

During training, subjects completed five daily motion direction discrimination training sessions. On each trial, two 100% coherent



random dot kinematograms (RDKs) with slightly different directions were presented sequentially at 9° eccentricity in one visual hemifield (left or right). Within a two-alternative forced-choice task, subjects judged the change in direction from the first to the second RDK (clockwise or counter clockwise) (Fig. 1*B*). A QUEST staircase was used to adaptively control the angular size of the change in direction within each trial and provided an estimate of each subject's 75% correct discrimination threshold.

Similar to the original finding by Ball and Sekuler (16), subjects' discrimination thresholds gradually decreased throughout training (Fig. 1C). The perceptual learning effect was quantified as the percentage change in performance from the pre-TMS psychophysical measures made at Pre to the pre-TMS measures made at Post (Fig. 1D). In the trained hemifield, training led to a significant decrease in discrimination threshold for both the trained stimulus [44%; t(19) = 11.46; P < 0.001] and the untrained stimulus [31%; t(19) = 5.95; P < 0.001]. The transfer from the trained to the untrained stimulus was substantial (71%, the percentage threshold decrease for the untrained stimulus/the percentage threshold decrease for the trained stimulus  $\times 100\%$ ). However, little learning occurred in the untrained hemifield for either stimulus [both t(19) < 0.66; P > 0.05]. Note that none of the learning effects differed significantly between the V3A and MT+ stimulation groups [all t(18) < 1.11; P > 0.05].

A Double Dissociation Between the Causal Contributions of V3A and MT+ to Motion Processing Before Training. Before training, we found a double dissociation between the effects of TMS delivered to V3A and MT+. For each hemifield (trained and untrained) and each stimulation group (V3A and MT+), subjects' motion discrimination thresholds were subjected to a two-way repeated-measures ANOVA with TMS (pre-TMS and post-TMS) and stimulus coherence level (100% and 40%) as within-subject factors (Fig. 2). Because the discrimination task with the 40% coherent RDK was much more difficult than that with the 100% coherent RDK, all the main effects of stimulus coherence level were significant. Therefore, here we focused on the main effects of TMS and the interactions between TMS and stimulus coherence level.

In the trained hemifield (contralateral to the hemisphere that received TMS), for the V3A stimulation group (Fig. 2A), the main effect of TMS [F(1,9) = 4.57; P = 0.06] and the interaction [(F(1,9) = 9.70; P < 0.05] were (marginally) significant. Paired t tests showed that after TMS, performance was impaired and discrimination thresholds were significantly elevated for the 100% coherent stimulus [t(9) = 3.30; P < 0.01]. However, performance for the 40% coherent stimulus was unaffected by TMS [t(9) = 1.29; P > 0.05]. For the MT+ stimulation group (Fig. 2B), we found the opposite pattern: The main effect of TMS and the interaction were significant [both F(1,9) > 10.32; P < 0.05]. After stimulation, discrimination thresholds were significantly elevated for the 40% coherent stimulus [t(9) = 3.71; P < 0.01), but not for the 100% coherent stimulus [t(9) = 2.24; P > 0.05). These results demonstrated that V3A stimulation specifically impaired the processing of 100% coherent motion, whereas MT+ stimulation specifically impaired the processing of 40% coherent motion. This effect was highly specific to the trained hemifield. In particular, there was no significant main effect of TMS or interaction for either the V3A or MT+ stimulation group in the untrained hemifield [ipsilateral to the hemisphere that received TMS; all F(1,9) < 0.94; P > 0.05; Fig. 2 C and D].

Training Changes the Causal Contributions of V3A and MT+ to Motion Processing. The same statistical analysis used for the pretraining data was applied to the posttraining data. In the trained hemifield, for the V3A stimulation group (Fig. 3*A*), the main effect of TMS and the interaction between TMS and stimulus coherence level were significant [both F(1,9) > 23.56; P < 0.01]. After TMS, subjects' discrimination thresholds were significantly elevated for both the 100% and 40% coherent stimuli [both t(9) > 3.14; P < 0.05]. For the MT+ stimulation group (Fig. 3*B*), the main effect of TMS and the interaction were not significant [both F(1,9) < 3.27; P > 0.05]. These results demonstrated that, after training, TMS of V3A disrupted motion processing not only for the 100% coherent stimulus but also for the 40% coherent stimulus. Surprisingly, TMS of MT+ no longer had any effect on task performance for the 40% coherent stimulus, which was in sharp contrast to the pronounced TMS effect for this stimulus before training.

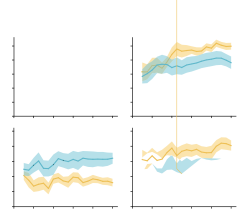
In the untrained hemifield, for the V3A stimulation group (Fig. 3*C*), the interaction was not significant [F(1,9) = 0.07; P > 0.05], but the main effect of TMS was significant [F(1,9) = 13.08; P < 0.01]. After TMS, subjects' discrimination thresholds decreased for the 100% coherent stimulus [t(9) = 3.58; P < 0.01]. This facilitation might reflect a TMS-induced disinhibition of contralateral cortical activity (17), which will be a topic for future investigation. For the MT+ stimulation group (Fig. 3*D*), the main effect of TMS and the interaction were not significant [both F(1,9) < 0.77; P > 0.05].

The TMS experiment demonstrated that before training, V3A and MT+ played causal and dissociable roles in the processing of the 100% and 40% coherent motion stimuli, respectively. In-triguingly, after training, the role of MT+ was replaced by V3A. A possible explanation for this phenomenon is that, before training,

decoders to classify two orthogonal directions and used decoding accuracy to quantify the representation quality. We reasoned that if training could improve the neural representations of the motion stimuli (especially in the trained direction), as suggested by the TMS and psychophysical results, it was possible that decoding accuracies for the orthogonal directions could be improved by training. Similar approaches have been used previously (19–21).

Before training, a repeated-measures ANOVA revealed a significant main effect of stimulus coherence level and a significant interaction between stimulus coherence level and area [V3A and MT+; both F(1,11) > 7.871 P < 0.05; Fig. 4*A*, *Left*). For the 100% coherent motion, the decoding accuracy in V3A was higher than that in MT+ [t(11) = 2.49; P < 0.05], and both were above chance level [both t(11) > 3.75; P < 0.01]. For the 40% coherent motion, only the decoding accuracy in MT+ was above chance level [t(11) = 2.52; P < 0.01], and it was significantly higher than that in V3A [t(11) = 3.19; P < 0.01].

After training, the decoding accuracies in V3A increased for both the 100% and 40% coherent motion [both t(11) > 3.09; P < 0.01]. ANOVA showed that the main effects of stimulus coherence level and area were significant [both F(1,11) > 11.32; P < 0.01; Fig. 4*A*, *Right*]. Furthermore, in stark contrast to the pretraining result, the decoding accuracies in V3A were higher than those in MT+,



not only for the 100% coherent motion [t(11) = 2.85; P < 0.05] but also for the 40% coherent motion [t(11) = 3.51; P < 0.01]. Therefore, the classification abilities of these two areas before and after training were in accordance with their dissociable contributions to the 100% and 40% coherent motion processing revealed in the TMS experiment, supporting our hypothesis.

It should be pointed out that the decoding result did not depend on the number of selected voxels. For V3A and MT+, we selected 20–200 responsive voxels and performed the decoding analysis. The decoding performance generally improved as the voxel number increased. ANOVAs with factors of area (V3A and MT+) and voxel number (160–200) revealed significant main effects of area when conducted separately on data corresponding to each combination of stimulus coherence level (100% and 40%) and training [pretraining and posttraining; Fig. 4B; all F(1,11) > 4.92; P < 0.05].

In addition to V3A and MT+, we also investigated how training changed decoding accuracy in other visual cortical areas (Fig. 4C). For the 100% coherent motion, V3A had the highest decoding accuracy before and after training [paired t tests between V3A and other areas, all t(11) > 2.21; P < 0.05]. In addition to V3A and MT+, the decoding accuracies in V2 and V3 were also significantly above chance level before training [both t(11) > 2.26; P < 0.05]. Notably, only the decoding accuracy in V3A increased significantly after training [t(11) = 5.99; P < 0.01]. For the 40% coherent motion, MT+ was the only area with decoding accuracy that was significantly above chance level before training [t(11) = 2.52; P < 0.05]. However, after training, decoding accuracy in V3A increased dramatically [t(11) = 7.01; P < 0.01], allowing V3A to surpass MT+ and become the area with the highest decoding accuracy [paired t tests between V3A and other ROIs, all t(11) > 2.15; P < 0.05]. Taken together, these results suggest that decision-making areas in the brain rely on the visual area with the best decoding performance for the task at hand, and crucially, that this process is adaptive, whereby training-induced changes in decoding performance across visual areas are reflected in decision-making.

Correlations Among Psychophysical, TMS, and fMRI Effects. To evaluate further the role of V3A in processing the 40% coherent motion after training, we calculated the correlation coefficients between the psychophysical and TMS/fMRI measures for the 40% coherent stimulus across individual subjects (Fig. 5). The coefficient between the behavioral learning effect and the posttraining TMS effect [(post-TMS threshold – pre-TMS threshold)/pre-TMS threshold  $\times$  100%] at V3A was 0.76 (P < 0.05), and the coefficient between the behavioral learning effect and the decoding accuracy change in V3A was 0.62 (P < 0.05), demonstrating a close relationship among the psychophysical, TMS, and fMRI effects. Specifically, the greater the improvement in direction discrimination of 40% coherent motion after training, the greater the involvement of V3A in the task. In addition, the correlation between behavioral learning and decoding accuracy change indicated that the use of orthogonal stimuli within the fMRI experiment allowed for the detection of learning-induced changes in stimulus representation.

**fMRI Linear Discriminant Analysis.** Training improved the decoding performance of V3A with the 40% coherent motion. Responses of V3A voxels to repeatedly presented motion blocks are noisy, fluctuating around a mean value. From the perspective of signal detection theory, there are two strategies to increase the signal-to-noise ratio to improve decoding performance: increasing the distance between the mean values of responses to the trained and the orthogonal directions, and decreasing the response fluctuations along the direction orthogonal to the decision line that separates the responses to the two directions (22). Here we asked which strategy V3A adopted during training.

Linear discriminant analysis (LDA) was used to project the multivoxel response patterns onto a linear discriminant dimension by weighting each voxel's response to maximize the ratio of the between-direction (trained direction vs. orthogonal direction) variance to the within-direction variance. Using this method, we characterized the distributions of the two response patterns in the direction orthogonal to the decision line. In V3A, training reduced the overlap between the patterns evoked by the trained and the orthogonal directions for both the 100% and 40% coherent stimuli (Fig. 6A). We fitted the projected patterns with two Gaussians and compared the signal distance (i.e., the distance between the two Gaussians) and the noise fluctuation (i.e., the variance of the Gaussians) before and after learning. After training, we found significant increases in signal distance at both coherence levels [both t(11) > 2.49; P < 0.05], but no change in noise fluctuation [both t(11) < 1.39; P > 0.05; Fig. 6C]. In MT+, no change occurred in either signal distance or noise fluctuation (Fig. 6 B and D). Notably, the signal distance in V3A at the 40% coherence level, which was almost zero before training [t(11) = 1.57; P > 0.05], surpassed that in MT+ [t (11) = 3.09; P < 0.01] after training. These results confirmed the findings from the decoding analysis and demonstrated that perceptual training increased the pattern distance between the trained and the untrained (orthogonal) directions, rather than reducing the noise fluctuation of neural responses to the two directions.

# Discussion

Whether functional differences exist between V3A and MT+ has been a long-standing question in visual neuroscience. Most previous studies have found that V3A and MT+ exhibit similar functional properties when processing motion (23, 24). In contrast, Vaina and colleagues (25, 26) provided neuropsychological evidence indicating that V3A and MT+ are dominant in local and global motion processing, respectively. Recently, we found that perceptual training with 100% coherent motion increased the neural selectivity in V3A (21). We also have data showing that training with 40% coherent motion increased the neural selectivity in MT+. Together with the results in the current study, these findings point to dissociable roles of V3A and MT+ in coherent and noisy motion processing. In a coherent motion stimulus, the local motion direction of individual dots is the same as the global direction of the stimulus. The specialization of V3A in coherent motion processing might be a result of its greater capacity to process local motion signals than MT+, which is underpinned by its relatively small receptive field sizes and narrow tuning curves for motion direction (27, 28). In a noisy motion stimulus, only some dots move in the global direction, whereas others move in random directions and can be treated as noise. The MT+ specialization for noisy motion processing is believed to be a result of spatial pooling of local motion, which averages out motion noise to reveal the global motion direction (29).

In this study, a substantial transfer of learning occurred from coherent motion to noisy motion, consistent with other studies demonstrating that learning transferred from stimuli without noise to those with noise (11, 13). We speculate that the transfer reported here is a result of an improved representation of the trained motion direction within V3A combined with an increased resilience to the noise present in the noisy motion stimulus. It has been suggested that local motion processing is a primary limitation for global motion sensitivity (30), and that perceptual learning of global motion tasks reflects changes in local motion processing (31). Because 40%of the dots (i.e., the signal dots) in the noisy motion stimuli traveled in the trained motion direction, training with the coherent motion stimuli could enhance the ability of the visual system to use the direction information provided by these signal dots. At the neural level, training with coherent motion resulted in an improvement of local motion representation in V3A. This improvement was characterized by an increase in the pattern distance between the trained direction and the orthogonal directions, which may have made the representation of the trained direction more resistant to the noise present in the noisy motion stimulus. Together, these changes enabled V3A to outperform MT+

V3A were weighted more heavily than those from any other visual cortical area for both kinds of motion. The popular reweighting theory of perceptual learning argues that perceptual learning is implemented by adjusting the weights between basic visual channels and decision-making areas. The visual channels are assumed to lie either within a single cortical area or across multiple cortical areas (34). In the context of the reweighting theory, our results suggest that reweighting can occur between different cortical areas for optimal decision-making. However, it is currently unknown whether training-induced changes in the relative contribution of V3A and MT+ to motion processing were associated with changes in connection "weights" between motion processing areas and decision-making areas, as assumed by the reweighting model. In our study, fMRI slices did not cover IPS. Therefore, we were not able to measure the connection weight changes. This issue should be investigated in the future.

Most perceptual learning studies trained and tested on the same task and stimuli, and assumed that the neural plasticity that accompanies learning is restricted to areas that mediate performance of the trained task. The functional substitution of V3A for MT+ in noisy motion processing induced by coherent motion training challenges this view and approach. Previously, studies of functional substitution or reorganization have mostly been restricted to subjects with chronic sensory disorders. For example, the "visual" cortex of blind individuals is active during tactile or auditory tasks (35, 36). In the area of perceptual learning, two studies attempted to investigate the functional substitution issue. Chowdhury and DeAngelis (12) found that fine depth discrimination training eliminated the causal contribution of MT to coarse depth discrimination. However, the

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visual areas that took responsibility for coarse depth discrimination after training were not identified. Using TMS, Chang et al. (37) demonstrated that perceptual training shifts the limits on perception from the posterior parietal cortex to the lateral occipital cortex (see also ref. 38). Here, we propose that perceptual learning in visually normal adults shapes the functional architecture of the brain in a much more pronounced way than previously believed. Importantly, this extensive cortical plasticity is only revealed when subjects are tested on untrained tasks and stimuli. In the future, investigating the neural mechanisms underpinning the transfer of perceptual learning will not only remarkably advance our understanding of the nature of brain plasticity but also help us develop effective rehabilitation protocols that may result in training-related functional improvements generalizing to everyday tasks through learning transfer.

### **Materials and Methods**

**Subjects.** T ent subjects (11 female, 20–27 old) participated in the TMS e periment, and 12 subjects (five female, 20–25 old) participated in the fMRI e periment. The ere naive to the purpose of the e periment and had never participated in a perceptual learning e periment before. All subjects ere right-handed and had normal or corrected-to-normal vision. The had no kno n neurological or visual disorders. The gave ritten, informed consent in accordance ith the procedures and protocols approved b the human subject revie committee of Peking Universit. Detailed methods are provided in *SI Materials and Methods*.

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