Current Biology

Perceptual Learning of Contrast Detection in the Human Lateral Geniculate Nucleus

Highlights

- Contrast learning shows specificity to the trained eye and visual hemifield
- Contrast learning boosts the activity of the M layers of the LGN
- Perceptual learning in human adults can occur as early as at the thalamic level

Authors

Qinlin Yu, Peng Zhang, Jiang Qiu, Fang Fang

Correspondence ffang@pku.edu.cn

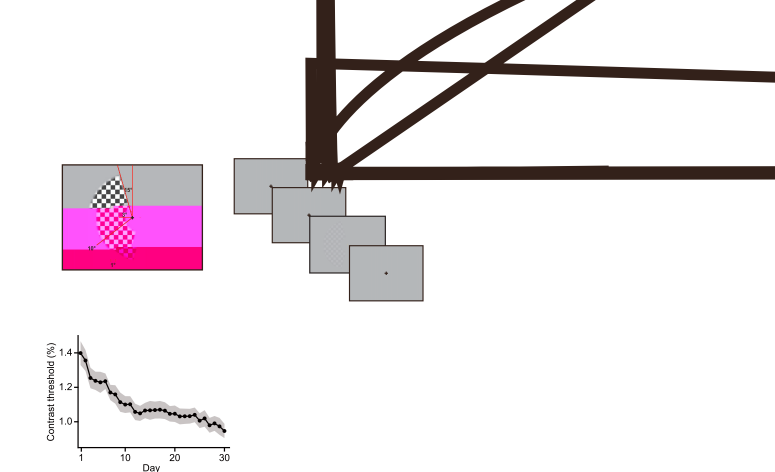
In Brief

Yu et al. reveal that perceptual learning of contrast detection leads to an eye- and hemifield-specific neural response increase to low contrast in the M layers of the LGN and suggest that visual training can induce plasticity in subcortical nuclei.









voxels was further constrained by the anatomical locations of the LGN based on high-resolution T1 images. On the T1 images in Figure 2A, which shows the LGN from a representative subject, the LGN appeared darker relative to surrounding brain tissues. The LGN is the thalamic component in the retinocortical projection and has been traditionally viewed as a passive relay station for retinal signals on their way to the primary visual cortex, or V1 [13]. This view has been challenged recently. There is growing evidence from human fMRI and monkey neurophysiology studies that neural responses in the LGN are influenced by perceptual and cognitive tasks (see [14] for a review).

Using the counterphase flickering checkerboard stimuli, we measured fMRI contrast response functions in the ROIs at three contrast levels (6%, 24%, and 96%). During scanning, subjects performed a demanding task to detect the color change of the fixation point (Figure 2B). Therefore, the peripheral checkerboard stimuli were task irrelevant. The fMRI contrast response functions are shown in Figure 2C. For each ROI and each test condition, blood-oxygen-level dependent (BOLD) amplitudes were submitted to a repeated measures ANOVA, with training (preand post-training) and contrast (6%, 24%, and 96%) as withinsubject factors. The main effects of contrast were significant (LGN: all Fs(2, 38) > 82.82, p < 0.001; V1: all Fs(2, 38) > 142.77, p < 0.001; V2: all Fs(2, 38) > 168.93, p < 0.001; V3: all Fs(2, 38) > 122.98, p < 0.001, Bonferroni corrected). The BOLD responses increased with contrast. The main effects of training were not significant (LGN: all Fs(2, 19) < 3.195, p > 0.36; V1: all Fs(2, 19) < 0.378, p = 1; V2: all Fs(2, 19) < 0.445, p = 1; V3: all Fs(2, 19) < 1.217, p = 1). The interaction effect between training and contrast was only significant in the THTE condition in the LGN (THTE: F(2, 38) = 6.839, p < 0.05; UHTE: F(2, 38) = 0.567, p = 1; THUE: F(2, 38) = 0.350, p = 1; UHUE: F(2, 38) = 1.408, p = 1, Bonferroni corrected). Furthermore, post hoc t tests showed that the BOLD response after training was significantly

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targets presented at the same retinal location to the untrained eye, which is in line with the eye specificity property of this kind of behavioral learning [5].

Is this LGN response enhancement a long-lasting change, and does it serve as a long-term mechanism of contrast detection learning? One recent study [35] measured the dynamics of subjects' behavioral performance with a texture detection task [5] and their V1 activation over a long time course of perceptual learning. Within the first few weeks of training, V1 activation in a subregion corresponding to the trained location and task performance both increased. However, while the improved performance was maintained 2 weeks after training, the V1 activation decreased to the level observed before training. Similar transient response enhancements were also found in the fusiform face areas immediately after training on a face discrimination task [11]. Both of the studies challenged the role of the transient response enhancements immediately after training in perceptual learning. In the present study, we did not measure brain signals after the post-training test to examine the persistence of the response enhancement to the low contrast. Nevertheless, the significant correlation between the behavioral and neural enhancements provides deterministic evidence for the crucial role of the M layers in the contrast detection learning, at least in the learning effect immediately after training.

Unlike previous studies [28–30], we did not observe traininginduced response increase at the cortical level (i.e., V1). Here are several possible reasons. First, the fMRI measurement is not sensitive enough to detect such small changes (if there are any) that might be also specific to the trained eye and M neurons. In V1–V3, BOLD signals from individual voxels reflect mixed neural signals from left and right eye neurons and from M and P neurons, which could not be separated due to the limit of the current fMRI spatial resolution. Second, subjects were trained for the glutamate receptor agonist to block visual responses in oncenter retinal ganglion cells and found that the inactivation led to a rapid emergence of off-center responses from on-center neurons in the LGN. A signibcant stride we made in the present study is that, without such abnormal visual experience (i.e., eyelid closure or pharmacological inactivation), even regular practice could profoundly change local receptive Þeld properties of the LGN neurons in human adults. Recently, it has been recognized that the LGN and other thalamic structures actively regulate information transmission to the cortex and between cortical areas using various mechanisms, thereby contributing to perception and cognition much more than we previously believed [14, 41]. Exploring the functional plasticity of the subcortical structures induced by training is an important research topic in the future, which is necessary for us to fully understand the adaptive nature of perceptual and cognitive information processing in the brain.

EXPERIMENTAL PROCEDURES

The procedures and protocols used in this study were approved by the human subject review committee of Peking University. Complete procedures can be found in the Supplemental Information.

SUPPLEMENTAL INFORMATION

Supplemental Information includes two Þgures and Supplemental Experimental Procedures and can be found with this article online at http://dx.doi. org/10.1016/j.cub.2016.09.034.

AUTHOR CONTRIBUTIONS

Q.Y. and F.F. designed the study. Q.Y. and J.Q. conducted the experiments. Q.Y., P.Z., and F.F. analyzed the data and wrote the manuscript.

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