Optogenetically Induced Behavioral and Functional Network Changes in Primates

Annelies Gerits,1,2,3 Reza Farivar,1,2 Bruce R. Rosen,1,2 Lawrence W. Wald,1,2,6 Edward S. Boyden,4,5,6 and Wim Vanduffel1,2,3,*

1Athinoula A. Martinos Center for Biomedical Imaging, Massachusetts General Hospital, Charlestown, MA 02129, USA
2Department of Radiology, Harvard Medical School, Boston, MA 02115, USA
3Laboratorium voor Neuro- en Psychofysiologie, KU Leuven Medical School, Campus Gasthuisberg, 3000 Leuven, Belgium
4Media Lab, Synthetic Neurobiology Group
5McGovern Institute, Department of Brain and Cognitive Sciences
6Department of Biological Engineering
Massachusetts Institute of Technology, 77 Massachusetts Avenue, Cambridge, MA 02139, USA

Summary

Optogenetics is currently the state-of-the-art method for causal-oriented brain research. Despite an increasingly large number of invertebrate and rodent studies showing profound electrophysiological and behavioral effects induced by optogenetics [1, 2], only two primate studies have reported modulation of local single-cell activity but with no behavioral effects [3, 4]. Here, we show that optogenetic stimulation of cortical neurons within rhesus monkey arcuate sulcus, during the execution of a visually guided saccade task, evoked significant and reproducible changes in saccade latencies as a function of target position. Moreover, using concurrent optogenetic stimulation and opto-fMRI [5, 6], we observed optogenetically induced changes in fMRI activity in specific functional cortical networks throughout the monkey brain. This is critical information for the advancement of optogenetic primate research models and for initiating the development of optogenetically based cell-specific therapies with which to treat neurological diseases in humans.

Results

We tested whether optogenetic stimulation of neurons in the posterior and anterior bank of the arcuate sulcus alters behavioral performance of monkeys performing a cognitive task. We first identified cortical patches within the arcuate sulcus that were activated by the saccade task using fMRI [7]. Then, using fMRI-guided neuronavigation, we transduced neurons in ventral premotor (F5) and prefrontal cortex (frontal eye fields, FEF) of two monkeys (M1 and M2) with adeno-associated viral vector-chicken β-actin promoter-channelrhodopsin-2-GFP (AAV5-CAG-ChR2-GFP) (Figure 1) and stimulated these cortical patches with blue light. We stimulated with two optic fibers simultaneously to increase the number of neurons that were activated within a functional network relative to previous monkey optogenetic studies [3, 4]. A visually guided saccade task with peripheral go cues, requiring divided attention across multiple peripheral visual field locations [8], was used to test for changes in behavioral performance (Figure 2, see Experimental Procedures for details). In short, this task started with a fixation period after which a green saccade target appeared either on the left or the right side of the fixation point together with four possible white go cues. The latter were presented either on the vertical meridian (lower or upper visual field) or on the horizontal meridian (left or right). A luminance change of these peripheral go cues indicated to the animal to make a saccade toward the green target in order to receive a reward. We compared behavioral performance during visually guided saccade trials with and without optical stimulation at cortical sites in the posterior and anterior bank of the arcuate sulcus that were previously injected with AAV-ChR2 (stimulation sessions) (Figure 1). In addition, we also compared the monkey’s behavioral performance between stimulated and nonstimulated trials of control sessions during which the optical fibers were inserted in prefrontal sites that were not previously injected with the viral vector construct—but otherwise using exactly the same experimental paradigm as during the stimulation sessions (see Experimental Procedures). Finally, brain-wide optogenetically induced functional changes were probed using opto-fMRI [5, 6], analogous to previous experiments combining monkey fMRI with electrical stimulation [9].

Behavioral Results

In total, we performed five stimulation sessions and four control sessions in each animal (excluding the lentiviral experiments in M1; see Experimental Procedures). Optogenetic stimulation experiments targeting the AAV sites started 130 and 122 days after injection of the viral vectors in M1 and M2, respectively. We obtained 32 and 31 runs from M1 and 52 and 57 runs from M2 during stimulation and control sessions, respectively. For the statistical analysis, we included only trials from runs in which the monkeys were able to correctly complete >60% of the trials within the time windows as listed in the Experimental Procedures. Furthermore, we analyzed only trials with saccadic latencies >50 ms and <400 ms, as well as trials with latencies >100 ms and <400 ms (which yielded the same results). This resulted in a total of 1,961 and 1,680 analyzed trials from M1 and M2, respectively. M1 successfully completed 83.9% ± 1.5% and 81.1% ± 1.9% of the fixation trials during the stimulation and control sessions, respectively. For M2, these figures were 85.6% ± 1.0% and 81.6% ± 1.3%. No statistical difference in fixation performance was observed between stimulated and nonstimulated fixation conditions (t test; p > 0.5 for both animals).

Saccade Latencies

In both animals, there was a significant interaction between target position and stimulation (two-way ANOVA, F = 7.92, p = 0.005 for M1 and F = 5.76; p = 0.016 for M2), but not during the control sessions (F = 2.87; p = 0.1 for M1 and F = 3.7; p = 0.06 for M2) (Figure 3). In M1, saccade latencies in stimulated versus nonstimulated trials were shorter for ipsilateral (median of 14 ms, t test, p = 0.016) but not contralateral targets (median of −4.42 ms, t test, p = 0.27). For M2, faster saccade latencies...
were observed for targets presented in both hemifields, although the effect was larger for contralateral saccades. Median differences in saccade latencies measured 20 and 28 ms in M2 for ipsi- and contralateral targets, respectively (t test, \( p < 10 \times 10^{-6} \) for both targets). A significant interaction for saccade latencies was observed between cue position and stimulation for M1 (F = 6.3, \( p = 0.01 \)), but not for M2 (F = 1.3, \( p = 0.25 \)). One could argue that the fixed order of conditions may have contributed to the observed saccade latency effects. This is very unlikely, however, because we used exactly the same order of conditions during the stimulation and control sessions, and there was a highly significant interaction between the type of session (stimulation or control) and trial type (stimulation or no stimulation) (F = 54.9; \( p = 1.5 \times 10^{-13} \) [group data], see also Figure 3).

**Target Detection Accuracy and Saccade Metrics**

Accuracy for target detection by the two monkeys was unaltered by light stimulation both during stimulation (median correctly executed trials across conditions = 77%); with less than 3% difference in accuracy between stimulation and control trials, t test, \( p > 0.43 \) for both animals), and control sessions (p > 0.24 for both animals). Finally, no changes were observed between saccade endpoints of stimulation and control trials (t test, \( p > 0.35 \) for both monkeys), nor were there any differences in the number of saccades (t test, \( p > 0.11 \) for either monkey) or eye blinks (t test, \( p > 0.24 \) for both monkeys) during stimulation and control trials. Thus, activation of light-sensitive depolarizing ion channels in transduced neurons of the arcuate sulcus results in faster saccadic reaction times as a function of target position.

**Functional MRI Results**

In a case where focal perturbation of brain activity is causally linked to behavior, one expects not only local but also more extensive network-specific changes in function. To test this prediction, we measured fMRI signals during epochs with and without optogenetic stimulation [5, 6] of the foci in the anterior or posterior bank of the arcuate sulcus (Figure 1). Significant optogenetically induced activations were observed near the site of stimulation in both monkeys (white arrows, Figure 4A). Moreover, clearly different, largely bilateral, functional networks were activated by stimulation of each of these two injection sites. Optical stimulation of control sites, not previously injected with AAV-ChR2, revealed virtually no local or remote fMRI activations (Figure 4A).

Although slightly different locations were stimulated during the various sessions, we observed reproducible activation patterns across sessions of the same subject. For example, Figure 4B shows optogenetically induced activity in several visual areas across two different sessions in M1 (e.g., MSTd [yellow arrow], MSTv [blue arrow], V4 [red arrow], and peripheral V1 [green arrow]). In M2, to take another example, we observed reproducible patterns of optogenetic induced activity across three sessions in the post central sulcus (orange arrow, Figure 4C).

**Discussion**

Optogenetics is a recently developed method to increase and decrease the activities of specific neurons with high temporal resolution in order to relate their function to behavior, and to make causal inferences about the role they play, both within local microcircuitry and across macroscopic functional brain networks [1, 2]. Critical with regard to future translational purposes, optogenetics presents a promising method for manipulating activity in distinct cell types that are relevant to specific neurological diseases [10, 11]. This novel method has been shown to evoke clear behavioral and neuronal effects in invertebrates [12] and rodents [13, 14]. However, despite some evidence that optogenetics can also alter single unit activity in macaques [3, 4], no study has reported optogenetically induced behavioral or functional network changes in primates—which is of critical importance for translational purposes. Possibly the number of optically stimulated neurons needs to be substantially greater in primates than in animals with simpler brains to evoke any behavioral effects. Alternatively, previous primate optogenetic studies may have failed to reveal psychophysical or motor changes, because behavioral tests used thus far have been insufficiently sensitive.

In this study, we attempted to overcome these potential issues by injecting a viral vector construct under the control of a cell-type nonspecific CAG promoter in order to stimulate...
In M1, two different constructs were injected, which both contained the reporter gene GFP and the membrane channel ChR2, but under the control of different promoters. The first was a lentiviral (LVV) construct consisting of a specific Ca²⁺/calmodulin-dependent protein kinase II (CaMKII) promoter to target only excitatory cells, as used previously in monkeys by Han and colleagues [3]. This LVV construct was injected into the posterior and anterior bank of the left arcuate sulcus. The second adeno-associated viral vector construct (AAV) serotype 5 under control of the nonspecific CAG promoter [4] was injected into the right frontal cortex (Figure 1). No behavioral or functional MRI effects were obtained after stimulation of the targeted LVV sites in the left arcuate sulcus of M1 (five stimulation sessions). Therefore, we injected only the AAV-construction in the right frontal cortex of M2 and we focused exclusively on results obtained after stimulating the sites injected with the AAV construct.

All injections were made through a grid placed in an implanted chamber over the right arcuate sulcus (Crist Instrument) (Figure 1). The three-dimensional coordinates of the injection targets were determined using BrainInsight. Neuronavigation and were based on prior fMRI activations for the visually guided saccade task. We injected only in cortical sites activated by the saccade task relative to the fixation task (i.e., in cortical tissue corresponding to voxels with p < 0.001, uncorrected). In the target sites, a total of 1 µl (per site) of the viral vector solution (1.1 × 10⁵ genome copies per µl) was slowly delivered with a 5 µl Hamilton syringe at a rate of 0.2 µl every 2 min at two cortical depths per site.

Behavioral Tasks
In the visually guided saccade task with multiple possible go cues, the animals had to maintain fixation in a virtual window of maximum 2 × 2 degrees around a small red spot in the center of the display for a fixed duration of 700–1,400 ms. Thereafter, a single green saccade target and four possible white go cues appeared (Figure 2). Target and go cues were equal in size (0.14°) and luminance (6 cd/m²). The saccade target appeared either on the left or right side of the fixation point at 10 degrees eccentricity along the horizontal meridian. The possible go cues appeared at 8 degrees eccentricity, either on the vertical meridian (lower or upper visual field) or on the horizontal meridian (left or right). After a variable delay of 700 to 1,400 ms, one of the randomly selected go cues would turn gray. This change was the go signal indicating to the animal to make a saccade toward the green target. Note that all four peripheral go cues had to be attended by the animal to detect the go signal as quickly as possible. The monkey was rewarded for making a saccade toward the green target within 700 ms after the go signal and for maintaining fixation within a 3–4 degree window around the target for 200 ms. To encourage rapid responses, we varied reward size as an exponential function of reaction time (RT) between 150 and 700 ms after the go signal. The time between target onset and the go signal was a random variable drawn from a unimodal Weibull distribution delayed by 500 ms [15].

Saccades were detected using a computer algorithm that searched first for eye position before and after the high-velocity gaze shift. The time point of saccade initiation and termination were then defined as the beginning and end of the monotonic change in eye position before and after the high-velocity gaze shift.

During the fixation task, the monkeys were rewarded with juice for fixating upon a red fixation point in the center of an otherwise empty screen within a 2 × 2 degree window. The average duration of the fixation trials was the same as the saccade trials. All procedures were approved by MGH’s Subcommittee on Research Animal Care (Protocol # 2010N000165; PI Vanduffel), and are in accordance with NIH guidelines for the care and use of laboratory animals.
Figure 3. Optogenetically Induced Changes in Behavior
Saccadic reaction times of optogenetically stimulated (blue) and nonstimulated (red) trials during optical stimulation (left) and control (right) sessions.
The first two bars show the median change in saccadic latencies of the two monkeys for the stimulation sessions. The two groups of four bars show the individual data per monkey and per target location (C = contralateral relative to stimulated hemisphere, I = ipsilateral). The last two bars show the median saccadic latencies acquired during the control sessions (single-subject data are indicated by the diamond and circles) (error bars represent SEM across trials; t test, * = p < 0.05).

Stimulation Paradigm
Stimulation sessions utilized two fibers simultaneously in the posterior and anterior banks of the arcuate sulcus, at sites where viral injections had been made. Light pulses (40 Hz, 8 ms pulses) were administered with two 473 nm blue lasers (Shanghai Laser & Optics Century), coupled to two optical fibers 200 μm in diameter. The light intensity at the fiber tip ranged from 80 to 300 mW/mm², as measured before and after each session with a PM 100D power meter (Thorlabs). We alternated between a fixation epoch without stimulation, a saccade epoch with stimulation, a saccade epoch without stimulation, and a fixation epoch with stimulation. This sequence was repeated 8 times within a given 512 s run. In each 16 s task epoch, both monkeys were able to complete 4 ± 1 saccade trials.

During control sessions, we stimulated with fibers inserted at nearby prefrontal locations (3.5–6.5 mm removed from the previous sites, mainly in the anterior-middle portion of the superior branch of the arcuate sulcus and in the principal sulcus) but without prior viral vector injections. We used exactly the same experimental paradigm as described above for the stimulation sessions and monkeys performed the same number of trials per task epoch as in the stimulation sessions (4 ± 1 trials/epoch).

fMRI Data Acquisition
Contrast-agent-enhanced functional images were acquired using a four-channel phased array coil (GRAPPA, acceleration factor 2) on a 3 Tesla TIM-Trio scanner (Siemens) with AC88 gradient coil, using a gradient-echo T2*-weighted echo-planar sequence (62 and 64 sagittal slices for M1 and M2, respectively; 102 x 106 matrix, TR = 2 s, TE = 17 ms, 1 mm isotropic voxels) [7]. We performed single-subject analyses and compared the stimulation condition to nonstimulation (p < 0.001 uncorrected, t > 3.09).

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