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Characterization of the Functional MRI Response Temporal Linearity via Optical Control of Neocortical Pyramidal Neurons

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The blood oxygenation level-dependent (BOLD) signal serves as the basis for human functional MRI (fMRI). Knowledge of the properties of the BOLD signal, such as how linear its response is to sensory stimuli, is essential for the design and interpretation of fMRI experiments. Here, we combined the cell-type and site-specific causal control provided by optogenetics and fMRI (opto-fMRI) in mice to test the linearity of BOLD signals driven by locally induced excitatory activity. We employed high-resolution mouse fMRI at 9.4 tesla to measure the BOLD response, and extracellular electrophysiological recordings to measure the effects of stimulation on single unit, multunit, and local field potential activity. Optically driven stimulation of layer V neocortical pyramidal neurons resulted in a positive local BOLD response at the stimulated site. Consistent with a linear transform model, this locally driven BOLD response summed in response to closely spaced trains of stimulation. These properties were equivalent to responses generated through the multisympathetic method of driving neocortical activity by tactile sensory stimulation, and paralleled changes in electrophysiological measures. These results illustrate the potential of the opto-fMRI method and reinforce the critical assumption of human functional neuroimaging that—to first approximation—the BOLD response tracks local neural activity levels.

Introduction

The blood oxygenation level-dependent (BOLD) functional MRI (fMRI) signal (Ogawa et al., 1990; Kwong et al., 1992) is widely used to indirectly measure neuronal activity in the human brain. Because the BOLD response measures hemodynamic fluctuations and not underlying electrophysiological signals directly, the relation between neural activity and the BOLD fMRI signal has been studied extensively (Boynton et al., 1996; Dale and Buckner, 1997; Heeger et al., 2000; Miezin et al., 2000; Rees et al., 2000; Logothetis et al., 2001; Mukamel et al., 2005; Viswanathan and Freeman, 2007). With some caveats, most evidence links spike rate and the local field potential (LFP) with the BOLD response in the cortex (for reviews, see Heeger and Ress, 2002; Logothetis, 2008). Several studies have further demonstrated that the BOLD response has a roughly linear relation to underlying spiking activity elicited by sensory stimulation (Boynton et al., 1996; Dale and Buckner, 1997; Miezin et al., 2000; Rees et al., 2000; Logothetis et al., 2001).

We built on these prior studies by characterizing the somatosensory neocortical BOLD response in a mouse model that allows precise temporal control of neural activity, an “opto-fMRI” approach. As in other recent demonstrations (Desai et al., 2011; Lee et al., 2010), we combined activation of the microbial opsin channelrhodopsin-2 (ChR2), a light-sensitive nonselective cation channel (Nagel et al., 2003; Boyden et al., 2005), with high-field functional imaging (9.4 tesla fMRI). We employed the Thy1-ChR2-yellow fluorescent protein (YFP) mouse, in which ChR2 is expressed in layer V pyramidal neurons throughout the neocortex (Arenkiel et al., 2007). By targeting primary somatosensory (SI) cortex, we characterized the properties of the BOLD response as a function of pyramidal neuron spiking and tested the hypothesis that the local excitation of pyramidal neurons by optogenetic means recruits a BOLD response that can be modeled with a linear transform from underlying neural activity.

We tested whether a BOLD response evoked by closely temporally spaced short trains of optically driven pyramidal cell activity would summate linearly. We found that the BOLD...
response closely matched local pyramidal neuron activity and added in an approximately linear fashion. These findings both inform our interpretation of the classic sensory driven BOLD neocortical response and provide a quantitative investigation of the opto-fMRI BOLD signal.

Materials and Methods

Animals. All procedures were conducted in accordance with the National Institutes of Health guidelines and with the approval of the Committee on Animal Care at Massachusetts Institute of Technology. ChR2 transgenic mice (Arenkiel et al., 2007) were purchased from The Jackson Laboratories (line 18, stock no. 007612, strain B6.Cg-Tg(Thy1-COP4/EYFP18GfngI) and subsequently crossed with C57BL/6J mice. Animals used were of either sex. Electrophysiological recordings and fMRI were obtained in separate animals.

Animal anesthesia and surgery. Animals were anesthetized with isoflurane mixed with oxygen for all procedures. During preparation and surgery anesthetia was kept at a level of 1–3%. Immediately before MRI scanning and electrophysiological recording, the anesthetic level was reduced to 0.5–1% expired isoflurane. For the MRI experiments, the craniotomy was restricted to a burr hole in the skull. For the electrophysiological recordings, the dura mater was opened after the craniotomy to allow positioning of the optical fiber and recording electrode.

Light stimulation. Light stimulation was generated by a 473 nm laser (Shanghai Dream Lasers) controlled by a computer. Optical stimulation at 40 Hz frequency and 8 ms pulse duration was delivered via a 200 µm diameter unjacketed optical fiber with 0.48 numerical aperture (Thorlabs) at a range of 320–470 mW mm⁻² irradiance at the tip of the fiber, irradiance that can evoke EMG responses when targeting primary motor cortex (Ayling et al., 2009). The fiber was positioned over SI barrel field (SIBf; from bregma: anteroposterior, −1.5 mm; mediolateral, 2.5 mm) at the cortical surface. We computed the dropout in light irradiance using a Monte Carlo simulation (Chow et al., 2010), with a simulated volume of 4 mm³, 20 µm³ voxels, 5 million photons, scattering coefficient in gray matter at 473 nm of 10 mm⁻¹ (Yaroslavsky, 2002), absorption coefficient of 0.07 mm⁻¹, anisotropy factor of 0.88, and a uniform laser emission pattern over a half-opening angle of 5 degrees, yielding blue light penetration to layer V of mouse cortex with about 90% irradiance loss, chiefly due to scattering of light (Bernstein et al., 2008).

Sensory stimulation. A piezoelectric (PZ) element (5.0 cm x 0.78 cm x 0.18 cm; Noliac) was used to deflect vibrissae. A glass capillary tube was used for extension (1 mm internal diameter and 5 cm long), and a nylon thread was looped through the glass capillary tube. All the posterior vibrissae of the right facial pad were pinched at −1 cm from the vibrissal pad. A 40 Hz sinusoidal wave was used to drive the PZ element with a peak-to-peak motion amplitude calibrated to 347 ± 12 µm along the dorsoventral direction.

MRI procedures. Mice were fixed to an animal bed (Bruker BioSpin MRI) using a standard tooth piece and ear bars. Breathing rate (Small Animal Monitoring Model 1025, SA Instruments) and expired isoflurane (V9004 Capnograph Series, SurgiVet, Smiths Medical PM) were continuously monitored. Imaging was performed on a 9.4 tesla BioSpec 94/20 USR MRI (Bruker BioSpin MRI) that operates at a maximum gradient strength of 675 mT/m, maximum slew rate 4673 T/m/s, and a rise time of 130 µs. A single transmit and receive surface coil consisting of a single copper loop and etched circuit board (20 mm diameter) was placed over the head. The fiber was placed on or ≤50 µm from the dura mater. In a subset of mice, the fiber slightly compressed the underlying neocortex, as was seen in the anatomical MRI. These animals produced data identical to that observed in other preparations, suggesting that it did not appreciably impact local cortical responses. BOLD-contrast signals were acquired using a spin-echo echo-planar sequence (repetition time [TR], 2.5 s; echo time, 12.17 ms; 10 coronal slices, 200 µm x 200 µm x 500 µm). High-resolution T₁- and T₂-weighted anatomical images were recorded using a rapid acquisition process with a relaxation enhancement (RARE) sequence in coronal orien-tations (10 coronal slices, 78 µm x 78 µm x 500 µm). Image reconstruction was carried out using ParaVision 5.0 (Bruker BioSpin MRI). Preprocessing included compensation of slice-dependent time shifts, rigid body correction for motion within and across runs, and smoothing with a full-width half-maximum kernel of 600 µm. For each mouse in the fMRI component of the study, we conducted interleaved “localizer” and “target” runs. The SIBf region of interest (ROI) identified in the localizer runs was subsequently used to test temporal linearity in the target runs. For each localizer run, the images were detrended and percentage signal change was computed by subtracting, from each voxel’s BOLD time series, the temporal average of all off periods for all scans in a session, dividing by the average of the signal across no-stimulation periods and then averaged across all localizer runs. Significant BOLD signal increases were determined by a voxelwise time series correlation of the percentage signal change with the protocol boxcar, delayed by one TR. A family wise error corrected p value of 0.05 was used. To achieve this threshold, only clusters with at least six contiguous voxels and uncorrected p values below 0.005 were considered. The cluster size and p value threshold were determined via Monte Carlo simulations performed in AFNI using AlphaSim (Forman et al., 1995; Xiong et al., 1995).

To estimate the location of activated voxels in SIBf, the EPI volumes were aligned to the T1 structural volumes using the AFNI align_epi_anat.py program. ROIs were identified by overlaying the structural images onto corresponding atlas plates (Paxinos and Franklin, 2001), aligning the anteroposterior coordinates, and scaling the atlas plates along the x- and y-axes until the borders corresponded.

Electrophysiology. A small craniotomy was made over SIBf. Stimulus control and single unit activity (SU) data acquisition were performed using software custom written in LabView (National Instruments) and Matlab (The Mathworks). Glass pipette-pulled electrodes (10–14 MΩ resistance) were filled with saline and lowered under positive pressure while injecting small amplitude current pulses. Cell-attached recording configuration was achieved when a medium resistance seal (≥50 MΩ) was formed and spikes were observed (MultiClamp 700B, Molecular Devices). LFP and multunit activity (MUA) recordings were obtained using 16 contacts (15 µm diameter spaced 100 µm apart) laminar electrodes (Neuronexus Technologies) connected to a Cheetah32 data acquisition system (Neuralynx). Recordings were performed at 0.1–9000 Hz filter settings and separated into LFP and MUA signals post hoc.

Electrophysiology analysis. Cell-attached recording data were bandpass filtered between 600 and 6000 Hz, and spike times were extracted where the potential exceeded 10 standard deviations of the signal calculated across the entire block. The MUA data were extracted from laminar recordings by bandpass filtering the signal between 600 and 6000 Hz, and spike times were extracted where the potential exceeded 4 standard deviations of the signal calculated across the entire block. All firing rate data were binned at 100 ms. The LFP amplitudes were calculated by subtracting any slow drift in the potential (piecewise linear fit to the potential at the beginning of each trial) and averaging the negative of the signal during the entire stimulus presentation. For LFP power, the signal was lowpass filtered at 300 Hz. Spectrograms were computed for each trial using a 250 ms Hamming window and frequencies from 7 to 200 Hz. Values reported represent the average over all frequencies and the entire stimulus presentation time.

Statistical procedures. We tested the data for normality with the Lilliefors test (Lilliefors, 1967). The electrophysiological measures exhibited normal distributions, but the fMRI BOLD data exhibited non-normal distributions (p < 0.005). Therefore, we conducted nonparametric tests throughout (Wilcoxon signed rank and rank-sum two-tail tests). Goodness-of-fit (R²) measures were derived using least-squares regression of the averaged time series from each animal against the predicted response.

Histology. Mice (n = 4; two from the MRI experiment and two from the electrophysiology experiment) were transcardially perfused with 100 ml phosphate buffer (PB) followed by 4% formaldehyde in PB, and brains were postfixed for 18 h at 4°C. Free-floating sections (50 µm) were cut using a freezing microtome (American Optical Model 820) mounted on glass slides with Vectashield (Vector Laboratories) and coverslipped. Spread and labeling efficiency were estimated by examination of 50 µm coronal sections near the SIBf for the presence of YFP using a Zeiss LSM.
Figure 1. Optogenetic fMRI. Neural activation of layer V ChR2-expressing neurons by light stimulation evokes a reliable fMRI response that is equivalent to somatosensory response evoked by vibrissa deflection. 

A. Opto-fMRI experimental design. Atlas (Paxinos and Franklin, 2001) sagittal (i) and coronal (ii) sections depict the placement of the fiber optic (blue line) over primary somatosensory cortex (SI) barrel field. Monte Carlo simulation of laser intensity contours are plotted (iii), demonstrating that light intensity at 750 μm was 10%, power sufficient to induce action potentials in layer V pyramidal neurons in vitro. MI, Primary motor cortex. 

B. Wide-field fluorescence depicts ChR2 expression of primarily layer V pyramidal neurons in a transgenic Thy1-ChR2-YFP mouse. Sagittal slice depicts expression throughout the neocortex, and an atlas section is superimposed on representative histological and anatomical MRI coronal sections, depicting the site of stimulation for a representative animal. 

C. Characteristic expression is shown demonstrating fluorescence in layer V pyramidal neurons (top). Maximum intensity projection of confocal laser scanning microscopy demonstrates ChR2 expression (bottom). Denoted by white triangles are apical dendrites approaching the surface, the cell bodies, and axons. 

D. “Localizer” runs consisted of 16 repetitions of 15 s light-pulses or deflections of the mystacial vibrissae followed by 15 s of no stimulation. A statistical parametric map of a positive BOLD response to light stimulation (p < 0.05 corrected for multiple comparisons) is overlaid on the average fMRI BOLD image of the sequential cross-sections demonstrating the response for the optical drive in a typical animal (“Optogenetic”). A similar BOLD response was observed to vibrissa deflections (“Sensory”).
5 Pascal (Carl Zeiss light microscopy) in wide-field fluorescence and laser scanning confocal modes.

Results

The SIbf was chosen as the optical stimulation site. Figure 1, A and B, shows typical fiber optic placement. The light irradiance at the tip of the fiber was at least 300 mW mm$^{-2}$, such that at layer V about 10% of the light was present (Fig. 1 Aiiii), a power level that fully activates ChR2 (Boyden et al., 2005). To confirm expression of the ChR2-YFP fusion protein under the control of the Thy1 promoter in the stimulated site, we carried out histological experiments in four mice. Robust ChR2-YFP expression was observed in layer V pyramidal neurons (Fig. 1 B, C) with extensive expression in apical dendrites and cell bodies. The pattern was consistent with previously reported expression in Thy1 promoter-driven lines (Feng et al., 2000; Arenkiel et al., 2007; Ayling et al., 2009).

We first tested whether a BOLD response could be observed in response to optical stimulation in neurons expressing ChR2 and the extent to which optical drive spatially corresponded to sensory driven responses. In each animal participating in the fMRI experiment, 2–8 trials were interleaved with target runs. In each localizer run, either light pulses (eight animals) or vibrissa deflections (four animals) were applied for 15 s followed by 15 s of no stimulation, and this sequence was repeated 16 times. BOLD responses were observed in SIbf for the optical and sensory conditions (see Fig. 1 D for typical optogenetic and sensory responses). Optically driven BOLD responses extended over a volume of 0.65 ± 0.29 mm$^3$ (mean ± SD; eight animals), consistent with the emitted light irradiance. Similarly, sensory driven BOLD responses extended over a volume of 0.60 ± 0.11 mm$^3$ (four animals).

Dale and Buckner (1997) observed that the BOLD response summed in an approximately linear fashion to pairs of brief visual stimuli (1 s) spaced by as little as 2 s. We sought to test the assertion that subtraction of $(n)$ 1 s trains from $(n + 1)$ 1 s trains of activity would yield a response equivalent to a single 1 s train. Employing cell-attached juxtacellular recording from single units (SUA; 5 animals) as well as MUA and LFP recording with 16-channel laminar electrodes (1 animal) we observed an equivalent level of activity by 1 s periods of stimulation in both a single (1 s interval of stimulation) and double (two 1 s intervals spaced by 2 s) trial conditions (Fig. 2). We next extracted the BOLD time series from voxels identified in localizer runs for all animals (Fig. 3). The single and double trial conditions demonstrated an approximately linear increase, where a subtraction of the single trial from the double trial yielded a time series matching the single trial condition (Fig. 3A; six animals). To test the generality of this result, we asked whether this pattern held when multiple 1 s trains were compared. We compared in a separate experiment (two animals) optical drive of five (quintuple) to six (sextuple) 1 s trains. The computed subtraction of these conditions yielded a response similar to that observed for the single 1 s train (Fig. 3B). To provide a bridge to BOLD responses observed in human fMRI, we compared the optical somatosensory-evoked responses to multisynaptic sensory-evoked responses such as those generated by vibrissal deflections. Both the single and double (two animals; Fig. 3C) and quintuple − sextuple (two animals; Fig. 3D) trial conditions exhibited what appeared to be linear increases.

We quantified the temporal linearity of the BOLD response using a variety of metrics. First, we verified that the integrated BOLD response of the $(n + 1)$ trials produced a significantly greater response relative to $(n)$ trials in the optical and sensory driven responses (Wilcoxon signed rank paired two-tailed test; Optogenetic (optical drive): single − double, $p < 0.0001$, and quintuple − sextuple, $p < 0.05$; Sensory (sensory drive): single − double, $p < 0.01$, and quintuple − sextuple, $p < 0.05$). Second, we...
confirmed that the temporal evolution of a single trial response yielded a response equivalent to a double trial by testing the goodness of fit of the double trial to the summation of a single trial and a single trial shifted by 3 s (Fig. 4A, B; Optogenetic: $R^2 = 0.64$, $F_{(1,23)} = 38.94, p < 0.0001$; Sensory: $R^2 = 0.88, F_{(1,7)} = 44.60, p < 0.001$). Third, we computed the ratio between the responses of the $(n + 1)$ and $(n)$ trial conditions (Fig. 4C, D). For the single – double conditions the BOLD response signal was integrated over the 2.5–12.5 s period for the optical and sensory driven responses; for SUA and MUA the total number of spikes was counted, and for LFPs the total power was calculated. The electrophysiological measures yielded a double to single response ratio close to 2: $1.99 \pm 0.02$ (mean $\pm$ SEM) for SUA, $2.05 \pm 0.07$ for MUA, and $2.05 \pm 0.04$ for LFP, and a similar ratio held for the BOLD responses (Optogenetic: $1.97 \pm 0.25$; Sensory: $1.70 \pm 0.18$). Similarly, for the quintuple – sextuple conditions BOLD integrated responses were computed over the 2.5–22.5 s period for the optical and sensory driven responses, yielding a response ratio close to 1.2 (Optogenetic: $1.25 \pm 0.08$; Sensory: $1.25 \pm 0.08$). Finally, we tested whether the linear model holds by computing the goodness of fit of the subtraction of $(n)$ from $(n + 1)$ trial conditions time series to the single trial response. A hemodynamic response function (HRF) was fitted to the single trial condition of the optical and sensory drive separately by a mixture of two Gamma functions ($f_1$ and $f_2$) fitting a delay of the response and undershoot, dispersion of response and undershoot, and ratio of response to undershoot [probability density function of Gamma distribution with shape ($h$) and scale ($l$) parameters; Optogenetic: $f_1(h = 5.55, l = 0.34); f_2(h = 10.42, l = 0.13)$; Sensory: $f_1(h = 5.05; l = 0.16); f_2(h = 8.33; l = 13)$]. The resultant HRF was tested for goodness of fit to the subtracted responses for the single – double and quintuple – sextuple. In the optical stimulation condition, the subtracted responses were well fitted by the HRF (Fig. 4E; single – double: $R^2 = 0.53, F_{(1,29)} = 31.06, p < 0.0001$; quintuple–sextuple: $R^2 = 0.68, F_{(1,9)} = 16.86, p < 0.0005$). In the sensory stimulation condition, only the single – double subtraction resulted in a reliable goodness-of-fit to the HRF (Fig. 4F; single – double: $R^2 = 0.585, F_{(1,9)} = 11.2778, p = 0.01$; quintuple – sextuple: $R^2 = 0.2773, F_{(1,9)} = 3.0046, p = 0.12$). Thus, consistent with spike activity in individual neurons, short trains of activity, when separated in time, linearly summed in the BOLD response. Furthermore, linearity of the hemodynamic response held for optogenetic drive consistently, while it failed for sensory drive in the quintuple – sextuple condition.

**Discussion**

We applied optogenetic stimulation to control the precise timing of neocortical pyramidal neuron activity and tested the spike, LFP, and BOLD signal responses to this stimulation. We demonstrate that when regular spiking pyramidal cells in layer V of the sensory neocortex were driven by stimulation of light-activated cation channels, a hemodynamic response was evoked reliably. We recorded SUA, MUA, and LFP activity to test whether the BOLD response followed the linear transform model. We observed a linear summation of the BOLD response for temporally spaced short trains of optical stimulation that was proportional to the cumulative neural activity recorded as SUA, MUA, and LFP signals.

A core assumption of brain imaging with fMRI using the BOLD-contrast mechanism is that local changes of neural activity elicit a hemodynamic response and that the hemodynamic response is approximately linearly related to the underlying neural
activity (Boynton et al., 1996; Dale and Buckner, 1997; Miezin et al., 2000; Rees et al., 2000; Logothetis et al., 2001; Heeger and Ress, 2002). Our results suggest that—to first approximation—this assertion is supported. Optically driven BOLD responses demonstrated temporal linear summation following closely temporally spaced trains of stimulation. Furthermore, our data indicate that similar analysis assumptions can be applied to opto-fMRI responses generated by direct optical activation of pyramidal neurons. Our findings suggest that a rodent model may be a valid approach to explore what inferences can be made about underlying neural activity from the indirect BOLD response.

Our results using opto-fMRI to probe the BOLD response suggest a general approach to study systematically the relation between local changes in neural activity and the BOLD response recorded by fMRI. The opto-fMRI approach we employed is similar to that taken by Desai et al. (2011) and Lee et al. (2010), who used opto-fMRI to study network connectivity, and activated CaMKIIα-expressing excitatory neurons that were virally transduced. We used the Thy1 transgenic mouse model expressing ChR2, providing an effective system for studying the relation between neural activity and fMRI. Critically, we observed a deviation from linearity using sensory drive when multiple 1 s intervals of stimulation were applied. Failure of linearity can, in principle, result from failure of linearity between the stimulus and neural responses or neural responses and the hemodynamic response. The present results eliminate the latter possibility. By focusing on understanding the local response, we have probed the neural and BOLD responses under conditions typically employed in human imaging, the summation of short duration, closely spaced, event-related signals.

References


