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Alpha-180 spin-echo based line-scanning method for high resolution 2 laminar-specific fMRI 3

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23 ABSTRACT

24 Laminar-specific functional magnetic resonance imaging (fMRI) has been widely used to study circuit-specific 25 neuronal activity by mapping spatiotemporal fMRI response patterns across cortical layers. Hemodynamic 26 responses reflect indirect neuronal activity given limit of spatial and temporal resolution. Previous gradient-echo based line-scanning fMRI (GELINE) method was proposed with high temporal (50 ms) and spatial (50 um) 27 resolution to better characterize the fMRI onset time across cortical layers by employing 2 saturation RF pulses. 28 However, the imperfect RF saturation performance led to poor boundary definition of the reduced region of interest 29 (ROI) and aliasing problems outside of the ROI. Here, we propose α (alpha)-180 spin-echo-based line-scanning 30 fMRI (SELINE) method to resolve this issue by employing a refocusing 180° RF pulse perpendicular to the 31 excitation slice. In contrast to GELINE signals peaked at the superficial layer, we detected varied peaks of laminar-32 33 specific BOLD signals across deeper cortical layers with the SELINE method, indicating the well-defined exclusion of the large drain-vein effect with the spin-echo sequence. Furthermore, we applied the SELINE method with 200 34 35 ms TR to sample the fast hemodynamic changes across cortical layers with a less draining vein effect. In summary, 36 this SELINE method provides a novel acquisition scheme to identify microvascular-sensitive laminar-specific 37 BOLD responses across cortical depth.

38 INTRODUCTION

39 Line-scanning fMRI has been successfully applied to investigate circuit-specific neuronal activity by measuring dynamic hemodynamic responses across cortical layers with high spatiotemporal resolution¹⁻⁹. This is initially 40 originated from Mansfield's line-profile mapping studies in early 1970s^{10,11}. The advantage of the current line-41 scanning fMRI method is to sample cortical layers with ultra-high spatial resolution. Meanwhile, the line-scanning 42 method only acquires a single k-space line per timepoint, enabling an ultrafast sampling rate. This high 43 spatiotemporal laminar fMRI sampling scheme has been being utilized for bottom-up and top-down blood-44 45 oxygenation-level-dependent (BOLD) fMRI mappings in both animal and human fMRI studies. Previously, Yu et al. developed a line-scanning fMRI method to delineate laminar fMRI onset time with distinct laminar-specific 46 neural inputs such as thalamocortical input and corticocortical input in the rat brain with high spatial (50 um) and 47 temporal resolution (50 ms)¹. Line-scanning fMRI has been also combined with optogenetic control to further 48 investigate the temporal features of the fast neural inputs across cortical layers in rodents². Beyond preclinical fMRI 49 50 studies, line-scanning fMRI for human brain mapping demonstrated a good correspondence with BOLD responses of 2D echo planar imaging (EPI) at the same temporal scale (200 ms)¹². This line-scanning fMRI also motivated 51 the cortical depth-dependent diffusion-based fMRI mapping schemes¹³. Lately, the ultra-fast line-scanning fMRI 52 53 with k-t space reshuffling scheme has even provoked some interesting investigation of direct neuronal activity 54 measurements¹⁴.

Typical gradient echo (GRE)-based line-scanning fMRI (GELINE) method needs to dampen signals outside of the 55 region of interest (ROI) to avoid aliasing artifacts along the phase encoding direction^{1,2,4,5,7–9}. Two saturation slices 56 with additional RF exposure are applied for this purpose. However, two issues should be further investigated. One 57 58 is the imperfect elimination of the aliasing artifacts (including inflow effects) due to imperfect RF performance and 59 inhomogeneous B0 field. The other is specific absorption rate (SAR) problem with high duty cycle sequences. Here, we developed α (alpha)-180 line-scanning fMRI method to solve these problems. We modified spin-echo (SE) 60 sequence by altering the refocusing 180° RF pulse perpendicular to the excitation slice^{3,10,11}. This adjustment allows 61 to only highlight a line-profile across the cortical layers without additional saturation RF pulses. In contrast to the 62 GELINE method, SE based line-scanning fMRI (SELINE) method effectively exclude the surface draining vein 63 effects. However, it should be noted that the laminar patterns of BOLD signals in SELINE can still be highly varied 64 across different cortical layers in anesthetized rats. Furthermore, we also shorten TR to 200 ms for the SELINE 65 method to sample the high resolution T2-weighted fMRI signals, demonstrating the feasibility of the fast sampling 66 67 of laminar fMRI with effective ROI selectivity in rodents.

68

69 **RESULTS**

70 Mapping the evoked BOLD fMRI signals with GELINE and SELINE

71 We developed the SELINE method to map laminar-specific BOLD responses across cortical layers at the primary forepaw somatosensory cortex (FP-S1) of anesthetized rats, which can be compared with the conventional GELINE 72 73 method¹. First, unilateral electrical stimulation of left forepaw of rats showed robust BOLD responses in the right 74 FP-S1 using EPI-fMRI method (Fig. 1A). Using the GELINE method, the selected FOV was defined by two 75 saturation slices to avoid aliasing problem along the phase encoding direction (Fig. 1B). In contrast, the same FOV 76 could be selected by applying a refocusing 180° RF pulse perpendicular to the excitation slice with SELINE (Fig. 1E). To compare ROI selectivity between GELINE and SELINE. 2D in-plane images were acquired by turning on 77 78 phase encoding gradient (Fig. 1C and 1F) and 1D profiles were plotted by averaging all readout voxels of the 2D 79 image (Fig. 1D and 1G). Background signals were estimated from the areas outside of the FOV (for details, see the 80 Method section): For GELINE, trial #1) 17.6 %, #2) 51.0 %, and for SELINE: trial #1) 2.3 %, #2) 3.0 %. This result indicated the efficiency of the SELINE method to produce sharper 2D slice profiles and lower background signals. 81

To study the laminar fMRI characteristics of GELINE and SELINE across the cortical layers, we calculated temporal signal-to-noise ratio (tSNR) with 1D line-profiles which were acquired by turning off the phase encoding gradient. The tSNR of SELINE was higher than those of GELINE (**Fig. 1H**). The tSNR graph of SELINE had gradually decreasing trend across the cortical depth while those of GELINE had gradually increasing trend. The difference was likely caused by different TRs (1000 ms vs. 100 ms) and flip angles (90° vs. 50°) of the transceiver surface coil.

As shown in Fig. 1I-L, we demonstrated dynamic BOLD responses across different cortical layers of FP-S1 from 88 the representative trial in individual GELINE (Fig. 1I and 1J) and SELINE (Fig. 1K and 1L) studies. Fig. 1I 89 demonstrated periodic evoked BOLD signals upon left forepaw electrical stimulation with the T2*-weighted 90 GELINE method, showing the dynamic laminar-specific BOLD responses as a function of time peaked around the 91 92 superficial layer in the FP-S1 (4 s on/16 s off for each 20 s epoch, total 32 epochs). Average BOLD time series and laminar-specific BOLD maps illustrated that the peak BOLD response is located at L1, highlighting large draining 93 vein effects at the cortical surface¹⁵⁻²¹ (Fig. 1J). In comparison to GELINE, SELINE also detected robust FP-S1 94 95 BOLD signals across different cortical layers (Fig. 1K), but showed peak BOLD signal located at L4, presenting improved spatial specificity to deeper cortical layers^{15–18,20,21} (Fig. 1L). 96

97 Comparison of the laminar-specific peak BOLD responses in GELINE and SELINE.

98 We further investigated the reproducibility of laminar-specific peak BOLD responses, as well as the variability of 99 laminar-specific BOLD response patterns, between two methods (14 trials from 3 animals). The GELINE method 100 detected peak BOLD signals primarily located at L1, but the peak BOLD signal detected by the SELINE method is 101 much deeper. In animal #3, the ultra-strong BOLD signal detected in the superficial voxel indicates a large draining 102 vein dominating the voxel BOLD signal. A similar BOLD response was also detected by the SELINE method, 103 which may be contributed by potential intravascular effect of which the large draining vein is not negligible in the 104 voxels with only 50 um thickness (Fig. 2G and 2H). Interestingly, the layer-specific BOLD signal varies largely 105 across animals in both GELINE and SELINE maps. Besides the primary peak BOLD in L1 of GELINE, a second 106 peak appeared in L4 in some animal (Fig. 2D). And for SELINE method, the primary peak also varies at L2/3 and L4, which present highly different laminar patterns from GELINE even acquired from the same animal with 107 interleaved trials during experiments. These results have suggested that the profile of laminar-specific BOLD 108 109 signals can vary largely across animals, which may present varied dynamic patterns of BOLD responses due to the 110 altered neurovascular coupling across different cortical layers.

111 Mapping the laminar BOLD responses with a 200 ms SELINE method.

112 We performed BOLD fMRI experiments with 200 ms time of repetition (TR) by applying optimized flip angles based on the Bloch equation^{22,23}. For comparison, we also performed GELINE method in the same anesthetized rat. 113 114 As shown in Fig. 3A-D, we demonstrated the evoked BOLD responses across the cortical layers upon the periodic electric stimulation with the GELINE (Fig. 3A and 3B) and SELINE (Fig. 3C and 3D) methods, showing the 115 116 average BOLD time series and percentage changes peaked at L1 in both GELINE and SELINE. To characterize the laminar-specific BOLD responses, the normalized BOLD signals were plotted across the cortical layers. As shown 117 118 in Fig. 3F, the GELINE method had the steep signal drop from L1 to L2, while the SELINE method had the gradual 119 signal drop across the cortical depth. It indicates that the high temporal SELINE method reduces the large vessel 120 contribution to the BOLD responses by minimizing magnetic susceptibility effects at the superficial layer (i.e., L1). 121 To select an optimized flip angle, the tSNR of different flip angles was plotted (Fig. 3G). Even though the Ernst 122 angle for TR 200 ms was ~150° and had the highest tSNR, the difference of the tSNR change was relatively small 123 in multiple trials with the different flip angles. This result was possibly caused by the long T1 effect (~2200 ms) in SELINE acquisition with a short TR (200 ms)²⁴. As same as the theoretical predictions based on the Bloch equation 124 22,23 , $e^{-TR/T1}$ was almost close to one and thus, the maximum intensity at the Ernst angle wasn't changed much. It 125 was noteworthy that the average tSNR of SELINE was higher at the superficial and middle layers than that of 126 GELINE (Fig. 3E and 3G) due to larger flip angle (100-150° vs. 50°) and longer TR (200 ms vs. 100 ms). In 127 128 summary, these results not only demonstrated less magnetic susceptibility effects at the superficial layer, but also highlighted both tSNR and laminar specificity enhancement in SELINE with high temporal resolution. 129

130 DISCUSSION

In this study, we applied the SELINE method to investigate laminar-specific evoked BOLD responses across cortical layers with high spatial and temporal resolution. The SELINE method has sharper and better ROI-selectivity than the GELINE method, employing the refocusing 180 RF pulse perpendicular to the excitation plane. Our results show that the peak signal of SELINE is spread across the cortical layers while those of GELINE is at the superficial layer^{25,26}By pushing the temporal resolution of SELINE to 200 ms, we also demonstrate the feasibility to map laminar-specific BOLD response with the suppression of the large draining vein effect^{15–18,20,21} in comparison to the GELINE method.

138 Significant effort with high magnetic field fMRI has been made to explore laminar fMRI responses corresponding to distinct information flows (e.g., top-down/bottom-up or feedforward/feedback) at high spatial and temporal 139 scales in both animals and humans. Among these efforts, cortical depth-dependent fMRI, detecting BOLD, cerebral 140 141 blood volume (CBV), and cerebral blood flow (CBF) signals with both SE and GRE methods, has identified 142 hemodynamic regulation, blood volume distribution, circuit-specific laminar responses, and hierarchical information streams across cortical layers in animal^{1,2,5,8,15,16,25-29} and human brains³⁰⁻³⁴. In particular, the high 143 144 resolution CBV-fMRI (based on the VASO mapping scheme) has been used to measure layer-specific directional functional connectivity across human motor cortex and somatosensory and premotor regions³⁰. It should be noted 145 146 that the cortical thickness of human brains is in the range of 1-4 mm, which is highly comparable to that of rodent brains in the range of 1-2 mm³⁵. Given the limited spatial resolution of the high field laminar-fMRI method (~600-147 700 um), the truly counted voxels across different cortical regions are in the single digit number, which could be 148 much better improved by the developed line-scanning fMRI method, as well as with ultra-fast sampling rate. 149

150 Recently, the GRE-based line-scanning BOLD mapping scheme has been implemented to investigate BOLD signals across cortical layers in human fMRI studies^{7,12}. Nevertheless, the required saturation RF pulses of the GELINE 151 method result in high specific absorption rate (SAR) and total RF power limits with short TRs, inducing more 152 complicated aliasing problems. For the SELINE method, the beam-like line-scan projection has been previously 153 154 applied for probing myeloarchitecture across cortical layers in the primary somatosensory cortex (S1) and primary motor cortex (M1) of the human brain¹³ and mapping irreversible and reversible transverse relaxation rates (i.e., R2 155 and R2') in primary visual cortex (V1), S1, and M1 of human brains³⁶. We thus applied this SELINE method to 156 better characterize layer-specific fMRI features across cortical depths at FP-S1 of rodent brains. The SELINE 157 method employed the spin-echo scheme to reduce the large draining vein effect, which can be further distinguished 158 159 from the deeper cortical layer responses given the high spatial resolution (Fig. 1I-L).

As reported in previous studies^{15–17,25,37–39}, GELINE is more sensitive to large veins at the pial surface and has poor 160 161 specificity across different cortical depths, whereas SELINE is less vulnerable to superficial large draining veins 162 and has good sensitivity to microvessel across cortical layers. However, the largely varied laminar patterns of the 163 BOLD responses were observed in both methods (Fig. 2). It may suggest that the varied patterns of laminar-specific 164 BOLD signals pertain on microvascular biases and baseline blood volume distribution across cortical layers ^{40,41}. Whereas the confounding observation of the varied peak profiles of BOLD responses across different cortical layers, 165 these results illustrate the feasibility of the line-scanning method to detect distinct laminar BOLD responses, 166 167 providing a high-resolution mapping scheme when investigating altered neurovascular coupling events across 168 cortical layers.

The limitation of SELINE is the slow sampling rate. We tried to shorten TR by adjusting excitation flip angle. Based on Bloch equation^{22,23}, we have estimated the appropriate angles with a short TR (i.e., 200 ms). Our results show the feasibility of the fast SELINE method which has a good sampling capability capturing dynamic BOLD signals from superficial to deeper layers. For the future work, simultaneous GRE- and SE-type fMRI acquisition can be applied to better characterize laminar-specific fMRI patterns and minimize time dependency of dynamic fMRI responses by employing GRASE⁴²-based line-scanning in rodents as already suggested for the human fMRI mapping⁶.

176 METHODS

177 Animal preparation. The study was performed in accordance with the German Animal Welfare Act (TierSchG) 178 and Animal Welfare Laboratory Animal Ordinance (TierSchVersV). This is in full compliance with the guidelines 179 of the EU Directive on the protection of animals used for scientific purposes (2010/63/EU) and the MGH Guide for 180 the Care and Use of Laboratory Animals. The study was reviewed by the ethics commission (§15 TierSchG) and approved by the state authority (Regierungspräsidium, Tübingen, Baden-Württemberg, Germany) and the MGH 181 Institutional Animal Care and Use Committee (Charlestown, MA, USA). A 12-12 hour on/off lighting cycle was 182 maintained to assure undisturbed circadian rhythm. Food and water were available ad libitum. A total of 4 male 183 184 Sprague–Dawley rats were used in this study.

Anesthesia was first induced in the animal with 5% isoflurane in the chamber. The anesthetized rat was intubated using a tracheal tube and a mechanical ventilator (SAR-830, CWE, USA) was used to ventilate animals throughout the whole experiment. Femoral arterial and venous catheterization was performed with polyethylene tubing for blood sampling, drug administration, and constant blood pressure measurements. After the surgery, isoflurane was switched off, and a bolus of the anesthetic alpha-chloralose (80 mg/kg) was infused intravenously. After the animal was transferred to the MRI scanner, a mixture of alpha-chloralose (26.5 mg/kg/h) and pancuronium (2 mg/kg/h)
was constantly infused to maintain the anesthesia and reduce motion artifacts.

192 EPI fMRI acquisition. All data sets from rats were acquired using a 14.1T/26 cm (Magnex, Oxford) horizontal 193 bore magnet with an Avance III console (Bruker, Ettlingen) and a 12 cm diameter gradient system (100 G/cm, 150 194 µs rising time). A home-made transceiver surface coil with a 10 mm diameter was used on the rat brain. For the 195 functional map of BOLD activation (Fig. 1A), a 3D gradient-echo EPI sequence was acquired with the following parameters: TR/TE 1500/11.5 ms, FOV $1.92 \times 1.92 \times 1.92$ cm³, matrix size $48 \times 48 \times 48$, spatial resolution 0.4×10^{-10} 196 0.4×0.4 mm³. A high order (e.g., 2nd or 3rd order) shimming was applied to reduce the main magnetic field (B0) 197 inhomogeneities at the region-of-interest. For anatomical reference of the activated BOLD map, a RARE sequence 198 199 was applied to acquire 48 coronal images with the same geometry as that of the EPI images. The fMRI design 200 paradigm for each trial comprised 200 dummy scans to reach steady-state, 10 pre-stimulation scans, 3 scans during 201 stimulation, and 12 post-stimulation scans with a total of 8 epochs.

202 GELINE acquisition. GELINE datasets (9 trials of 4 rats) were acquired with a 6-mm diameter home-made transceiver surface coil in anesthetized rats for evoked fMRI. GELINE was applied by using two saturation slices 203 to avoid aliasing artifacts in the reduced field-of-view along the phase encoding (*i.e.*, from left to right) direction 204 (Fig. 1B and 1C). 2D line profiles were acquired to evaluate saturation RF pulses performance (Fig. 1D). Laminar 205 206 fMRI responses were acquired along the frequency-encoding direction (Fig. 1I and 1J). The following acquisition parameters were used: TR/TE 100/12.5 ms, TA 10 min 40 sec, FA 45°, slice thickness 1.2 mm, FOV 6.4 × 3.2 mm², 207 208 and matrix 128×32 . The fMRI design paradigm for each epoch consisted of 1 second pre-stimulation, 4 seconds 209 stimulation, and 15 seconds post-stimulation with a total of 20 seconds. A total of 6400 lines (*i.e.*, 10 m 40 s) in 210 each cortex were acquired every single trial in evoked fMRI. Evoked BOLD activation was identified by performing 211 electrical stimulation to the left forepaw (300 µs duration at 2.5 mA repeated at 3 Hz for 4 seconds).

212 SELINE acquisition. SELINE datasets (18 trials of 4 rats) were acquired in anesthetized rats for evoked fMRI. SELINE was applied by the 180° RF pulse oriented perpendicular to the α° excitation RF pulse as moving the 213 214 refocusing gradient to phase encoding gradient in order to obtain high spatial resolution without reduced FOV aliasing problem along the phase encoding (*i.e.*, from left to right) direction (Fig. 1E and 1F). 2D line profiles were 215 216 also acquired to evaluate the refocusing RF pulses performance (Fig. 1G). Laminar fMRI responses were acquired 217 along the frequency-encoding direction (Fig. 1K and 1L). The following acquisition parameters were used: TR/TE/FA 1000/20 ms/90°, 200/10 ms/ 100° or 130° or 150°, TA 10 min 40 sec, slice thickness 1.2 mm, FOV 3.2 218 \times 1.2 mm² for TR 1000 ms, FOV 6.4 \times 1.2 mm² for TR 200 ms, and matrix 64 \times 32. The fMRI experiment set-up 219 220 was identical to those of the GELINE in evoked fMRI.

221 **Data Analysis**. All signal processing and analyses were implemented in MATLAB software (Mathworks, Natick, 222 MA) and Analysis of Functional NeuroImages software⁴³ (AFNI, NIH, USA). For evoked fMRI analysis for **Fig.** 223 **1A**, the hemodynamic response function (HRF) used was the default of the block function of the linear program 224 3dDeconvolve in AFNI. BLOCK (L, 1) computes a convolution of a square wave of duration L and makes a peak 225 amplitude of block response = 1, with $g(t) = t^4 e^{-t}/[4^4 e^{-4}]$. Each beta weight represents the peak height of the 226 corresponding BLOCK curve for that class. The HRF model was defined as follows:

227
$$HRF(t) = int(g(t - s), s = 0..min(t, L))$$

228 Cortical surfaces were determined based on signal intensities of fMRI line profiles as described in the previous work. The detailed processing was conducted as provided in the previous line-scanning studies ^{1,5,8}. For Fig. 1I and 229 1K, demeaned fMRI time courses were used as follows: $(x - \mu)$, where x was the original fMRI time courses and μ 230 was the mean of the time courses. The line profile map concatenated with the multiple fMRI signals was normalized 231 by a maximum intensity. The Z-score normalized time courses were calculated as follows: $(x - \mu)/\sigma$, where x was 232 original fMRI time courses and μ , σ were the mean and the standard deviation of the time courses, respectively 233 234 (zscore function in MATLAB). Average BOLD time series and percentage changes were defined as $(S-S0)/S0 \times$ 100 %, where S was the BOLD signal and S0 was the baseline. S0 was obtained by averaging the fluctuation signal 235 in the 1-second pre-stimulation window in evoked fMRI that was repeated every 20 seconds with the whole time 236 series (640 sec). The BOLD time series in each ROI were detrended ('polyfit' function in Matlab, order: 3) and 237 bandpass filtered (0.01-0.1 Hz, FIR filter, order: 4096). The bandpass filtering was performed as a zero-phase filter 238 239 by 'firl' and 'filter' functions in Matlab, compensating a group delay ('grpdelay' and 'circshift' functions in Matlab) 240 introduced by the FIR filter. Temporal signal-to-noise ratio (tSNR) values were calculated across the cortical depths to compare tSNR differences between GELINE and SELINE. Student t-test was performed with the tSNR values 241 of GELINE and SELINSE (Fig. 1H). The p-values < 0.05 were considered statistically significant. 242

Bloch stimulation. To optimize the α° excitation flip angle with short TR (i.e., 200 ms) in SELINE, signal intensities were calculated as a function of excitation flip angle by simulating the Bloch equation^{22,23}, by employing the refocusing 180° RF pulse. The maximum signal intensity occurred at the Ernst angle which was defined as follow:

247
$$S_{xy}(\alpha,\beta) = \frac{\sin(\alpha) \cdot [1 - \cos(\beta) \cdot e^{-TR/T1} - \{1 - \cos(\beta)\} \cdot e^{-(TR - TE/2)/T1}]}{1 - \cos(\alpha) \cdot \cos(\beta) \cdot e^{-TR/T1}} \cdot e^{-TE/T2}$$

- where α , β indicate excitation and refocusing flip angles, respectively and T1, T2 indicate longitudinal and
- transverse magnetization parameters, respectively. T1 and T2 values were estimated from the previous
- study²⁴.
- Data availability. All other data generated during this study are available from the corresponding author upon
 reasonable request.
- Code availability. The related image processing codes are available from the corresponding author upon reasonable
 request.
- 255 Competing interests. The authors declare no competing interests.

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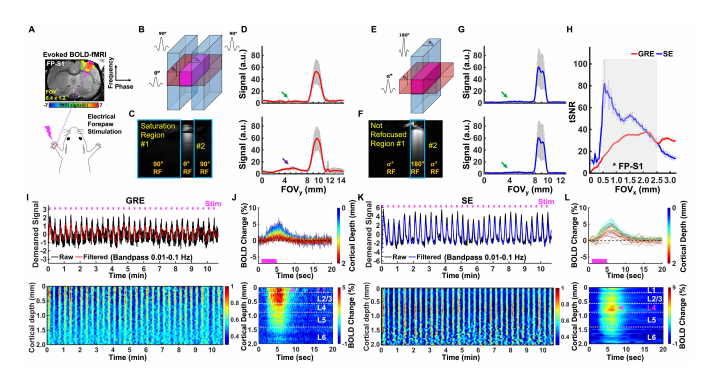
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362 Figure 1. Evoked BOLD responses upon left forepaw stimulation using the GELINE and SELINE methods. A. Schematic illustration of the 363 evoked fMRI experimental design on the EPI-BOLD activation map of FP-S1 region overlaid on an anatomical RARE image. B-C. Schematic 364 drawing of GELINE imaging (B) and an acquired 2D image of GELINE (C). D. two representative 2D line-profiles of GELINE (average of 365 40 voxels): good saturation (green arrow) and bad saturation (purple arrow). Error bars represent mean \pm SD across the cortical depths (0-2 366 mm). E-F. Schematic drawing of SELINE imaging (E) and an acquired 2D image of SELINE (F). D. two representative 2D line-profiles of 367 SELINE (average of 40 voxels): good saturation (green arrows). Error bars represent mean \pm SD across the cortical depths (0-2 mm). H. 368 tSNR comparison between GELINE and SELINE (t-test: $*p < 10^{-12}$). I-J. A representative trial of GELINE. I. Top: Demeaned fMRI time 369 series (32 epochs, 10 min 40 sec) of raw (black) and filtered (red) data (average of 40 voxels, bandpass: 0.01-0.1 Hz) in the FP-S1 region 370 during electrical stimulation (3 Hz, 4 s, 2.5 mA) to left forepaw. Bottom: Normalized spatiotemporal map of the laminar-specific responses 371 along the cortical depths (0-2 mm, 50 µm resolution). J. Top: Average BOLD time courses and Bottom: Average percentage change map 372 across the cortical depths (0-2 mm, 40 lines in total) in the FP-S1. K-L. A representative trial of SELINE. K. Top: Demeaned fMRI time 373 series (32 epochs, 10 min 40 sec) of raw (black) and filtered (red) data (average of 40 voxels, bandpass: 0.01-0.1 Hz) in the FP-S1 region 374 during electrical stimulation (3 Hz, 4 s, 2.5 mA) to left forepaw. Bottom: Normalized spatiotemporal map of the laminar-specific responses 375 along the cortical depths (0-2 mm, 50 um resolution). L. Top: Average BOLD time courses and Bottom: Average percentage change map 376 across the cortical depths (0-2 mm, 40 lines in total) in the FP-S1. Pink arrows indicate peak BOLD signals across the cortical layers.

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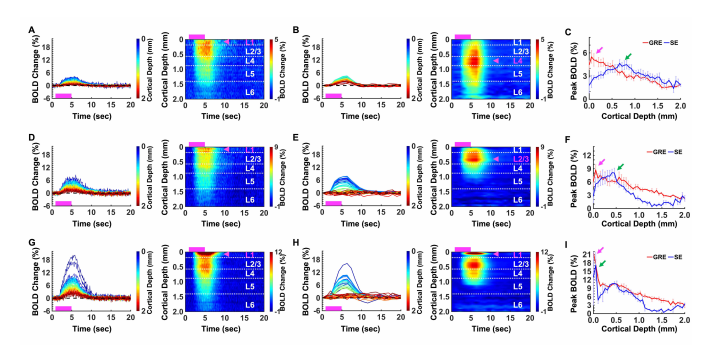
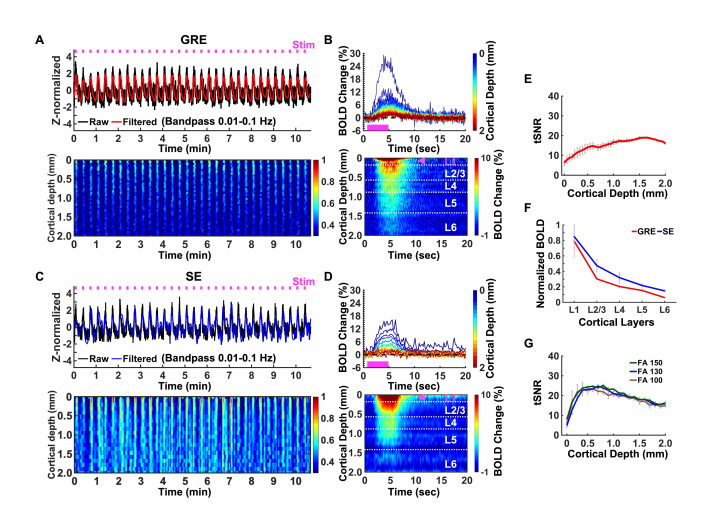




Figure 2. Evoked fMRI time series and percentage change maps of GELINE and SELINE in rat brains (14 trials of 3 rats). A-C. Rat #1 (3
trials of each). D-F. Rat #2 (2 trials of each). G-I. Rat #3 (2 trials of each). A, D, G. *Left*: Average BOLD time courses and *Bottom*: Average
percentage change map of GELINE across the cortical depths (0-2 mm, 40 lines in total) in FP-S1 region. B, E, H. *Left*: Average BOLD time
courses and *Bottom*: Average percentage change map of SELINE across the cortical depths (0-2 mm, 40 lines in total) in FP-S1 region. Pink
boxes indicate stimulation duration and pink arrows indicate peak BOLD signals across the cortical layers. C, F, I. Comparison of peak
BOLD signals between GELINE (pink arrows) and SELINE (green arrows). Error bars represent mean ± SD of peak BOLD signals.



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386 Figure 3. Evoked fMRI responses with GRE (TR 100 ms) vs. SE (TR 200 ms). A-B. GELINE (2 trials) A. Top: Z-score normalized fMRI 387 time series (average of 40 voxels) of FP-S1. Bottom: Normalized spatiotemporal map of the laminar-specific responses along the cortical 388 depths (0-2 mm, 50 µm resolution). B. Top: Average BOLD time courses and Bottom: Average percentage change map across the cortical 389 depths (0-2 mm, 40 lines in total) in the FP-S1. C-D. SELINE (3 trials) C. Top: Z-score normalized fMRI time series (average of 20 voxels) 390 of FP-S1, Bottom: Normalized spatiotemporal map of the laminar-specific responses along the cortical depths (0-2 mm, 100 um resolution). 391 D. Top: Average BOLD time courses and Bottom: Average percentage change map across the cortical depths (0-2 mm, 20 lines in total) in 392 the FP-S1. E. tSNR of GELINE (2 trials) across the cortical depths (0-2 mm). F. Comparison of normalized BOLD signals between GELINE 393 and SELINE across cortical layers. G. tSNR comparison of SELINE with three excitation flip angles (calculated by the Bloch simulation) 394 across the cortical depths (0-2 mm): FA 100 (5 trials), FA 130 (3 trials), and FA 150 (3 trials). Pink boxes indicate stimulation duration and 395 pink arrows indicate peak BOLD signals across the cortical layers. Error bars represent mean \pm SD.