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Abstract. Rod-dominated transient retinal phototropism (TRP) has been observed in freshly isolated retinas, promising a noninvasive biomarker for objective assessment of retinal physiology. However, in vivo mapping of TRP is challenging due to its subcellular signal magnitude and fast time course. We report here a virtually structured detection-based super-resolution ophthalmoscope to achieve subcellular spatial resolution and millisecond temporal resolution for in vivo imaging of TRP. Spatiotemporal properties of in vivo TRP were characterized corresponding to variable light intensity stimuli, confirming that TRP is tightly correlated with early stages of phototransduction. © 2018 Society of Photo-Optical Instrumentation Engineers (SPIE) [DOI: 10.1117/1.JBO.23.5.050502]

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It has been well established that many eye diseases, such as age-related macular degeneration (AMD) and retinitis pigmentosa (RP), can impair retinal photoreceptors and inner neurons to cause vision loss.1,2 As physiological abnormalities may occur before detectable morphological distortions, functional imaging of retinal physiology is essential for early diagnosis of eye diseases and reliable assessment of treatment outcomes. Although psychophysical and electrophysiological methods can be used for functional examination of the visual system, they suffer from low signal specificity or insufficient spatial resolution.3,4 Therefore, a high-resolution method for objective examination of photoreceptor physiology is desirable to advance early diagnosis of AMD, RP, etc. Particularly, rod photoreceptors are known to be more vulnerable, compared with cone photoreceptors, in early AMD and RP.1,2

Rod-dominated transient retinal phototropism (TRP) has been recently observed in freshly isolated amphibian and mammal retinas stimulated by oblique visible light illumination.5 Functional OCT of living eye-cups and time-lapse microscopy of retinal slices revealed that TRP has an anatomic origin within the outer segment (OS),6 presumably caused by localized shrinkage of rod OSs.9 Comparative electrophysiological investigations into isolated retina further identified the physiological source of TRP to the phototransduction processes before hyperpolarization of the rod photoreceptor.10 Therefore, TRP provides a noninvasive biomarker for objective assessment of rod function, promising a high-resolution method for early detection of AMD, RP, etc. However, in vivo mapping of the rod-dominated TRP is challenging due to its rapid time course and subcellular movement magnitude.7,9,10

We recently demonstrated a super-resolution scanning laser ophthalmoscope for in vivo imaging of frog retina.11 The custom-designed ophthalmoscope employed virtually structured detection (VSD) to achieve subcellular level spatial resolution, and it combined a rapid line-scan strategy to realize millisecond-level temporal resolution. In this paper, we report the first in vivo observation of TRP using the VSD-based line-scan super-resolution ophthalmoscope. Temporal dynamics of in vivo TRP correlated with variable stimulus intensities was characterized to verify the physiological origin of TRP.12

Adult northern leopard frogs (Rana Pipiens) were used for this study. The frogs were first dark-adapted for at least 4 h prior to in vivo imaging and then anesthetized through the skin. After confirmation of anesthesia, the frog was fixed in a custom-built holder and the pupils were fully dilated with topical atropine (1%) and phenylephrine (2.5%) for in vivo imaging. All experiments in this research were performed following the protocols approved by the Animal Care Committee at the University of Illinois at Chicago, and conformed to the statement on the use of animals in ophthalmic and vision research, established by the Association for Research in Vision and Ophthalmology.

Figure 1(a) shows a schematic diagram of the line-scan super-resolution ophthalmoscope. The light source is a near-infrared superluminescent diode (SLD-35-HP; Superlum Ireland, Inc.) with a center wavelength at 830 nm and a bandwidth of 60 nm. The light entering the frog pupil had a ∼2-mm beam diameter, with a ∼2.5-mW power. A focused line in the X-direction, produced by a cylindrical lens, scanned across the retina in the Y-direction under the control of a scanning galvanometer mirror (GV5001; Thorlabs, Inc.). Due to the line focus illumination and microsecond-level exposure time, the illumination power applied to the retina was well below the maximum permissible exposure determined by the ANSI and IEC laser safety standards.13–15 The line profile reflected from the retina was recorded by a high-speed two-dimensional (2-D) CMOS camera (FastCam Mini AX50; Photron, Inc.). To achieve fast recording speed for in vivo imaging, the line scanning was performed in one dimension (Y-direction) for super-resolution imaging. A total of 255 line-profiles were acquired to reconstruct one super-resolution image. In this experiment, the imaging speed of the camera was set at 30,000 frames/s (fps), corresponding to a 100-fps speed for VSD-based super-resolution imaging.

A fiber-coupled light-emitting diode with a central wavelength at 505 nm (M505F1; Thorlabs, Inc.) was employed to produce the green flash used for retinal stimulation. The visible stimulation was obliquely delivered to the retina to elicit TRP [Fig. 1(b)] and its incident angle on the retina was adjusted by...
a kinetic mount (KC1; Thorlabs, Inc.) that held both the fiber tip and the collimator. To better distinguish the stimulus-evoked photoreceptor movement from background, only the retinal area at the center of the field of view was illuminated by the visible stimulation [middle and left panel in Fig. 1(b)]. The localized stimulation was achieved by placing a slit at the conjugate plane of the retina in the stimulation path [Fig. 1(a)]. The stimulating power was first measured by a power meter (PM200; Thorlabs, Inc.) placed at the rear focal plane of the lens before the eye [L4 in Fig. 1(a)] and was then converted to stimulation intensity on the retina.16

Figure 2 shows in vivo TRP correlated with oblique visible stimulation. Figure 2(a) shows a representative image of the photoreceptor layer acquired by the VSD-based super-resolution ophthalmoscope with an imaging speed of 100 frames/s and a field of view of 100 μm x 200 μm on the retina. Individual photoreceptors were clearly imaged with subcellular spatial resolution and millisecond temporal resolution, which allowed quantitative measurement and dynamic monitoring of photoreceptor movements. To elicit TRP, a rectangular stimulation pattern was obliquely projected onto the retina and stimulated a retinal area that has a width of ~50 μm at the center of the field of view [red-dashed rectangle in Fig. 2(a)].

To quantify TRP, we calculated the magnitude and direction map of photoreceptor movements based on the retinal images using the optical flow code developed by Sun et al. Optical flow is a well-established method for measuring object movements between two images. Therefore, it can accurately identify the process of photoreceptor movement through sequential retinal images. A 3-σ threshold and a temporal window generated from the movement magnitude maps were applied to exclude potential spatial and temporal noise in the movement direction map. Figures 2(b) and 2(c) are representative magnitude and direction map of the photoreceptor movements evoked by the localized stimulation. As Figs. 2(b) and 2(c) match Fig. 2(a), the red-dashed rectangles shown in Figs. 2(b) and 2(c) also represent the stimulated retinal area. Therefore, Figs. 2(b) and 2(c) reveal that robust photoreceptor movements were primarily confined within the stimulated retinal region. The distribution of movement direction in Fig. 2(c) was further analyzed to illustrate the correlation between the directions of photoreceptor movements and oblique stimulation. As shown in Fig. 2(d), most stimulus-evoked photoreceptor movements had a direction close to ~90 deg, indicating the photoreceptors moved toward the incident stimulation, which was consistent with our previous in vitro observation of TRP.7,8,10

To illustrate the spatiotemporal dynamics of TRP, representative magnitude and direction maps of photoreceptor movement selected from different phases of the experiment are shown in Fig. 3. As the stimulation onset was set as 0 s and the stimulation period lasted for 0.5 s, Fig. 3 covers the photoreceptor movements before, during, and after the stimulation. Figure 3(a) demonstrates that robust photoreceptor movements were only observed after the onset of stimulation and within the stimulated retinal area. Figures 3(b) and 3(c) further proved the direction of stimulus-evoked photoreceptor movement was consistently toward the direction of incident stimulation during and after the stimulation period. It was also noticed that some of the stimulated retinal regions did not present reliable movements during the recording period and some other regions presented opposite movement direction at certain time points [pink areas in Figs. 2(c) and 3(b)]. Similar phenomena were observed in our previous in vitro TRP studies using different imaging modalities, including OCT, confocal, and light microscopies. These phenomena are speculated to be related to the complex structure of the retina, and accurate oblique stimulation and actual intensity control for each single photoreceptor are difficult due to inevitable light scattering and coherent mechanical interaction among neighboring photoreceptors.

The correlation between photoreceptor movement magnitude and stimulation intensity was further investigated. Three stimulation intensities, i.e., 1.97 x 103, 0.67 x 104, and 0.197 x 105 photons · μm² · ms⁻¹, were tested as a pilot study.
Previous in vivo intrinsic optical signal (IOS) studies showed such stimulations could activate significant but different retinal responses.\textsuperscript{19} The flash duration was set to 500 ms for eliciting robust photoreceptor movements. As the line-scan modality fulfilled a fast recording speed, a time-magnitude course was used to reflect the dynamics of photoreceptors with a 10-ms temporal resolution.\textsuperscript{10} Figure 4 shows the results obtained from a group of six samples. Each trace in Fig. 4(a) is the mean of 12 time-magnitude courses and is accompanied by the standard deviations of data about the mean (colored area). The relationship of the standard deviation amplitudes to the waveform in each trace indicates general similarity between the results obtained from different samples. The waveforms of all three traces are flat and stable in the prestimulus phase and exhibit an immediate and significant rise upon the initiation of the stimulation. However, the waveforms also show that different stimulation intensity remarkably changed the response of the photoreceptor.

To better illustrate the difference caused by increasing stimulation intensity, two parameters, peak amplitude (the maximum value of photoreceptor movement magnitude) and time-to-peak (time taken to reach the peak amplitude) of the waveforms, were compared. The results in Fig. 4(b) show that the difference between peak amplitudes of the three traces was statistically insignificant, suggesting the peak amplitude is irrelevant to the variation of stimulation intensities within a certain range. However, as shown in Fig. 4(c), the time-to-peak values were significantly reduced by higher stimulus intensity, indicating brighter stimulus results in earlier saturation of the photoreceptor movement. These results suggested an intimate correlation between photoreceptor movement and phototransduction as the accelerated photoreceptor response might be the consequence of more rhodopsin and additional amounts of cascaded reactions activated by strong stimulation.\textsuperscript{20} Similar effects were observed in our previous IOS and rod OS shrinkage studies, supporting that
photoreceptor movement is a major component of IOS and the unbalanced rod OS shrinkage is the mechanical source of TRP.\(^{18,21}\)

In summary, this study demonstrated the feasibility of using a line-scan VSD-based super-resolution ophthalmoscope for in vivo imaging of TRP. Compared with traditional structure illumination microscopy, the VSD-based approach provides a compact, cost-efficient, and phase-artifact-free strategy to achieve super-resolution retinal imaging. The line-scan imaging modality achieved a fast recording speed for retinal activity. Corresponding to variable light intensity stimuli, it was observed that the TRP peak amplitude was not significantly sensitive to the stimulus light intensity, whereas TRP time-to-peak value was significantly sensitive to stimulus light intensity, i.e., reduced time-to-peak value corresponding to enhanced stimulus intensity. This observation indicates that the rod-dominated TRP is closely related to phototransduction in rod photoreceptors, and therefore reflected the functionality of rod photoreceptors. Further investigation of the in vivo properties of TRP may provide a high spatial resolution IOS imaging method for functional mapping of rod physiology. Our recent study has revealed a tight correlation between stimulus-evoked TRP and photoreceptor IOS changes in freshly isolated retinal tissues.\(^{7}\) Previous studies have revealed IOS changes in both photoreceptors and inner retinal layers,\(^{14,15}\) and emerging functional OCT enabled depth-resolved detection of photoreceptor IOS from inner retinal IOS.\(^{18,21}\) We anticipate that better investigation of the in vivo properties, including physical and physiological mechanisms and spatial and temporal dynamics, of stimulus-evoked TRP in animal models will provide valuable information for advanced instrument development and better stimulation protocol designs for pursuing in vivo IOS imaging of human retina, enabling early detection of AMD, in which rod photoreceptor dysfunction occurs first, and other eye diseases.

Disclosures
No conflicts of interest, financial or otherwise, are declared by the authors.

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