Arbitrary multisite two-photon excitation in four dimensions

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We demonstrate dynamic and arbitrary multisite two-photon excitation in three dimensions using the holographic projection method. Rapid response (fourth dimension) is achieved through high-speed noniterative calculation of the hologram using a video graphics accelerator board. We verify that the projected asymmetric spot configurations have sufficient spatiotemporal photon density for localized two-photon excitation. This system is a significant advance and can be applied to time-resolved photolysis of caged compounds in biological cells and complex neuronal networks, nonlinear microfabrication and volume holographic optical storage. © 2009 American Institute of Physics. [DOI: 10.1063/1.3216581]

Advanced microscopy and diagnostic techniques require full three-dimensional (3D) access to the sample with fast dynamic control. Developments in array illumination schemes have facilitated simultaneous control and manipulation of multiple microparticles.1–3 Two-dimensional (2D) patterned illumination techniques have also been used for high-speed 3D image rendering of a sample. Techniques based on programmable array microscopy,4 structured illumination,5 and rotating Nipkow disks6–8 are methods using 2D excitation patterns, confocal detection schemes, and postprocessing algorithms to render fluorescent samples in 3D.

However, some experiments require an intrinsic 3D localized excitation response prior to detection. This can be achieved via nonlinear two-photon excitation (2PE).9 Such experiments (e.g., photolysis of biomolecules) also require multiple excitation sites at arbitrary positions in 3D. The challenge is therefore to design a system that provides for arbitrary multisite 2PE and meets the response time of biological tissues and cells, typically ~1 ms.

In this letter, we have expanded and combined the techniques of high-speed holographic beam control and nonlinear multiphoton absorption to demonstrate a fully controllable multisite 2PE in four dimensions. Our system significantly advances the implementation of 3D nonlinear microscopy,7,8 volume holographic storage,10 microfabrication,11 and nanosurgery.12 In neuroscience, our system allows for realistic simulation for summating simultaneous synaptic signals from multiple sites within complex dendritic trees of a neuron. Recently, multisite single-photon (1P) photolysis in neuronal networks has been demonstrated.13 However, 1P absorption is a linear process and suffers from poor 3D localized excitation.

Highly localized 2PE within a 3D diffraction-limited focal spot is achieved at high spatiotemporal photon densities provided by a focused femtosecond pulsed laser. Time-multiplexed multisite 2PE has been shown to work with high-speed scanning via acousto-optic modulators (AOMs).14,15 However, dispersion through an AOM crystal degrades the spatiotemporal quality of the femtosecond pulsed laser, especially when four AOMs are used to project the focal spot in arbitrary 3D positions.16 Multisite 2PE along a plane has also been demonstrated with the holographic method that uses an iterative-adaptive optimization algorithm to produce photon efficient spot arrays.17 They also incorporate a general lens function to shift the whole plane containing the spot array along the optical axis. On the contrary, we show a fast and arbitrary multisite 2PE in all three dimensions. Each excitation site has a localized response in 3D. We derive the hologram using a noniterative method and achieve optimal speeds by taking advantage of the parallel computing capability of a graphics accelerator board.

A phase hologram for 2PE at N arbitrary sites can be calculated with no iterative optimization procedure by the superposition of N fields.12 Each field is a combination of prism and lens phase functions that describe the 3D position of a focal spot. The localized 2PE profile is derived by the square of the intensity distribution of the excitation focal spot. Aberrations disturb the excitation spots when they are arbitrarily positioned around the sample. To visualize these aberrations, we use the nonoptimized holograms as input and calculate the 3D intensity profile using the Fresnel diffraction integral.18 Figure 1(a) shows the intensity distribution along the xz-plane with two spots (N=2) symmetrically positioned (0,0,f±5/ξ) with respect to the nominal focus (f) of the lens. The normalized transverse (x,y) and axial (z) coordinates are related to spatial coordinates by η=NAλ−1 and ξ=NA2λ−1, respectively, where NA is the numerical aperture of the objective lens and λ is the wavelength of the laser. For N=2, the maximum normalized intensity of each spot is 0.405. The axial intensity distribution of the spots is characterized by aberrations with uneven side lobes along the z-axis. These aberrations increase as the spots are positioned away from the focal plane of the lens. Figure 1(b) shows N=4 spots positioned at (±2.25/ η,0,5.25/ξ) at the top layer and (±2/ η,0,−5/ξ) at the bottom layer. For N=4, the max-

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which makes use of a linearly polarized TEM$_{00}$ laser beam. Lenses L1 and L2 expand the beam to illuminate the 16 × 12 mm$^2$ area of the phase-only SLM (Hamamatsu Photonics). The overall efficiency of the SLM is 92% with a fill factor of 95%. The update rate is currently limited by the SLMs response time of 30 ms. Electronic addressing of each pixel in the SLM is achieved via the video output of the graphics accelerator board.

The SLM operates in reflection mode, and separating the incident and the phase-encoded reflected beam with some degree of tilt optimizes the throughput. We minimize the tilt by reusing lens L2 to serve as the Fourier transform lens. The spots are formed at the focal region of L2 and relayed to microscopic scale using L3 and the objective lens. A dichroic mirror reflects the phase encoded near-infrared beam to the microscope while allowing visible light to pass through for bright field and fluorescence imaging. A standard microscope configuration with an appropriate filter is used to view the fluorescence along the $x\z$-plane.

To test arbitrary multisite 2PE along the focal plane, we prepared 2 μm diameter fluorescence latex microspheres (Molecular Probes, Inc.) diluted with water. The microspheres emit green fluorescence following 2PE from a focused near-infrared femtosecond pulsed laser. Figure 3 shows multiple excitation site configurations. The excitation spots are arbitrarily positioned around the optical axis, which is at the center of the micrograph. The average power of each excitation spot, including system losses, is approximately 2.9, 2.0, 1.5, and 1.2 mW for $N=15, 20, 25$ and 30, respectively. System loss of around 10% is associated with light diffracted due to the physical constraints of the SLM. Tolerable variations in fluorescence intensity are due to slight off-center illumination on the beads as well as slight differences in the maximum spot intensity as described earlier.

Next, we demonstrate multisite 2PE along the optical axis. Figure 4(a) shows a photograph of the setup where we used a 63× (NA=0.9) objective lens for projecting the excitation spots and a 10× lens for viewing the fluorescence...
from the side. We prepared a dye solution from carboxyfluorescein dissolved in water and sodium hydroxide. Figure 4(b) shows two localized 2PE sites positioned symmetrically at $x = \pm 10 \mu m$. Also shown are two [Fig. 4(c)] and four [Fig. 4(d)] 2PE sites projected along the optical axis. With NA $=0.9$, the maximum displacement along the $z$-axis is around $\sim 100 \mu m$. Since there is an overlap between the absorption ($\lambda_{abs} = 492$) and emission ($\lambda_{em} = 517$ nm) spectra of carboxyfluorescein, the emitted fluorescence from the 2PE process induces 1P fluorescence at neighboring regions around the excitation spot. Hence, the fluorescence detected from the side is smeared and does not reflect the true 3D localized excitation provided by the 2PE process. Nonetheless, such demonstration establishes our capacity to provide arbitrary and localized 2PE along the optical axis.

When applied to photolysis of caged neurotransmitters, irregular and asymmetric spot distributions match the arbitrary dendritic trees of neurons. Hence, high variations in spot intensities associated with symmetric spot arrays will not pose a major problem. It is also important to note that the programmability of the SLM should allow for appropriate wavefront corrections to rectify aberrations at $z$-positions further from the focal plane. Such corrections will be worked out in future studies.

We have demonstrated multisite and highly localized 2PE in four dimensions. Noniterative calculations carried out with the graphics accelerator board generate holograms on-the-fly with the update rate currently limited by the response time of the SLM. Multisite fluorescence emission indicates that the spots have sufficient spatiotemporal photon density for 2PE. For quantitative measurements, standard optical theory can account for slight intensity differences introduced by nonoptimal holograms. Immediate applications of our system point to neuroscience, where photostimulation of neurotransmitters at arbitrary sites provides fundamental answers to the mechanisms of information processing in the human brain.

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