High-speed laser microsurgery of alert fruit flies for fluorescence imaging of neural activity

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Intravital microscopy is a key means of monitoring cellular function in live organisms, but surgical preparation of a live animal for microscopy often is time-consuming, requires considerable skill, and limits experimental throughput. Here we introduce a spatially precise (<1-μm edge precision), high-speed (<1 s), largely automated, and economical protocol for microsurgical preparation of live animals for optical imaging. Using a 193-nm pulsed excimer laser and the fruit fly as a model, we created observation windows (12- to 350-μm diameters) in the exoskeleton. Through these windows we used two-photon microscopy to image odor-evoked Ca^{2+} signaling in projection neuron dendrites of the antennal lobe and Kenyon cells of the mushroom body. The impact of a laser-cut window on fly health appears to be substantially less than that of conventional manual dissection, for our imaging durations of up to 18 h were ∼5-20 times longer than prior in vivo microscopy studies of hand-dissected flies. This improvement will facilitate studies of numerous questions in neuroscience, such as those regarding neuronal plasticity or learning and memory. As a control, we used phototaxis as an exemplary complex behavior in flies and found that laser microsurgery is sufficiently gentle to leave it intact. To demonstrate that our techniques are applicable to other species, we created microsurgical openings in nematodes, ants, and the mouse cranium. In conjunction with emerging robotic methods for handling and mounting flies or other small organisms, our rapid, precisely controllable, and highly repeatable microsurgical techniques should enable automated, high-throughput preparation of live animals for optical experimentation.

Significance

Microscopy and neurophysiology experiments in live animals commonly involve complex surgical preparations, which are often time-consuming, demand considerable manual dexterity, and can sharply limit experimental throughput. Here we present a spatially precise laser microsurgical technique using a pulsed UV laser. Our approach reduces surgical time by up to two orders of magnitude while substantially improving reproducibility. Using the fruit fly as a model, we show that laser microsurgery leaves complex behaviors intact and allows us to visualize brain activity in live flies for up to 18 h, more than four times longer than reported previously using hand dissection. We also demonstrate laser microsurgery on nematodes, ants, and the mouse cranium, illustrating broad potential utility for both optical and electrophysiological studies.

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Either 0.2 and -nm wavelength excimer laser. To Sinha et al.

Table 1. Approximate durations for each step. Red font indicates steps at which our protocol improves over conventional methods. (A) Schematic of the laser microsurgery station based on a 193-nm-wavelength excimer laser. To monitor surgical progress, light from a blue LED enables imaging of green fluorescence from the fly. Photodiodes (PD) provide readouts of laser power and fluorescence emissions. A mercury lamp with a green fiber output enables bright-field imaging. A flow of air prevents ejected debris from the surgery from depositing on the focusing objective. A syringe pump enables bright-field imaging that we are also developing to further speed experimental throughput.

Results

We chose our laser wavelength and pulse duration to affect fly behavior minimally and avoid photobleaching fluorescent markers just beneath the cuticle. Initial tests with a 355-nm laser showed that this wavelength induced substantial photobleaching. However, UV laser ablation with an excimer laser of 193-nm wavelength seemed suited to our needs (Fig. 1B) because photons of this wavelength can break covalent bonds but have an absorption length in water of <1 μm (13). Thus, tissue absorbs the UV energy within a very shallow surface layer, and the superheated material is ejected before thermal energy diffuses to surrounding areas (14). We reasoned that this should avoid photobleaching beneath the ablated layer because fluorophores more than a few microns below the cut surface receive a tiny fraction of the incident light intensity.

Microfabricated, Silicon Mounts Facilitate Rapid Presurgical Preparation.

For the first step of the surgical protocol we developed means to quickly fix a fly’s body in place while permitting its legs and wings to move freely (Figs. S1 and S2 and Movie S1). To do this, we used UV-cured epoxy to attach the fly to a 125-μm-diameter silica fiber. The fiber lay in a precision-etched V groove created within a silicon wafer (SI Materials and Methods). The fixture was designed to be compatible with a set of robotic techniques for fly handling that we are also developing to further speed experimental throughput.

We positioned the fly with ~25-μm repeatability in all three spatial dimensions relative to fiduciary marks etched in the silicon wafer and the well-defined depth of the V groove (Figs. S1 and S2). We placed a metal strip with an 800-μm-diameter opening on top of the fly. This opening provided optical and fluidic access to the fly, allowing us to immerse exposed tissue in saline immediately after surgery (Materials and Methods).

whether these lasers can open optical windows in an organism’s exoskeleton for in vivo microscopy.

Here we report a largely automated microsurgery protocol that uses a nanosecond-pulsed UV excimer laser to gain optical access to the brain. Our approach enables swift (<1 s) removal of the exoskeleton and precise cuts of high repeatability. These are general capabilities that we illustrate here in three invertebrate species. We also demonstrate surgical incisions in the mouse cranium with sizes and spacings that are infeasible using drills and scalpels.

In flies, we used a phototaxis assay to show that our surgical technique can be performed with minimal to no measurable impact on fly behavior. Microsurgical removal of the cuticle facilitates in vivo imaging of neuronal Ca++ dynamics, as illustrated here in olfactory neurons. Microsurgery is also sufficiently gentle to permit intravital imaging over extended time periods, as demonstrated by our Ca++-imaging studies of Kenyon cells in the fly mushroom body for up to 18 h.

Fig. 1. Laser microsurgery of fruit flies is faster and more precise than manual dissection. (A) (Left) Work flow for imaging flies in vivo. (Right) Approximate durations for each step. Red font indicates steps at which our protocol improves over conventional methods. (A) Schematic of the laser microsurgery station based on a 193-nm-wavelength excimer laser. To monitor surgical progress, light from a blue LED enables imaging of green fluorescence from the fly. Photodiodes (PD) provide readouts of laser power and fluorescence emissions. A mercury lamp with a green fiber output enables bright-field imaging. A flow of air prevents ejected debris from the surgery from depositing on the focusing objective. A syringe pump enables bright-field imaging that we are also developing to further speed experimental throughput.

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the fly shown in E, H and I show the holes cut in the left and right retinae, respectively. (Scale bar, 25 μm.) (J) Two-photon image of UAS-TublNLH:Y237M6B fly following a microsurgery in which 150 pulses (350-μJ pulses; 100 Hz) drilled through the retina of a fly mounted on its side (spot size 400 μm × 350 μm). The image is a maximum-intensity projection of a 3D image stack (143 slices acquired at 1-μm intervals). (Scale bar, 25 μm.) (K) Signal intensity from fluorescently labeled neurons lying under the cuticle rises with the number of laser pulses used (200 μJ each) to drill a 300-μm-diameter circular spot. Each data point represents the median ±1 SD for 10–12 female flies (UAS-GFP;OK107-Gal4) that underwent laser surgery 6 d following eclosion. (L and M) Two-photon images of UAS-GFP;OK107-Gal4 flies after manual dissection (L) or after laser surgery (M). For both images the laser power (920-nm wavelength) at the specimen plane was 3 mW. (Scale bars, 5 μm.)
A Flexible Microdissection Protocol Based on Laser Percussive Drilling. In mounted flies positioned under a UV-compatible objective lens (Fig. 1B), we examined whether nonscanning, laser percussive drilling (15) allowed dissection of the fly cuticle or if a laser-scanning approach was necessary. Percussive drilling offers superior cutting speed but requires sufficient pulse energy to drill in parallel all subregions within the area demarcated for ablation. We found that the thickness of the fly cuticle is sufficiently uniform that percussive drilling can precisely create a window in the cuticle over the entire ~0.1-mm² area illuminated by the surgical laser (275 mJ/cm²) (Fig. 1 C–F), without ablating underlying brain tissue. With <1 s of drilling, laser pulses of 200 μJ delivered at 100 Hz removed a 300-μm-diameter window in the cuticle. We thus chose this mode of microsurgery rather than laser-scanning approaches, which would have required additional hardware for scanning and much higher laser repetition rates (e.g., ~3 orders of magnitude higher for a ~10-μm-diameter laser spot) to achieve the same rate of tissue removal. Our surgery station allowed us to select the size and shape of the ablation by using different apertures or masks that were placed in the laser beam’s delivery pathway and projected onto the cuticle surface (Fig. 1 B–E). The edges of the laser cuts in tissue were locally straight; electron microscopy studies of postsurgical specimens revealed ≤100 nm of orthogonal deviation from the straight edges (Fig. 1G). Tissue structures as near as 2 μm to the surgical cuts appeared unaltered (Figs. 1 G–I), and laser microsurgery readily created holes as tiny as ~12 μm in diameter (Fig. 1 H and I).

For tailoring surgeries to different parts of the fly brain, we attached flies to the silica fiber in different orientations and adjusted the laser pulse energy and beam size as needed (Fig. 1B). With the fiber on the side of the fly’s neck, we accessed visual neurons that are challenging to expose manually in live flies (Fig. 1J and Movie S2). To access olfactory neurons, we attached the fiber at the nape of the fly’s neck. To monitor the depth of surgical ablation, we used a 470-nm-wavelength, fiber-coupled light-emitting diode (LED) to excite green fluorescent protein (GFP) markers. For example, flies expressing GFP in mushroom bodies (UAS-GFP;OK107-Gal4), fluorescence signals rose monotonically across the mushroom bodies in different orientations and shapes of the ablation by using different apertures or masks (Fig. 1G–I), and laser microsurgery readily created holes as tiny as ~12 μm in diameter (Fig. 1 H and I). Laser Microsurgery Requires Seconds for Dissections with Submicron Precision. We tested the laser surgery system in several ways. As a test of cutting speed and throughput, we mounted four flies (UAS-GFP;OK107-Gal4) on an individual silicon wafer (Fig. S2B). Using motorized translation stages to rapidly address each of the four flies, we opened a 300-μm-diameter window in each cuticle and acquired a fluorescence image of the mushroom bodies. Cutting and imaging took 22 s for all four flies in total. A second test of our surgical system concerned the aspect ratio of the holes it can cut in tissue (Fig. S3). We successfully drilled a constant, 35-μm-diameter hole through the entire ~400-μm-thick fly, without adjusting the fly’s position en route. This aspect ratio is generally unachievable by laser cutting systems that rely on high-numerical aperture (NA) objective lenses, as typically used with near-infrared ultrashort-pulsed laser sources (8, 9).

A third test assessed whether excimer laser surgery induced photobleaching of the GFP markers in the brain. We used two-photon microscopy to inspect five flies (UAS-GFP;OK107-Gal4) expressing GFP in mushroom body olfactory neurons. These cells are among the most commonly examined in flies for studies of olfactory sensing, learning, and memory (7). We manually dissected the cuticles of two of the flies and used laser microsurgery on the other three (120 pulses). Using identical imaging parameters for all five flies, fluorescence signals from all of the flies were comparable to within ~15%. The difference in mean signal intensity (averaged in each fly over several dozen Kenyon cells of the mushroom bodies) between the two manually dissected samples (~15%) was comparable to the variation between the mean intensity from the two manually dissected flies and that from the three laser-cut flies (~13%), suggesting that the variation in GFP expression between flies was greater than any bleaching potentially induced by laser surgery (Fig. 1 L and M).

Last, we tested the applicability of our methods to other species. We attained comparable precision when cutting windows in the cuticle of the harvester ant (Fig. S4) and Caenorhabditis elegans (Fig. S5). Beyond invertebrates, manual dissection can be equally challenging in vertebrates and is often a stumbling block in mammalian neuroscience. Thus, we examined whether excimer laser surgery was effective in the mouse cranium. Strikingly, we found that with ~3 s of pulsed illumination we could cut precise, square openings (~100-μm across) through the entire depth of the cranium (Fig. S6).

Optical Parameters for Minimal Impact on Fly Behavior. To gauge the impact of laser surgery on fly behavior, we subjected hundreds of wild-type flies to various numbers of laser pulses, ranging from 0 to 100. We delivered all pulses at the same repetition rate and with identical spot sizes (275-μm diameter) and energies (170 μJ) to a portion of cuticle centered over the antennal lobes. Following microsurgery we used a T-maze to monitor the flies’ phototactic responses to a light source that is normally attractive (Fig. 2). This test showed that the sensory integration motor pathway (16). We included in this test some flies in which we additionally used laser surgery to cut muscle 16, which reduces brain movement during in vivo imaging.

Flies that received up to 90 laser pulses maintained statistically significant phototactic responses, whether or not muscle 16 was cut [for <80 pulses, P ≤ 3 × 10⁻⁴; n = 141 flies; for 80 pulses,
Imaging of Neural Dynamics over Extended Time Periods. We also used two-photon microscopy to record odor-evoked Ca\(^{2+}\) transients in the antennal lobe, to assess if laser microsurgery impacted flies’ odor-evoked neural responses. We imaged flies (GH146-Gal4, UAS-GCaMP3) expressing the Ca\(^{2+}\) indicator GCaMP3 in olfactory projection neurons (17), after using 85 UV laser pulses (spot size of 325 μm × 225 μm) to remove cuticle tissue (Fig. 3A). GCaMP3 expression was readily visible before odor delivery, and the antennal lobe tissue appeared as normal and healthy as in our prior two-photon imaging studies of manually dissected flies (18) (Movie S3). The ~25-μm repeatability of our mounting protocol allowed us to quickly locate an optical plane showing several different glomeruli. We then delivered multiple odors to the flies. Analysis of the resulting two-photon images revealed patterns of odor-evoked neural excitation and inhibition that were consistent with prior studies on manually dissected flies (4, 19, 20) (Fig. 3B). Odorific responses were prominent, with changes in GCaMP3 fluorescence (∆F/Φ) generally >60% (Fig. 3C).

To demonstrate the extension of imaging duration that is attainable via laser microsurgery, we developed a version of the surgical protocol that was particularly minimally invasive and prohibitive by hand dissection. This involved the creation of a smaller opening in the cuticle (120 μm × 90 μm elliptical opening) plus a surrounding annular region in which we thinned but did not remove the exoskeleton (Fig. 4A). In this way, we attained optical access to the Kenyon cells of the fly’s mushroom bodies (UAS-GCaMP3/+;OK107-Gal4/+via a hole that minimized the impact on fly health while providing adequate transparency across the broader thinned area. This thinned-cuticle strategy allowed us to observe odor-evoked Ca\(^{2+}\) dynamics in these neurons even 18 h after laser microsurgery (Fig. 4B and C) and to track the same individual cells across 6- to 18-h intervals (Fig. S7). By comparison, we were unable to reproduce these results using manual dissection.

Discussion

We developed an excimer laser microsurgery protocol that can rapidly and precisely remove fly cuticle with minimal impact on fly behavior or underlying fluorescent markers. Our approach obviates the highly variable outcomes of manual dissection by providing repeatable, user-friendly, and rapid laser-cutting capabilities with ~100-nm-level precision.

Traditional scanned UV laser ablation setups that operate at a wavelength of 355 nm (21) suffer from longer absorption depths, causing fluorophore bleaching in the brain. In comparison with surgical systems based on ultrafast laser sources, the cost of our instrumentation was ~2–5% the price of a commercial two-photon microscope equipped with a Ti:sapphire laser amplifier. These amplifiers typically require considerable effort to maintain intracavity alignment, but excimer lasers are widely used in medicine and exist in user-friendly versions.

Our surgical success with flies, ants, and nematodes indicates that research involving a range of small model organisms should benefit from our methods. Moreover, our capability to cut through the mouse cranium in ~3 s suggests that excimer laser surgery has substantial utility for research on mammals. Excimer lasers with higher energies are commercially available, so cranial openings larger than those illustrated here could also be created percussively. The cranial openings we cut had sharp corners that would be nearly impossible to create manually, certainly not within seconds.

Indeed, a striking advantage of our laser microsurgery system over manual dissection is our ability to cut holes of arbitrary shape, size, and depth. Our low-NA system can readily create high-aspect ratio holes of small diameter. In flies, mice, or other species, this capability may ease minimally invasive injections or electrical recordings. In imaging experiments, the traditional manual approach of removing the cuticle generally requires making an opening larger than the imaging field of view, to avoid damaging the brain area of study. The larger window in the cuticle constitutes a greater insult to the fly, and the resulting declines in the fly’s health reduce the total time available for imaging neural responses, typically 1 to several hours (1, 2, 7, 18).

Fig. 3. Two-photon imaging reveals normal odor-evoked neural Ca\(^{2+}\) responses in live flies following laser microsurgery. The first column shows baseline two-photon images of three different GH146-Gal4, UAS-GCaMP3 flies expressing the Ca\(^{2+}\) indicator GCaMP3. Several glomeruli are visible, three of which are demarcated with dotted lines. Across flies, glomeruli are labeled consistently by color: VM2, red; VM7, yellow; VMSV, brown; DM2, green; DL1, blue; DL2, orange; and DA1, pink. (Scale bars, 20 μm.) In the second and third columns, maps of odor-evoked fluorescence changes (∆F/Φ) reveal glomerular activation following a 2-s pulse of odor delivery, for different pairs of odors. Both excitatory and inhibitory glomerular responses occurred. Odor-evoked fluorescence changes are indicated in color for regions where the response absolute value was at least three times greater than the standard deviation of baseline fluctuations. The fourth column shows traces of odor-evoked changes (∆F/Φ) for the three encoded glomeruli and two odors indicated in each row. Colors match those of the glomeruli in the first column. Solid and dashed lines are for the odors indicated in the second and third columns, respectively. Odor delivery occurred at times marked by gray bars. Dashed black line indicates the time point shown in the second and third columns. Vertical and horizontal scale bars are 100% ∆F/Φ and 5 s, respectively.
The minimally invasive capabilities of our surgical approach have allowed the longest time-lapse imaging studies of neural activity in live flies reported to date. For these studies we thinned the cuticle over a 200-μm-wide region and created a central opening (~100 μm) smaller than possible by hand. The resulting 6- to 18-h imaging durations so attained represent a ∼4- to 18-fold extension compared with prior studies; this is a notable improvement because the durations attainable via laser microsurgery now suffice to study many unanswered questions in neuroscience, such as those involving neuronal plasticity or the neurobiology of learning and memory.

After mounting a fly, our entire surgical protocol takes only ∼10 s, of which ∼1 s is devoted to laser pulse delivery. With four flies mounted on the same silicon wafer, we completed all four surgeries in 22 s, indicating the amenability of our methods to high-throughput experimentation. By capitalizing on silicon wafers of larger diameter than those used here, our four-fly mount should be extendable to greater numbers of flies, such as for parallelization of imaging studies. Our protocol also allows flies to be mounted with high precision, which reduces the alignment time at subsequent surgery and imaging steps.

Given this speed and efficiency of throughput, our methodology should be able to facilitate future high-throughput screens for flies with mutant cellular, functional, or anatomical phenotypes. We are developing robotic techniques that use automated machine vision algorithms to further reduce the time it takes to pick and mount flies for surgery and imaging. Further, by integrating interferometric measurements of the cutting depth into the surgical process, it might be possible to remove the fly’s perineural sheath without damaging underlying brain structure, such as for electrophysiological studies. Similarly, it should be possible to create a preparation in which the fly cuticle is left entirely unopened and merely thinned for imaging, to further extend the duration of time-lapse experiments. We chiefly focused here on fruit flies, but our precise and repeatable capabilities to create optical windows in nearly any part of an animal’s body should be widely applicable in multiple biological disciplines and to invertebrates and vertebrates alike.

Materials and Methods
Fly Stocks. We crossed GH146-Gal4 and UAS-GCaMP3 flies (2) to label olfactory projection neurons in the antennal lobe. We crossed OK107-Gal4 with UAS-GFP and UAS-GCaMP3 lines to label mushroom body neurons. We raised all flies on standard cornmeal agar media under a 12-h light/dark cycle at 25 °C and 50% humidity. All experiments used adult mated females.

Fabrication and Use of Silicon Mounting Fixtures. We made silicon (Si) fixtures in Stanford’s Ginzton microfabrication facility. We used ∼100-μm-crystallographic orientation, 4-inch-diameter, 500-μm-thick, single-side polished Si wafers with 100 nm of low-stress nitride on both sides (University Wafer). SI Materials and Methods contains detailed microfabrication methods and description of how we mounted the fly to the silica fiber and the fiber to the Si wafer. It took ∼60 s to fix the fly’s position with ∼25-μm accuracy relative to the etched holes in the silicon. The fly was free to move its legs and wings when attached to the fiber (Movie S1).

Laser Surgery System. We used Optex and GamLaser EX5 Arf 193-nm excimer lasers, each capable of generating ∼12-muJ pulses of ∼10-ns duration. The total exposure time of the cuticle to the laser was <1 s, so the fly’s hemolymph would not have time to flow into the machined hole and shield the fly’s air sacs from the laser pulses. A 250-μL/min flow of air from a 6-mm-diameter tube aimed just above the fly prevented deposition of ejected debris on the objective lens. To control the beam shape and spot size on the fly, we used various pupil masks consisting of gold-coated thin copper substrates that were laser machined to have the desired aperture (National Ginzton microfabrication facility). To control the cuticle opening (Movie S1).

Fly Surgical Protocol. After transferring a mounted fly to the surgery station, we measured fluorescence signals using a photoreceiver (Newport 2151) and a 515/20 band pass optical filter (Semrock). For imaging certain brain regions, we reduced brain movement by laser drilling a 50-μm-diameter hole (6 μJ, 1,000 pps, 200 μs per pulse) between the fly antenna to cut muscle 16. We then opened the optical window in the cuticle. Immediately after surgery, we applied 90 μL of adult haemolymph-like (AHL) saline (103 mM NaCl, 3 mM KCl, 5 mM TEA, 1 mM NaH2PO4, 1.5 mM CaCl2·6H2O, 4 mM MgCl2·6H2O, 26 mM NaHCO3, 10 mM sucrose, 10 mM trehalose, and 10 mM glucose; all from Sigma in the highest purity). This prevented the brain from desiccating, and the surface tension of the AHL on the hydrophobic fly cuticle prevented AHL from spilling on the fly’s body. The entire surgery took ∼10 s.

Behavioral Testing. SI Materials and Methods provides details of the phototaxis studies.

Two-Photon Microscopy. To image the antennal lobe, we used a custom-built two-photon microscope and a 20× water-immersion objective (0.95 NA; Olympus). The excitation powers (920-nm wavelength) at the specimen were 3 mW and 15 mW for Figs. 1 L and M and 3, respectively. The pixel dwell time was 2 μs. For imaging mushroom bodies, we used a 40× water-immersion objective (1.0 NA; Zeiss) and a commercial two-photon microscope (Prairie Technologies). The excitation power (927 nm) at the specimen was 10–18 mW for Fig. 4. The pixel dwell time was 1.2 μs for all images.

Time-lapse imaging. Our protocol requires that to maximize reproducibility for a given laser microsurgery recipe (pulse energy, number of pulses, repetition rate, size of the window, and size of the window), the experimental conditions should vary as little as possible from fly to fly. Thus, we used flies of identical age (2-d-old mated females) and reared them under uniform growth conditions, with the same number of parent flies in each vial (10 female Oregon-R and 3 male flies).
We converted the raw 16-bit images to 32-bit (4 pixels × 256 pixel images) or 4 pixels (512 × 512 images). We additionally applied a temporal smoothing filter (sliding window of one frame) to the mushroom body images.

To analyze the time-lapse Ca²⁺-imaging data in the mushroom bodies, for each session we computed the ΔF/ΔF₀ traces. We median filtered (3 × 3 array) the ΔF/ΔF data and applied a uniform threshold to discard pixels at or below background fluorescence levels. In Fig. 4, pixels with >30% ΔF/ΔF odor responses are superimposed on the basal fluorescence images. We manually selected the three cells with the largest odor-evoked fluorescence signals (ΔF/ΔF) for each image sequence. We used a 6 × 6 pixel window and a custom MATLAB script to average the signals in this window at each frame across an odor delivery trial. This window size helped average out pixel noise within a frame but remained within the area of one cell (~20 × 20 pixels). All calcium-imaging experiments used four odor puff cycles, and we considered responses as valid only if they were present in at least two cycles. For z stacks, we averaged eight frames per slice. We performed all statistical tests in MATLAB (MathWorks).

**Mice.** The Stanford Administrative Panel on Laboratory Animal Care approved all procedures. See SI Materials and Methods for surgical details.

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