

# Battery-free, lightweight, injectable microsystem for in vivo wireless pharmacology and optogenetics

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Pharmacology and optogenetics are widely used in neuroscience research to study the central and peripheral nervous systems. While both approaches allow for sophisticated studies of neural circuitry, continued advances are, in part, hampered by technology limitations associated with requirements for physical tethers that connect external equipment to rigid probes inserted into delicate regions of the brain. The results can lead to tissue damage and alterations in behavioral tasks and natural movements, with additional difficulties in use for studies that involve social interactions and/or motions in complex 3-dimensional environments. These disadvantages are particularly pronounced in research that demands combined optogenetic and pharmacological functions in a single experiment. Here, we present a lightweight, wireless, battery-free injectable microsystem that combines soft microfluidic and microscale inorganic light-emitting diode probes for programmable pharmacology and optogenetics, designed to offer the features of drug refillability and adjustable flow rates, together with programmable control over the temporal profiles. The technology has potential for large-scale manufacturing and broad distribution to the neuroscience community, with capabilities in targeting specific neuronal populations in freely moving animals. In addition, the same platform can easily be adapted for a wide range of other types of passive or active electronic functions, including electrical stimulation.

### neuroscience | pharmacology | optogenetics

**N** eurological diseases affect the quality of life of millions of people worldwide (1). Emerging tools for the manipulation of neural activity in the deep brain create important avenues for basic research into the operation of neural circuits and the development of associated insights for the underlying causes and treatments of neurological diseases (1, 2). Pharmacology and optogenetics represent two highly informative and widely used approaches in neuroscience research (3–7). While facets of each technique have improved (e.g., more-selective drugs, uncaged ligands, light-sensitive proteins with varied on/off kinetics), the chronic implant and its connection to a secondary tethered system for drug/light delivery have effectively remained unchanged (3). The hardware associated with these approaches can 1) in-

troduce damage to fragile tissues during implantation, 2) result in irritation at the device/tissue interface, 3) affect studies of natural behaviors of untethered, freely moving animals, 4) frustrate investigations of social interactions due to entanglements, and 5) prevent

### Significance

Neuroscience studies using optogenetics have greatly improved our understanding of brain circuits. Advances in the combined use of optogenetics and pharmacology to further probe important neurochemical signals has lagged, however, in large part due to the inconvenience of conventional cannulated approaches, as well as the difficulty in controlling, powering, and manufacturing optofluidic devices that are reliable and scalable for distribution to the neuroscience community. Here, we present a battery-free, wireless, lightweight optofluidic device that allows adjustable infusion rates, hands-free operation, and unlimited power supply, and is compatible with existing near-field communication (NFC) technology. We show that this device not only reproduces optogenetic and pharmacological experiments in vivo but, additionally, allows far greater freedom of movement in small animal models compared to existing, comparable platforms.

Competing interest statement: M.R.B., J.A.R., and A.B. are cofounders in a company, Neurolux, Inc., that offers related technology products to the neuroscience community.

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movements in complex 3-dimensional environments (8–12). These drawbacks, coupled with recent advances in flexible electronics and microfluidics (13–15), provide strong motivation for the development of innovative engineering platforms to improve experimental fidelity and expand the possibilities for advanced brain research studies in awake, behaving animals.

In addition to improving the technology for optogenetic and pharmacological studies, there is a growing interest in combining different or multiple methodologies for neuronal circuit manipulation. For example, optogenetics and fiber photometry or miniaturized microscopy combine light and genetics to both control and monitor neurons in vivo (16). Studies using such techniques indicate that different optogenetic stimulation parameters engage different neurotransmitter systems (17), and modulate different neuronal ensembles during discrete behavioral epochs (18). Although sophisticated and powerful in dissecting circuit activity, these combined platforms are constrained by the inability to cohesively incorporate pharmacology, thereby leaving many of the neurochemical mechanisms driving changes in behavior or neural activity unresolved (16). This limitation can be overcome by using systems that combine independently controlled pharmacological and optogenetic functions within a single platform. Ideally, such devices would be wirelessly controlled, programmable, soft, and lightweight to adapt to any experimental design while also avoiding damage to brain tissue or disruption of behaviors of freely moving animals.

In 2015, we introduced a wirelessly controlled optofluidic device for delivery of both fluidic compounds and optogenetic stimulation (19). Although an important advance, the device retained several nonideal features, including a battery-powered, thermal-mechanical micropump, nonnegligible heat generation (~87 °C) during the operation, limited lifetimes, inability to reuse the device, complexity in system design, and absence of accessibility beyond specially trained laboratories. Recent advances offer options in wireless drug delivery, with or without optogenetic functionality, that bypass certain drawbacks associated with the traditional cannula approach, but most are still constrained by requirements for 1) rigid stainless steel channels and large feature sizes (weight > 6.6 g; dimension > 14.4 mm  $\times$  15 mm  $\times$  24.8 mm) and 2) wireless operation in radio frequency (RF) bands that are sensitive to signal reflection, interference, and absorption by biological tissue, metallic objects, water features, and other obstructions (6, 20).

Recently, we introduced a soft, fully implantable optofluidic cuff system designed to deploy on peripheral nerves, with ultralow power requirements by use of miniaturized electrochemical micropumps for drug delivery (21). Here, we improve upon and adapt those ideas to establish a functional device that is able to interface with the brain. The resulting system maintains the same ultralow power operation and wireless, battery-free functionality, but also includes additional capabilities in drug refillability and programmable control over flow rate. Lastly, the design of the device is compatible with existing large-scale manufacturing practice, further underscoring its potential for widespread use across the neuroscience community. The system is 1) refillable and reusable; 2) capable of independent, multimodal operation through the infusion of multiple types of drugs and/or the delivery of optical stimulation; 3) lightweight (0.29 g) and compact in its construction; and 4) entirely wireless, with battery-free operation, thereby bypassing the physical burden normally imposed by larger implantable devices. The results described in the following demonstrate multiple ways in which this easy-to-use device and control platform can be broadly applied to behavioral conditions used in modern neuroscience research. Collectively, this device represents an important advance in wireless optofluidic technology for neuroscience in a format that is accessible and practical.

# Results

Device Design and Operational Features. The microsystem detailed here includes a wireless, battery-free electronics module, 4 miniaturized electrochemical micropumps and reservoirs, a thin (~100 µm), soft (modulus ~3 MPa) microfluidic probe for fluid delivery, and a microscale inorganic light-emitting diode (referred to here as  $\mu$ -ILED, 270  $\mu$ m  $\times$  220  $\mu$ m  $\times$  50  $\mu$ m) light source for optogenetics (Fig. 1). A refilling port (diameter  $\sim$ 350 µm) on the sidewall of each reservoir allows for multiple fluidic infusions with the same device. Fig. 1A and B presents schematic diagrams of a complete system that integrates each of these components together with a magnetic loop antenna for wireless power harvesting via magnetic resonant coupling protocols. The design is fully compatible with our recent wireless electronic platform and control software for the optogenetics (22). Fig. 1 C and Dshows optical images of front side and backside views. Fig. 1E presents a scanning electron microscope (SEM) image of the injectable tip end of a probe that includes one µ-ILED for optical stimulation/inhibition and 4 microchannels for the programmable delivery of up to 4 drugs. Demonstration experiments using colored dyes and phantom brain tissues (0.6% agarose gel) illustrate the operation. Following a wirelessly triggered event, the microcontroller (µC) in the electronics module activates a selected electrochemical micropump to initiate fluidic delivery or the µ-ILED for optogenetic stimulation/inhibition, or both simultaneously (Fig. 1 F-H). A red indicator LED provides a convenient indication for the operation of device. The miniaturized dimensions (radius 0.5 cm; thickness ~0.4 cm), the lightweight construction (~0.29 g; SI Ap*pendix*, Fig. S1 A and B), wireless, programmable operation, and the mechanical compliance of the probes represent key characteristics, as detailed below.

Magnetic resonant coupling at 13.56 MHz transfers power wirelessly to the device from a primary antenna that encircles the area of interest (Fig. 11), thereby eliminating the need for a battery and its associated bulk and weight. These magnetically coupled antennas operate in the high-frequency (HF, 3 MHz to 30 MHz) band to enable full wireless coverage across many cage types and environments, with little sensitivity to the presence of objects or physical obstructions, including those environments that involve metal plates, cages, or meshes or those with water pools, baths, or mazes (22). This same wireless link supports a programmable control strategy through a graphical user interface on a computer. Wireless control relies on a custom protocol based on on-off keying, as schematically outlined in Fig. 11, with a predetermined parameter set, as described in detail in ref. 23. This scheme enables digital control over optogenetic stimulation frequency, duty cycle, and pumping parameters. SI Appendix, Fig. S1 C and D shows the complete system, which includes a computer with customized control software, a multicomponent power distribution control box, an antenna tuner, and a customizable transmission antenna.

**Electrical, Fluidic, and Optical Characterizations.** A thin, flexible printed circuit board (PCB) supports the electronic components for this system, including a  $\mu$ C, capacitors, rectifiers, and an RF antenna (Fig. 1 *B* and *C* and *SI Appendix*, Fig. S2 and Table S1). Here, a capacitor of 75 pF tunes the secondary antenna to 13.56 MHz (*SI Appendix*, Fig. S3*A*). *SI Appendix*, Fig. S3*B* shows the rectification behavior with decreasing ohmic loads. Peak power of over 18 mW is achieved at a voltage of 11 V and a load of 6.8 kohm at the center of a home cage with 6W input power. Harvested power changes with relative angle to the primary coil, shown by the unrectified power decreasing with increasing angle in *SI Appendix*, Fig. S3*C*. Fig. 2*A* shows the power distribution as a function of position in a 25 × 25 cm conditional place preference (CPP) cage powered at 6 W at physiologically relevant heights of 3 and 6 cm. Power distribution for a home cage powered at 6 W is

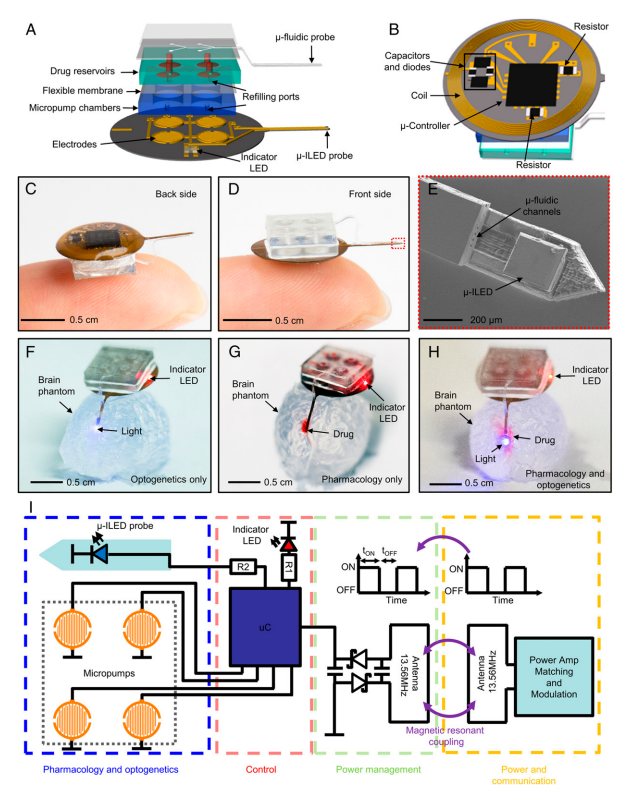
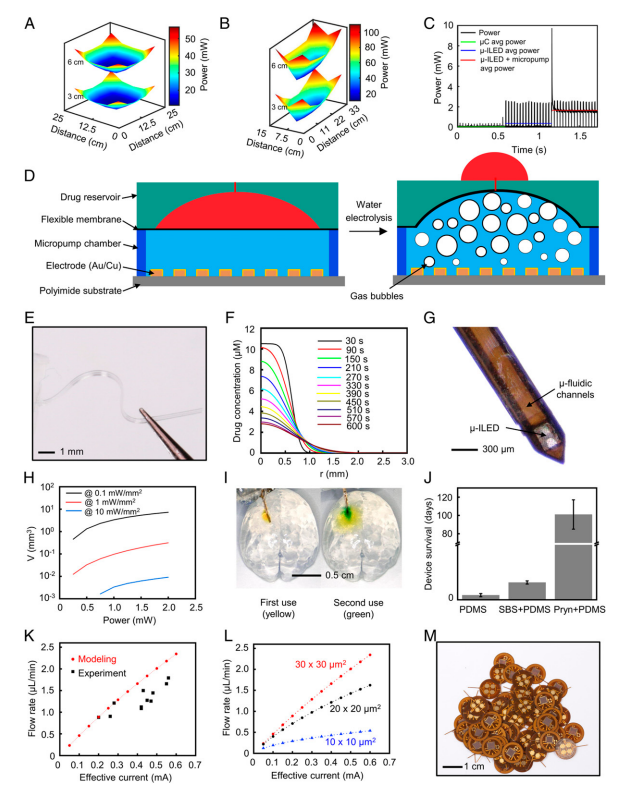


Fig. 1. Wireless, battery-free, injectable microsystems for programmable pharmacology and optogenetics. (A and B) Exploded view illustration and schematic diagram of the system. (C and D) Photographic images of the back and front sides of the device. (E) A scanning electron micrograph of the tip end of the injectable probe, showing the microfluidic channels for pharmacology and a  $\mu$ -ILED for optogenetics. (F–H) Demonstration of wireless fluid delivery and optical stimulation in a brain tissue phantom. (I) Schematic diagram of electronics design and wireless communication schemes.

also in Fig. 2*B*. Minimum power outputs of 11 and 18 mW occur at the center of the CPP cage and home cage, respectively. The power distributions for lower RF powers are shown in *SI Appendix*, Fig. S3 *D* and *E*. Fig. 2*C* summarizes a characterization study of the power consumption of the device. Here power consumption varies with time and is modulated by duty cycle and modality. Fig. 2*C* shows data for 3 states: low power mode of the  $\mu$ C (0.04 mW and peak power of 0.40 mW),  $\mu$ -ILED active (0.38 mW and



**Fig. 2.** Electrical, optical and fluidic characterization results. (*A*) Rectified power at the implant measured in a 25 cm  $\times$  25 cm behavioral chamber with 6W input power at implant heights of 3 and 6 cm with respect to cage floor. (*B*) Rectified power at the implant measured in a 15 cm  $\times$  33 cm home cage with 6W input power at implant heights of 3 and 6 cm with respect to cage floor. (*C*) Power consumption of implant with input voltage of 3.3 V for multiple states including a low-power  $\mu$ C idle state,  $\mu$ -ILED with 10 mW/mm<sup>2</sup> irradiance output at 30 Hz, and simultaneous  $\mu$ -ILED and drug micropump activation. (*D*) Schematic diagram of the electrochemical micropump based on water electrolysis. (*E*) Picture that illustrates the soft, compliant mechanics of the microfluidic probe. (*F*) Distribution of drug concentration along radial direction for t = 30, 90, 150, 210, 270, 330, 390, 450, 510, 570, and 600 s. (*G*) Optical micrograph of an integrated optofluidic probe. (*H*) Effective illumination volume at 3 different irradiance thresholds and input optical powers from 0.25 mW to 2 mW. (*l*) First and second use of a device, demonstrating the ability to refill and reuse the microfluidic system. (*J*) Device survival lifetime in 1× PBS at 37 °C when encapsulated with a monolayer of PDMS alone, a bilayer of SBS and PDMS, or a bilayer of PDMS. (*K*) Maximum flow rate in a microfluidic channel (cross-sectional area of microchannel  $\sim 30 \times 30 \, \mu$ <sup>m<sup>2</sup></sup>) as a function of effective current. (*L*) Geometry-dependent maximum flow rate in microfluidic channels as a function of effective current. (*M*) Picture of devices manufactured using flexible PCB technology.

peak power of 2.5 mW with a duty cycle of 13.5%, and stimulation frequency 30 Hz), and micropump in conjunction with the µ-ILED (1.66 mW and peak power of 2.5 mW, µ-ILED duty cycle 13.5%, micropump duty cycle 86.5%, and stimulation frequency 30 Hz). The initial power consumption peak when activating the micropump can be attributed to the initiation of the electrochemical process and the capacitive nature of the interdigitated electrode structure. The average power consumption is well below harvesting capabilities for both the  $25 \times 25$  cm CPP cage and the home cage, resulting in a tolerance to conditions such as rearing, which results in an increased implant angle with respect to the primary antenna. For experiments where small variations in output power and system voltage are not tolerable, an actively regulated system can be achieved via the integration of a low dropout voltage linear regulator (23). An example is shown in SI Appendix, Fig. S3F, where the output voltage with a load that represents the device under simultaneous electrochemical pumping and µ-ILED stimulation.

The miniaturized electrochemical micropump serves as an alternative to previously reported thermally activated versions (19, 21, 24–27). This approach allows for low power operation (<1 mW), negligible heat generation (<0.2 °C), high delivery efficiency (~90%) of the fluid housed in the reservoirs, simple construction, and large driving force (21). Each reservoir has an associated pumping chamber (cylinder shape; height 1 mm; diameter 2.45 mm, SI Appendix, Fig. S4), a set of interdigitated electrodes (Au/Cu, thickness 200 nm/18 µm, ~50 µm width, ~50 µm space), and a flexible separating membrane of polystyrene-blockpolybutadiene-block-polystyrene (SBS) (Fig. 2D). At room temperature, pressure from the water hydrolysis reaction deforms a flexible membrane, thereby driving the flow of pharmacological agents out of the reservoir to the targeted area of the brain through corresponding microfluidic channels (Fig. 2D). The micropump chamber and reservoir are both milled from a block of cyclic olefin copolymer (thickness: 1 mm), selected for its efficiency as a moisture barrier (water vapor permeability ~0.045 g·mm/m<sup>2</sup>·d at 23 °C and 85% relative humidity) and resistance to hydrolysis, acids, alkalis, and polar solvents such as methanol, ethanol, acetone, and isopropanol (28), thereby allowing use of drugs that are soluble in water, dimethyl sulfoxide, ethanol or sodium hydroxide (50%). A bilayer of Ti/SiO<sub>2</sub> (5 nm/20 nm) deposited by sputtering (AJA International) renders the surface of the SBS film hydrophilic. The positively charged drug molecules may attach on the surface of the drug reservoir because of electrostatic interactions. This issue can be solved by functionalizing with a positive amine group.

Here, the flexible membrane of SBS offers a combination of soft mechanical characteristics (modulus ~13 MPa) and effective barrier properties (water permeability  $8.52 \times 10^{-8}$  g·m/m<sup>2</sup>·h·Pa; oxygen permeability 11.88 barrier; hydrogen permeability 21.66 barrier) (21). Each microchannel lies along the length of a thin ( $\sim 100 \,\mu$ m), mechanically soft, and flexible (modulus  $\sim 3 \,\text{MPa}$ ) probe (4 channels, each with  $10 \times 10 \,\mu\text{m}^2$  cross-sectional areas) for minimally invasive, targeted fluid delivery into targeted regions of the deep brain (Fig. 2E and SI Appendix, Fig. S5 A and B). Previous studies quantified the "functional" spread of a drug by measuring changes in Fos activity (29-32). SI Appendix, Table S2 shows a few examples of this analysis for injection volumes sized similarly to our device. Overall, the functional diffusion volume is in the range between 0.05 and 3.92 mm<sup>3</sup>. An analytical model of drug diffusion into the brain tissue following wireless infusion captures the physics. The injected drug is modeled to have initial uniform distribution with concentration  $C_0$ in a sphere of radius  $r_0$  in the brain tissue. With an extracellular space fraction of the brain of  $\alpha = 0.21$  (33), and injected volume of 0.25 µL at 50 µM concentration of drug in a typical pharmacology study,  $C_0 = 10.5 \,\mu\text{M}$  and  $r_0 = 0.66 \,\text{mm}$ . With a given diffusion coefficient, the model provides a quantitative prediction of the distribution of drug concentration after wireless microfluidic delivery. For diffusion of small molecules, Fig. 2*F* shows the drug concentration distributed along the radial direction at t = 30, 90, 150, 210, 270, 330, 390, 450, 510, 570, and 600 s for D-2-amino-5-phosphonovalerate acid (D-APV), a widely used competitive *N*methyl-D-aspartate (NMDA) antagonist, with apparent diffusion coefficient  $D = 2.8 \times 10^{-6}$  cm<sup>2</sup>·s<sup>-1</sup> (34), and the contours of concentration are shown in *SI Appendix*, Fig. S5*D* and *E* show the diffusion of NGF, a 26.5-kDa nerve growth factor protein, with apparent diffusion coefficient  $D = 2.95 \times 10^{-7}$  cm<sup>2</sup>·s<sup>-1</sup> (35). Integration of the probe into these calculations will yield different distributions of drug concentration, as the substrate of the probe prevents the diffusion to its backside (Fig. 2*G*).

The soft microfluidic probe has a width of ~350 µm (crosssectional area: 0.035 mm<sup>2</sup>). These dimensions are much smaller than those of conventional 26-gauge metal cannulas, where the diameters are 460  $\mu$ m (cross-sectional area: 0.166 mm<sup>2</sup>). The probe also integrates an µ-ILED system with similar overall geometry to provide functionality relevant for optogenetics (22), in vivo optopharmacology, or other applications that require light sources (36, 37) (Figs. 1E and 2G). RF coupling of the primary antenna to the secondary antenna on the injectable microsystem results in harvested electrical powers ranging from 11 mW to 115 mW (Fig. 2A and B). The harvested power operates the  $\mu$ C and  $\mu$ -ILED, which typically requires an average electrical power of 0.38 mW (Fig. 2C). Electrical power to the µ-ILED is regulated through a current limiting resistor, resulting in a controlled conversion to optical stimulation output (23). In this configuration, optical powers of 0.25 mW to 2 mW can be realized, with corresponding irradiances ranging from 4.21 mW/mm<sup>2</sup> to 33.67 mW/mm<sup>2</sup>, at the  $\mu$ -ILED's surface, and produce illumination volumes from 0.012 mm<sup>3</sup> to 0.315 mm<sup>3</sup>, respectively, at 1 mW/mm<sup>2</sup> illumination threshold (Fig. 2H and SI Appendix, Fig. S6 and Table S3 provide µ-ILED illumination volume of brain tissue predicted by Monte Carlo simulations). Irradiance output is well within typical optogenetic parameters of 1 mW/mm<sup>2</sup> to 10 mW/mm<sup>2</sup>.

Drug Refillability. Fig. 2I highlights the ability to refill and reuse the reservoir. The electrochemical pumping mechanism and refilling ports on the sides of the reservoirs allow multiple cycles of use, thereby enabling studies over long periods of time with multiple drug delivery events. Here, a Hamilton syringe or 30G needle attached to a 1-mL syringe allows loading of electrolyte (50 mM potassium hydroxide, KOH, or 50 mM sodium hydroxide, NaOH) and drug into the micropump chamber and reservoir, respectively, through the refilling ports (SI Appendix, Fig. S7). Sealing the ports with a removable layer of silicone elastomer (Kwik-Sil; World Precision Instruments) or paraffin wax prevents leakage or evaporation. The power consumption-voltage characteristics and the maximum displacement of the flexible membrane serve as additional means to quantify the performance after refilling and resealing. The results, shown in SI Appendix, Figs. S8 and S9, indicate negligible difference (<10%) for the first, second, and third uses. The flow rate depends on the power applied to the electrochemical micropumps (power consumptionvoltage characteristics), the deformation of flexible membrane, and the sizes of microfluidic channels. Since the refilling does not change the geometry of the microfluidic channels, these measurements suggest a stable flow rate of devices for reuse.

**Device Encapsulation.** A bilayer coating of Parylene C and polydimethylsiloxane (PDMS) encapsulates the associated wireless electronics and  $\mu$ -ILED probe. Fig. 2*J* shows the survival lifetime of injectable part of the  $\mu$ -ILED probe (6 mm long, adjustable to match the targeted brain region) in 1× phosphate-buffered saline (PBS) at 37 °C when encapsulated with a single layer of PDMS (dip-coating, mixed at 10:1 ratio, ~20  $\mu$ m thick), a bilayer of SBS (4 g/40 ml in toluene, dip-coating, ~20  $\mu$ m thick) and PDMS (dip-coating, mixed at 10:1 ratio, ~20  $\mu$ m thick), and a bilayer of Parylene C (Pryn, chemical vapor deposition, ~5  $\mu$ m thick) and PDMS (dip-coating, mixed at 10:1 ratio, ~20  $\mu$ m thick). Although ultrathin, transferred layers of thermally grown silicon dioxide can provide superior barrier characteristics under physiological conditions (38, 39), an encapsulation lifetime of about 100 d is sufficient for most optogenetic studies, considering the optimal period of testing for virally expressed opsins and age of the test animal.

**Programmable Control over Flow Rate.** Control of the rates of delivery can be important in many neuroscience studies. Excessively high rates may induce tissue damage, while insufficient rates may increase the potential for blockage at the exit ports of soft microfluidic channels. The rates of gas generation during electrochemical pumping determine the delivery rates. The pumping rate under constant current operation can be roughly written as (40)

$$Q = \frac{3}{4} \frac{i}{F} V_m, \qquad [1]$$

where i is the current applied to the electrodes, F is Faraday's constant, and  $V_m$  is the molar gas volume at 25 °C and atmospheric pressure. This equation suggests that the rate can be controlled most easily through the time-averaged applied current (effective current), or, more specifically, in practice, the duty cycle of the pulse width modulation current. However, due to 1) the flow resistance in microfluidic channel, and 2) the resistance from the stiffness of the flexible membrane, the actual flow rate is typically lower than the value predicted by Eq. 1, and a more accurate model must be used to predict the pumping rate. Fig. 2K shows the maximum flow rates of a device inserted into a brain phantom (0.6% agarose gel) and operated at 3 V as the wireless transmitted power serves as a constant voltage source, with pulses at 490 Hz and various time-averaged applied currents. In modeling, constant time-averaged applied current is assumed during the operation of the electrochemical pumping process, and the predicted flow rate matches well with experiments when small current is applied. However, the gas bubbles generated during the electrochemical pumping reduce the contact area between the electrolyte solution and the interdigitated electrodes, thereby causing a decrease in applied current and, therefore, a mismatch between experiment and modeling at higher applied currents (Fig. 2K and SI Appendix, Table S4). SI Appendix, Table S5 shows the relationship between applied current and time required to reach maximum flow rate or total time to deliver >80% of fluid housed in the reservoir, which could be important for time-sensitive experiments. Importantly, the maximum infusion rates also depend on channel size, which can be customized (Fig. 2L). We note that microfluidic channel sizes of  $30 \times 30 \ \mu m^2$  (cross-sectional areas) minimize the potential for blockage in in vivo studies.

**Compatibility with Large-Scale Manufacturing Practice.** The components and systems described previously form the foundations for a manufacturable, low-cost device for wireless programmable pharmacology and optogenetics that can be used by neuroscientists with little or no background in RF electronics. The designs align with standard manufacturing practice in electronics, fluidics, and microsystems technologies. Specifically, the fabrication exploits scalable techniques such as laser structuring and microfabrication, using equipment readily available in academic laboratories. Manufacturing of each subsystem can occur in an automated process using flexible PCB technology and microfluidics. Results of an exploratory manufacturing procedure for the wireless power

supply, control electronics, and electrodes for the electrochemical micropumps appear in Fig. 2*M* and *SI Appendix*, Fig. S10.

Implanting Devices in the Brain. The process for surgical insertion of the device is similar to that of a conventional guide cannula or optic fiber. SI Appendix, Fig. S11 illustrates the steps for implanting the needle portion of the device into the brain and fixing the body of the device to the top of the skull. Briefly, an incision is made on top of the skull, and a small craniotomy is performed to allow probe insertion. The top of the device is mounted to a stereotaxic arm with a flat surface, and gently lowered into the brain. A small amount of medical device adhesive (Prism Medical Device Adhesive, 4541; Loctite) is applied near the point of insertion, bonding the needle to the skull. When the adhesive is fully hardened, the stereotaxic arm is raised while the secured device remains attached to the skull. SI Appendix, Fig. S11 G and H shows that the mice are in good health and that the devices cause no obvious adverse effects at 3 d and 30 d after surgery, respectively. Implantation does not disrupt normal locomotion, even as soon as 3 d after surgery. The high degree of mechanical compliance and the minimal tissue displacement associated with the probes are highlighted via some reduced lesioning and immunoreactive glial responses compared to those associated with conventional metal cannulas (Fig. 3). These findings are consistent with those of other thin, flexible deep brain implants (7, 19).

Wireless Pharmacology. This system operates with a remotely controlled interface to independently infuse multiple fluids in various environments, such as home cages or behavioral testing chambers. To test the operability of these devices in the deep brain across multiple test sessions, we targeted the brainstem ventral tegmental area (VTA) of wild-type (C57BL/6J) mice (Fig. 4A). Previous studies show that unilateral opioid stimulation in the VTA causes a stereotypical increase in locomotion and/or turning behavior in which the animals spin 360° in the contralateral direction relative to the hemisphere of injection (Fig. 4B) (19). We recapitulated these studies by wirelessly triggering the delivery of artificial cerebrospinal fluid (ACSF) as a vehicle control or the mu opioid receptor agonist [D-Ala<sup>2</sup>,N-Me-Phe<sup>4</sup>,Gly-ol<sup>5</sup>]-enkephalin (DAMGO) into mice 7 d following surgical implantation. Drug conditions were counterbalanced across mice, and infused via the same drug reservoir. While the device has the capability to infuse up to 4 drugs through its 4 separate reservoirs, we chose to demonstrate the reusability and refillability of the reservoirs by using the same reservoir for both test days. Specifically, mice were gently placed into a home cage-sized chamber and immediately received a wireless infusion of the ACSF or DAMGO. Mice were allowed to freely move in the chamber for the duration of the infusion ( $\sim 2 \text{ min}$ ). After 4 min in the chamber (to allow full diffusion of the drug), mice were moved directly into an open field chamber (50 cm  $\times$ 50 cm) where they were allowed to freely explore for 1 h (Fig. 4C). At the end of the hour session, mice were removed from the open field and placed back into their home cage. To record locomotor and rotation behaviors, a camera was mounted to the ceiling, and total distance traveled was recorded. Total number of contralateral turns was scored offline by an observer blind to the test condition. There was no obvious change in the overall locomotor behavior after vehicle or DAMGO infusions (Fig. 4D). As predicted, however, DAMGO stimulation led to a doubling of the number of stereotypical rotations compared to the ACSF test day (Fig. 4E) (Wilcoxon, P = 0.031, Cohen's d = -0.64). While previous demonstrations of this experiment have shown both locomotor and rotation effects of DAMGO (19), the differences here could be related to the lightweight nature of this device (0.29 g vs. 1.855 g). Previous reports using a battery-operated optofluidic device showed substantially lower baseline locomotion compared

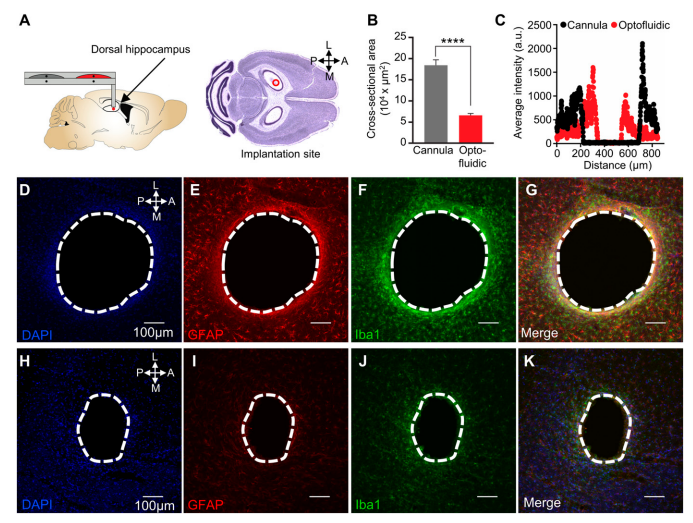


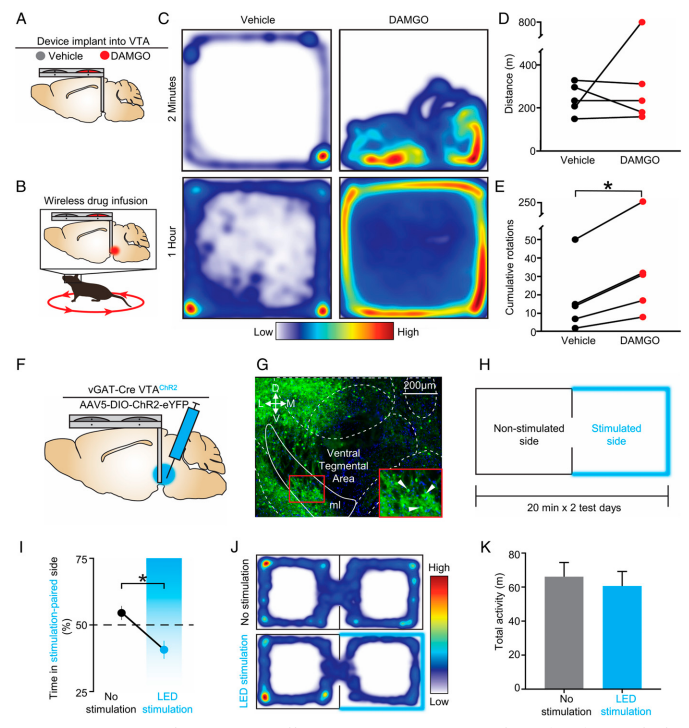
Fig. 3. Device implantation impact on the brain. (A) Schematic and mouse brain library atlas. Red spot indicates approximate implantation site. (B) Mean  $\pm$  SEM cross-sectional area of lesions from cannulas and optofluidic probes (Student's t test, \*\*\*\*P < 0.0001, n = 3/group). (C) Representative linescan of fluorescence intensity from cannula (black) and optofluidic probe (red) lesions. (D-K) Representative confocal fluorescence images of horizontal striatal slices show immunohistochemical staining for DAPI (blue), astrocytes (GFAP, red), and activated microglia (lba1, green) and overall lesion from a cannula and an optofluidic device. All histological and confocal settings were kept consistent across groups. Lesioned tissue from either cannula (*Top*) or device (*Bottom*) implantation is outlined in white dashed lines. (Scale bars, 100  $\mu$ m.)

to this device (19), and even the DAMGO stimulated locomotion often only reached the baseline locomotor activity observed in mice implanted with this device. These differences highlight how the weight of a head-mounted device can dramatically affect normal behavior.

Wireless Optogenetics. To evaluate whether the optogenetic capabilities of the device would also work in deep tissue, we stimulated local GABAergic neurons in VTA expressing channelrhodopsin (ChR2) in a real-time place test (RTPT) (Fig. 4 F and G) (41). Previous reports have shown that GABA neuron stimulation is aversive, reduces responding for rewards, and is actively avoided (41, 42). First, vGAT-cre mice received an injection of AAV5-EF1a-DIO-hCHR2-(H134R)-eYFP into the VTA. Two weeks later, mice were implanted with the device, then allowed to recover for 1 week before behavioral testing. Mice were tested in a 20-min RTPT, in which they were able to freely roam between 2 indistinguishable chambers (Fig. 4H). Each mouse was tested twice. On one test day, they never received stimulation. On the other test day, they received stimulation in one chamber, but not the other. Time spent in each chamber was calculated to determine an overall preference score for each day. As expected, mice

showed a robust avoidance of the stimulation-paired chamber, spending ~15% less time in the stimulation side compared to the nonstimulated test day (Wilcoxon test, P = 0.031, Cohen's d = 2.05) (Fig. 4 *I* and *J*). This aversion did not appear to be due to changes in locomotor activity between the stimulated and non-stimulated test days (total distance, P = 0.103; velocity, P = 0.103) (Fig. 4K). Like the DAMGO-stimulated rotations experiment, these mice showed demonstrably higher general locomotion compared to previous wireless optofluidic models, and results are comparable to traditional tethered optogenetic studies (19, 41). These results demonstrate that the wireless optofluidic devices can activate neurons in vivo while mice perform a frequently used behavioral assay.

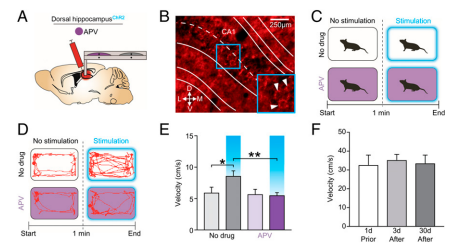
**Combining Wireless Pharmacology and Optogenetics in Vivo.** One of the most compelling uses for this optofluidic device is to combine pharmacology and optogenetics. Here, we sought to determine whether this device was capable of effectively integrating these two approaches. We first injected a panneuronal channelrhodopsin expressing virus into dorsal hippocampus (DH) (Fig. 5 A and B). Three weeks later, we implanted the device into the same site. After 3 d of recovery from surgical implantation, animals



**Fig. 4.** Deep tissue demonstrations of in vivo wireless delivery of fluidic compounds and optogenetic stimulation of neurons. (*A* and *B*) Schematic of the fluid delivery and subsequent contraversive rotations. (*C*) Representative traces of movement from the same animal over the course of 2 min and 1 h after either vehicle or DAMGO infusions. (*D*) Overall locomotor behavior after vehicle or DAMGO infusions. (*E*) Cumulative contraversive rotations after vehicle or DAMGO infusions (F < 0.05 Wilcoxon test). (*F*) Schematic of viral injection of channelrhodopsin into vGAT-cre mice. (*G*) Representative image showing robust viral infection in VTA. White arrows indicate individual infected neurons. (*H*) Schematic of Real-Time Place Test. (*I*) Optogenetic stimulation of VTA GABA neurons causes a place avoidance (F < 0.05 Wilcoxon test). (*J*) Representative trace of cumulative time spent in each zone on test day with no stimulation (*Top*) versus stimulation (*Bottom*). (*K*) Optogenetic simulation did not disrupt overall locomotor activity.

were habituated in a chamber similar in size to their home cage, in which they were able to freely explore for 2 min (day 1). On day 2, the animals were again placed in the chamber and allowed to explore for 2 min. At the onset of the second minute, mice received optical stimulation for 1 min, and locomotor activity was recorded (Fig. 5*C*). Here we show that 4-Hz stimulation of DH results in a significant increase in locomotion (*t* test, P = 0.023) (Fig. 5 *D* and *E*). In a subsequent, counterbalanced experiment, wireless pharmacological delivery of an NMDA receptor antagonist (APV) was given prior to the photostimulation. We found that pretreatment with





**Fig. 5.** Optogenetic stimulation of hippocampal neurons increases locomotion, and is blocked by the wireless infusion of the NMDA receptor antagonist APV. (A) Schematics of viral injection and the implantation of optofluidic device. (B) Fluorescent image showing expression of pAAV.CAG.hChR2(H134R)mCherry.WPRE.SV40 in DH. Solid lines depict outline of hippocampal tissue, and dashed line indicates subdivisions within CA1. (*Inset*) Blue box in center of image is enlarged in right corner, with white arrows pointing to individual infected neurons. (C) Schematic of locomotor test, with and without optical stimulation or APV infusion (D) Representative traces of mice across experimental conditions. (E) The 4-Hz optical stimulation of DH increases locomotion (\*P < 0.05, t test), and is blocked by the wireless infusion of APV (\*\*P < 0.01, t test), n = 7 to 8. (F) Baseline locomotor activity prior to and after device implantation. Implantation does not disrupt normal locomotion, even as soon as 3 d after surgery (n = 4).

APV (wirelessly infused via the device) completely blocked optical stimulation-induced increases in locomotor activity (paired *t* test, P = 0.0071) (Fig. 5 *D* and *E*). These results are consistent with previous reports that photostimulation of neurons in DH causes a glutamate-dependent increase in locomotion (43–45), demonstrating that this device can effectively replicate a well-known optogenetic and pharmacological phenomenon. Notably, baseline locomotor activity was not affected by implantation of the device, even as soon as 3 d after surgery, nor were there any discernable deficits observed when behavior was recorded 30 d later (Fig. 5*F*). These results indicate that the optogenetically induced locomotion and the subsequent APV blockade function as expected, with no unusual or disruptive features preventing mice from behaving similarly to fiber or cannula implanted mice.

# Discussion

The wireless, battery-free, injectable microsystem presented here represents a versatile technology for manipulation of deep brain tissue in freely moving animals. The platform offers multifunctional capabilities in a single, soft implant with options for in vivo wireless pharmacology and optogenetics, many of which would be impossible to reproduce with conventional metal cannulas and/or optical fibers (37). There are several benefits of this platform compared to existing technologies (SI Appendix, Table S6). First, fully wireless and battery-free operation eliminates the need for external or bulky connections to provide power for fluid or light delivery. The device is powered, controlled, and operated within the industrial, scientific, and medical radio band, and is highly compatible with widely available 13.56-MHz technology, thereby facilitating worldwide dissemination. The HF (3 MHz to 30 MHz) band antenna designs can be easily operated inside almost any environmental arena by novice operators, and are largely unaffected by the presence of objects or physical obstructions (e.g., walls, novel objects, bedding). In effect, this type of technology allows the device to be as functional as any currently available optical stimulation system (22, 42, 46-48), with the additional option of complementary pharmacology. Finally, since this device is directly and continuously powered through near-field communication (NFC) technology, there is no limit on power lifetime, thereby eliminating concerns about battery life/charging status during behavioral test sessions. This feature is especially

useful for studies that have extended time courses, such as overnight sleep studies or long-access ( $\sim$ 6 h) self-administration sessions. Similarly, this device can be turned on and off at will or through transistor-transistor logic for more longitudinal studies without the need to interact with the animal at any time point during each test session.

Second, the lightweight construction (0.29 g, including reservoirs and micropumps, soft microfluidics, µ-ILED, and electronics for wireless power supply and control) bypasses physical constraints that can limit the natural movements of small animal models, allowing them to interact with the environment uninhibitedly. In fact, other drug delivery devices, with or without optogenetic functionality, are 6.4 to 22.8 times the weight of the device reported here (SI Appendix, Table S6), including the batteryoperated device we have previously published (19). Such freedom is critical for many behavioral neuroscience studies (drug self-administration, social interactions) (49), where performance can be disrupted by stressful distractions or impediments (e.g., tethers, heavy weights). The battery-free operation allows the platform to operate with a much smaller feature size and weight than with battery power (the primary power source in other commercially available devices). The removal of an external battery and net reduction in device size (SI Appendix, Table S6) also allows our device to be implanted flush against the skull. This option enables unimpeded entry into nose ports and reward magazines in operant chambers, which can be difficult with some tethered protocols (extended tethers can prevent animals from triggering infrared beams or reaching rewards). These devices also lend themselves to locomotor tasks that can be difficult to test with tethered or bulky head mounts, such as wheel running, rotarod, or gait assays. Indeed, we observed a large difference in baseline locomotion between this device and other batterypowered iterations, indicating that weight alone is sufficient to greatly influence behavior. Using lighter, less obtrusive devices, like the one described here, will allow for greater versatility in behavior/optofluidic experiments.

Third, the device design uses a thin ( $\sim 100 \ \mu m$ ), soft (modulus  $\sim 3$  MPa) microfluidic probe that avoids many of the issues in chronic inflammation and necrosis of neurons that are associated with traditional cannulas made of rigid materials like stainless steel (50). The thin geometry and soft mechanics adapt to brain

micromotions associated with physiological, behavioral, or mechanical sources (50). This behavior minimizes brain tissue damage during implantation and reduces the formation of glial scars for stable chronic operation (Fig. 3) (51). By our calculations, the area of lesioned tissue in a 26-gauge metal cannula that is 460  $\mu$ m in diameter is ~0.184 mm<sup>2</sup>, whereas the damage produced by our device is ~0.068 mm<sup>2</sup> (Fig. 3*B*). Perhaps even more striking, lesioned tissue from a thin optic fiber (~240  $\mu$ m in dimeter) results in ~0.045 mm<sup>2</sup>, indicating that our device produces comparable overall damage, but with double the functionality. Given that microglia can strongly impact neuronal activity (52, 53), features that minimize their recruitment, as found in our device, are critically important.

Fourth, multimodal operation permits independent infusions of multiple drugs or optical stimulation, for combined modes of use. The conventional approach in pharmacological and optogenetic experiments, such as optopharmacology, requires multiple tethers, tubing, and specialized commutators that can impede or make difficult otherwise straightforward experiments (37). With the system introduced here, separate tubing is no longer necessary, facilitating the execution and completion of behavioral studies. Eliminating tubes, injectors, commutators, and fiber optics reduces the number of steps, at which failure to perform can disrupt an experiment. Such a streamlined approach has gained traction in recent years, with several groups capitalizing on "singlestep" optogenetics to minimize potential failures and maximize usability (54). Fittingly, our device is also capable of infusing viral vectors, allowing for more targeted optogenetic approaches. With the 4 available reservoirs, it would even be possible to infuse multiple viruses or tracers to further gain information on circuit dynamics or connectivity, in conjunction with causal optogenetic stimulations. The field of optopharmacology has made great strides in recent years, not only with the number of available compounds but also with the versatility of these compounds (37, 55). These advances are especially relevant for peptide neurotransmitter systems, which have historically been difficult to quantify in vivo. While some advances have been made in observing temporally distinctive peptide events (36), subminute time scales remain out of reach. Optopharmacology provides one means to tackle this issue by allowing investigators to infuse an inactive compound, then experimentally induce a peptide signal at a specific behaviorally relevant moment with optical stimulation. As newer compounds continue to be developed (e.g., bistable uncaging/recaging), the ability to simulate neurotransmitter signaling patterns in vivo will allow for exciting new ways to investigate neural systems. Therefore, having a device with both fluidic and optical capabilities will allow for the effective use of these compounds in vivo.

The system presented here is refillable and reusable. This design represents a major distinction compared to other available devices, and will encourage the adoption of combined optogenetic and pharmacology approaches into mainstream experimental designs. Traditionally, intracranial pharmacology is accomplished via the insertion of a microinjector through chronically implanted stainless steel cannulas. For experiments that require multiple injections, each insertion is accompanied by fresh injury to the brain, thereby limiting the total number of injections that can be performed. Our system is only implanted once, and the filling/refilling of the device is performed through the ports on the side wall of the reservoirs. In other words, this device bypasses the detrimental effects of multiple instances of acute tissue trauma while preserving the security of a chronically implanted fluid delivery system. An additional advantage is in the electrochemical pumping mechanism coupled with reaccessible drug reservoirs. Unlike other systems, in which experimenters are limited in the total number of infusions (e.g., traditional cannulations, optofluidic device published in 2015) (19), this device offers reusability beyond the 4 independent reservoirs already available, making it ideal for long-term experiments. For example,

it could be used for experiments that have multiple testable time points (e.g., fear extinction, drug sensitization, neurodegenerative diseases). Another potential use for the multiple reservoirs is the injection of multiple agents at the same time or at discrete time points during a test session (e.g., agonistic stimulation of a system followed by selective antagonism later in the same session). Regardless of the particular experiment, the adaptable user interface for controlling drug and/or light delivery allows for tailored experimental designs without having to reprogram the device. The freedom to program and initiate drug or light delivery, both of which our device accomplishes, is essential for an effective and widely usable device.

This platform also provides some potential as part of an integrated closed-loop system for in vivo recording and manipulation. For example, sensors might monitor neural activity or organ behaviors to guide control of a second system to enhance or disrupt those patterns (e.g., via optogenetic illumination) (56– 59). Alternatively, the device and platform described here could be extended to include wireless photometric recordings of neural activity to guide localized pharmacological and/or optogenetic interventions (49). Similarly, genetically encoded receptor indicators could be combined with triggered drug release to quantify how the local neurochemical milieu affects receptor activity, with subsecond resolution (60). These and other possibilities provide multiple exciting avenues of study.

Overall, this lightweight optofluidic device with refillable/usable drug reservoirs, straightforward implantation procedures, an easy-to-use interface can accommodate wide-ranging experimental designs, and an essentially endless source of power via magnetic resonant coupling technology. The device is well suited for investigations of interactions between optogenetically activated circuits and subsequent neurochemical signaling, as well as for use with more advanced and cutting-edge technologies like optopharmacology. Because of its lightweight construction, it can be used in essentially any behavioral paradigm, as well as for either acute or long-term experiments. Its straightforward and streamlined manufacturing protocol also makes it readily accessible for nearly all engineering or neuroscience laboratories. Future iterations of the device could include multiple activatable LEDs, photometric sensing (49), or even more efficient NFC coupling to further boost its power output.

### **Materials and Methods**

Details of fabrications and assembly of wireless optofluidic device, power characterization, theoretical modeling of drug diffusion, Monte Carlo simulations on brain tissue, and flow rate measurement and modeling appear in *SI Appendix*. Experimental animals, surgery, immunohistochemistry, immunoglial response in implanted tissues, and behavioral experiments are also in *SI Appendix*. All procedures were approved by the Animal Care and Use Committee of Washington University and Northwestern University and conformed to US National Institutes of Health guidelines.

**Data and Materials Availability.** All data needed to evaluate the conclusions in the paper are present in the paper and/or *SI Appendix* and Movie S1.

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