Imaging Striatal Dopamine Release Using a Non-Genetically Encoded Near-Infrared Fluorescent Catecholamine Nanosensor

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Abstract

Neuromodulation plays a critical role in brain function in both health and disease. New optical tools are needed that can image neuromodulation with high spatial and temporal resolution, which will add an important new dimension of information to neuroscience research. Here, we demonstrate the use of a catecholamine nanosensor with fluorescent emission in the 1000-1300 nm near-infrared window to measure dopamine transmission in ex vivo brain slices. These near-infrared catecholamine nanosensors (nIR Cats) represent a broader class of nanosensors that can be synthesized from non-covalent conjugation of single wall carbon nanotubes (SWNT) with single strand oligonucleotides. We show that nIR Cats can be used to detect catecholamine efflux in brain tissue driven by both electrical stimulation or optogenetic stimulation. Spatial analysis of electrically evoked signals revealed dynamic regions of interest approximately 2 microns in size in which transients scaled with simulation intensity and lasted 5-10 seconds. Optogenetic stimulation of dopaminergic terminals produced similar transients, while optogenetic stimulation of glutamatergic terminals showed no effect on nIR Cat signal. Furthermore, bath application of nomifensine prolonged nIR Cat fluorescence signal, consistent with reuptake blockade of dopamine. These nanosensors may be advantageous for future use because they i) do not require virus delivery, gene delivery, or protein expression, ii) their near-infrared fluorescence facilitates imaging in optically scattering brain tissue and is compatible for use in conjunction with other optical neuroscience tool sets, and iii) the broad availability of unique near-infrared colors have the potential for simultaneous detection of multiple neurochemical signals. Together, these data suggest nIR Cats and other nanosensors of this class can serve as versatile new optical tools to report dynamics of extracellular neuromodulation in the brain.
The catecholamines dopamine and norepinephrine are neuromodulators known to play an important role in learning and attention and are implicated in multiple brain disorders.\textsuperscript{1–5} Dopamine, in particular, is thought to play a critical role in learning\textsuperscript{6}, motivation\textsuperscript{7,8}, and motor control\textsuperscript{9}, where aberrations in dopamine neurotransmission are implicated in a wide range of neurological and psychiatric disorders including Parkinson’s disease\textsuperscript{10}, schizophrenia\textsuperscript{11}, and addiction.\textsuperscript{12}

Neuromodulatory neurotransmission is thought to occur on a broader spatial scale than classic neurotransmission, whereby classic neurotransmission is mediated by synaptic release of glutamate and \(\gamma\)-aminobutyric acid (GABA) in the central nervous system. In synaptic glutamatergic and GABAergic neurotransmission, neurotransmitter concentrations briefly rise in the synaptic cleft to mediate local communication between the pre- and postsynaptic neurons through the rapid activation of ligand-gated ion channels.\textsuperscript{13} In contrast, neuromodulators (catecholamines, neuropeptides) may diffuse more broadly beyond the synaptic cleft and act via extrasynaptically-expressed metabotropic receptors.\textsuperscript{14–19} Thus, modulatory neurotransmitter activity extends beyond single synaptic partners and enables small numbers of neurons to modulate the activity of broader networks.\textsuperscript{20} The larger spatial scale and often absence of direct change in ionic flux across cell membranes makes quantification of neuromodulation a challenge.

Ideally, to understand how neuromodulation sculpts brain activity, we need to develop tools that can optically report modulatory neurotransmitter concentrations in the brain extracellular space (ECS) in a manner that is compatible with other available tools to image neural structure and activity. This goal motivates our effort to design an optical sensor that can report extracellular catecholamine dynamics with high spatial and temporal fidelity within a unique near-infrared spectral profile. In this work, we describe the design, characterization, and implementation of a nanoscale near-infrared (nIR) non-genetically encoded fluorescent reporter that allows precise measurement of catecholamine dynamics in brain tissue. This technology makes use of a single wall carbon nanotube (SWNT) non-covalently functionalized with single strand (GT)$_6$ oligonucleotides to form the near-infrared catecholamine nanosensor (nIRC). nIRCats respond to dopamine with \(\Delta F/F\) of up to 24-fold in the fluorescence emission window of 1000-1300 nm\textsuperscript{48}, a wavelength range that has shown utility for non-invasive through-skull imaging in mice.\textsuperscript{42} First, we show \textit{in vitro} data characterizing the specificity of the nanosensor for catecholamines dopamine and norepinephrine, and demonstrate the nanosensor’s insensitivity to GABA, glutamate, and acetylcholine. Second, we use \textit{ex vivo} brain slices containing the dorsal striatum to demonstrate that nIRCats exhibit a fractional change in fluorescence that has the dynamic range and signal-to-noise ratio to report dopamine efflux in response to brief electrical or optogenetic stimulation of dopaminergic terminals. Optogenetic stimulation of this preparation is also used to demonstrate the selectivity of the nIRCat nanosensor response to dopaminergic over glutamatergic terminal stimulation. Finally, the presence of a dopamine reuptake inhibitor yields a prolonged nIRCat fluorescent signal indicating that the sensors report a change in the time course of dopamine diffusion and reuptake in striatal
brain tissue. These data indicate that nIRCats provide a unique tool to interrogate the release, diffusion, and reuptake of neuromodulators in intact neural tissue.

**A Near-Infrared Dopamine and Norepinephrine Nanosensor**

We report near-infrared fluorescent catecholamine nanosensors (nIRCats) that enable imaging of extrasynaptic catecholamines and their release and re-uptake dynamics in the ECS of brain tissue. Using a previously established nanosensor generation platform\(^{21,22}\), synthetic bio-mimetic polymers were pinned onto the surface of intrinsically near-infrared fluorescent single-wall carbon nanotubes (SWNT). The resulting non-covalent nanometer-scale conjugate produced the catecholamine-selective nIRCat. In *in vitro* solution phase experiments (Methods), nIRCats exhibited a maximal change in fluorescence (\(\Delta F/F\)) of up to 24 (Figure 1b, 1c) with a dynamic range of 4 orders of magnitude, reporting detectable fluorescence changes from 10 nM to 100 \(\mu\)M dopamine concentration (Figure S1). nIRCats can also serve as norepinephrine sensors with a maximal response of \(\Delta F/F=35\) and a similar dynamic range (Figure S1). We showed that nIRCats are insensitive to GABA, glutamate, and acetylcholine (Figure 1c), and can measure fluctuations in dopamine concentration in the presence of ascorbic acid, which is present in cerebrospinal fluid (Figure S1). Single-molecule imaging revealed that the nanosensors were reversible upon exposure to repeated perfusions of 10 \(\mu\)M dopamine (Figure S2), an important feature for measuring neuromodulator kinetics. In previous work, we performed stochastic simulations that suggest nIRCats have sufficient sensitivity to detect physiologically relevant fluctuations in dopamine concentration in brain tissue arising from the activity of a single dopaminergic terminal, which can briefly exceed concentrations of 1 \(\mu\)M from the release site in a distance-dependent manner.\(^{23}\)
Figure 1. Synthesis and testing of near-infrared catecholamine nanosensors (nIRCats). (a) Schematic of optical catecholamine reporters, nIRCats. Pristine SWNT are functionalized with (GT)$_6$ oligonucleotides to generate turn-on optical reporters for dopamine (DA) and norepinephrine (NE) (b) Fluorescence spectra of nIRCats before (black) and after (red) addition of 10 µM of dopamine in an in vitro preparation in phosphate buffered saline (without tissue). Multiple emission peaks correspond to unique SWNT chiralities contained within the multi-chirality mixture. (c) Nanosensor optical response to 100 µM dopamine (DA), norepinephrine (NE), glutamate (GLU), γ-aminobutyric acid (GABA), and acetylcholine (ACH) (data from in vitro testing). Black bars represent averages from n=3 independent measurements and error bars are calculated as standard deviations of the n=3 measurements.

Imaging of electrical stimulation-evoked dopamine release in acute striatal brain slices

To determine the efficacy of nIRCats for imaging dopamine in brain tissue, we used brain slices from the dorsal striatum of the mouse. The dorsal striatum is densely innervated by dopaminergic projections from the substantia nigra pars compacta (SNC) but lacks significant innervation from cells that release norepinephrine (NE)$^{24}$, therefore we leveraged nIRCats capacity to serve as a dopamine sensor in this striatal brain region. The majority of neurons within the striatum are GABAergic medium spiny neurons (MSNs) with a minority fraction of interneuron populations that include GABAergic and cholinergic interneurons.$^{25}$ Glutamatergic inputs from the cortex and thalamus are the major drivers of MSN activity and dopaminergic terminals in close proximity to these inputs are thought to play an important role in modulating the activity of MSNs and plasticity at striatal synapses.$^{26}$ Due to the composition of local axons, intrastriatal
electrical stimulation is predicted to drive release of GABA, glutamate, acetylcholine, and dopamine, but negligible amounts of other catecholamines like norepinephrine.

Coronal mouse brain slices were prepared as described previously. Slices were subsequently incubated with 2 mg/L nIRCat for 15 minutes to enable sensors to diffuse into the brain tissue (Figure 2a). Slices were subsequently rinsed to remove excess nIRCat and incubated in standard artificial cerebrospinal fluid (ACSF) for another 15 minutes before imaging. This method, modified from Godin et al., who demonstrated that SWNT localize in the brain ECS of acute slices, enabled even and widespread labeling of the coronal slice, including the dorsal striatum (Figure 2c). Imaging of nIRCat fluorescence modulation in dorsal striatum was accomplished with a custom-built visible and near-infrared microscope to enable concurrent imaging of both visible (400 nm – 750 nm) and near-infrared (750 nm – 1700 nm) wavelengths on the same detector (Figure 2b). Briefly, a 785 nm laser for excitation of nIRCat and mercury bulb for generating brightfield images were directed onto the back focal plane of an epifluorescence upright microscope, and imaging channels were selected using a sliding mirror. Serially, either brightfield or near-infrared images were collected on a Ninox Vis-SWIR 640 broadband camera (Raptor Photonics) with appropriate dichroic filters (Methods) and a 60X water dipping objective (Nikon) providing an imaging field of 178 µm by 142 µm, containing hundreds of dopaminergic terminals.
Figure 2. Brain slice nIRCat loading protocol and schematic of visible and near-infrared fluorescence microscopy for imaging nIRCats in acute slices. (a) Experimental schematic depicting preparation of acute brain slices and subsequent incubation in 2 mg/L nIRCat solution to load the nanosensors into brain tissue. (b) Schematic of visible/near-infrared microscope. A 785 nm CW laser is beam-expanded and co-aligned with a mercury vapor lamp and directed into the objective with dichroic filter cubes. Emitted photons are filtered through a 900 nm long-pass filter and are relayed onto the sensor of a broadband InGaAs camera that is sensitive to both visible and near-infrared wavelengths. (c) Dorsomedial striatum from mouse acute slice imaged in brightfield (top) and near-infrared (bottom) after tissue nanosensor loading. Scale bars = 10 µm.

To investigate striatal neuromodulator release with temporal control of tissue stimulation, we used a bipolar stimulating electrode to evoke terminal release within the dorsomedial striatum (stimulus protocol: 3 millisecond wide single square pulses, over 5 biological replicates). We found a single pulse could elicit a nIRCat signal transient, and found that increasing the strength of the stimulus led to larger evoked changes in nIRCat \( \Delta F/F \) signal, \((\Delta F/F)_{\text{max}} \) 0.1 mA = 0.0465 ± 0.025; 0.3 mA = 0.122 ± 0.026; and 0.5 mA = 0.2 ± 0.033; mean ± s. d., n=5 for all measurements ) (Figure 3b).

To further test if evoked nIRCat signals tracked striatal dopamine release and reuptake kinetics, we investigated the effect of nomifensine, a dopamine reuptake inhibitor that slows the clearance of dopamine from the ECS by competitively binding to dopamine transporters (DATs). Addition of 10 µM nomifensine to the bath yielded nIRCat signal
with higher peak fluorescence modulation \((\Delta F/F)_{\text{max}} = 0.108 \pm 0.029 \text{ vs. } 0.189 \pm 0.023; \text{ mean } \pm \text{ s. d., } n=3\) and a prolonged fluorescent signal compared to signals obtained in normal ACSF from the same field of view (decay time constant, \(\tau = 2.43 \pm 0.24 \text{ s vs. } 10.95 \pm 1.15 \text{ s; mean } \pm \text{ s. d., } n=3\) (Figure 3a top vs. bottom row, Figure 3c).

Figure 3. Imaging and spatiotemporal analysis of dopamine release evoked by electrical stimulation in acute striatal slices. (a) Nanosensor \(\Delta F/F\) imaging of the same acute brain slice field of view upon electrical stimulation by 0.3 mA in standard ACSF (top row) and in ACSF plus 10 \(\mu\)M nomifensine (bottom, +Nomifensine). Three still frames are presented from each movie. “Pre” is a representative frame before electrical stimulation is applied, “Stim” represents frame corresponding to peak \(\Delta F/F\) following stimulation, and “Post” is a representative frame after nIRC at fluorescence has returned to baseline. Scale bars = 10 \(\mu\)m. (b) Nanosensor fluorescence modulation scales with single pulse electrical stimulation amplitudes. Field of view mean traces and standard deviation bands are presented for three stimulation amplitudes of 0.1 mA, 0.3 mA, and 0.5 mA (n=5) (c) Time traces of \(\Delta F/F\) for 0.3 mA single pulse stimulation in standard ACSF (red) and in ACSF plus 10 \(\mu\)M nomifensine (purple, +NOMF). Mean traces with standard deviation bands are presented (n=3). (d) A single frame from a time series of imaging in the dorsomedial striatum showing the entire field of view, overlaid with ROIs identified from an automated program from per-pixel \(\Delta F/F\) stack
projections of nIRCat fluorescence modulation (see Methods). Color bar represents nIRCat fluorescence intensity of labeled tissue. Scale bar = 20 µm. (e) Frequency histogram of ROI sizes depicted in (d), exhibiting a log-normal distribution with median ROI size of 2 µm. (f) ROI size distribution from three different fields of view (representing n = 3 biological replicates) in the dorsal striatum, each stimulated separately by 0.3 mA stimulation. In each of these fields of view, ROIs showed similar median size and size distribution even when compared across biological replicates. (g) A higher magnification view of an ROI with an effective radius of 5 µm. Maximum ΔF/F projection of the ROI shows presence of smaller fluorescence hotspots within the ROI. Scale bar = 5 µm. (h) Overlay of representative normalized FSCV (gray) and nIRCat (blue) traces showing that nIRCat ROI signals exhibit a wider diversity of decay kinetics. Inset: An example of nIRCat experimental data (blue dots) fitted to first order kinetics (red line) to compute decay time constants (τ). (i) Normalized frequency histogram of τ’s computed from FSCV and nIRCat individual ROI time traces. Data from n = 4 fields of view representing n=2 biological replicates is pooled. Medians of each distribution are: nIRCats τ = 1.1 s and FSCV τ = 0.4 s.

To identify nIRCat fluorescence change hotspots (i.e., regions of high ΔF/F), we analyzed our video-rate acquisitions using a custom-built program that accounted for background fluorescence and identified regions with fluctuations in fluorescence intensity in the post-stimulation epoch (see Methods for details). We defined nIRCat ΔF/F hotspots as regions of interest (ROIs) based on a per-pixel stack projection of maximal ΔF/F in the imaging time series. The ROIs identified were found to vary in size from 1 µm to 15 µm (Figure 3d, 3e). We found that ΔF/F hotspots do not necessarily correspond to high nIRCat labeling of the brain tissue, suggesting that the hotspot is a consequence of variation in dopamine release not nanosensor loading in the tissue (Figure 3d, Figure S3). Using data from single pulse electrical stimulation experiments, the program identified ROIs whose size distribution was observed to exhibit a log-normal distribution with median ROI size of 2 µm (Figure 3e). Repeat stimulations with the same stimulation amplitude in fields of view of the dorsomedial striatum across biological replicates generated similar size distributions (Figure 3f). Closer examination of several larger ROIs (> 5µm) suggested these may be comprised of smaller hot-spots in close proximity (Figure 3g).

For further examination of the temporal resolution of nIRCats, we compared the temporal profile of evoked transients measured with nIRCats to transients measured with fast scan cyclic voltammetry (FSCV). FSCV is a technique that has been widely utilized to measure catecholamine dynamics both in vivo and in vitro in the striatum and other brain areas.29–31 FSCV and nIRCat experiments were conducted on separate experimental rigs with the same solutions, temperature settings, electrodes, and stimulation parameters. Evoked transients measured with FSCV and nIRCat fluorescence emission showed comparable temporal profiles in the rising phase (latency to peak: FSCV = 0.25 ± 0.0 s vs. nIRCat = 0.40 ± 0.18 s; mean ± s. d., n=4 fields of view from 2 biological replicates). Meanwhile, nIRCat signals exhibited a wider diversity of decay kinetics (τ: FSCV = 0.51 ± 0.08 s vs. nIRCats = 2.43 ± 0.24 s; mean ± s. d. n=4 fields of view from 2 biological replicates). A subset of ROIs exhibited decay time constants that overlapped with those of FSCV signals (Figure 3h, 3i).

**Imaging of optogenetically-evoked dopamine release in acute striatal brain slices**

To further confirm striatal nIRCat nanosensor signals were indeed reporting dopamine release, we compared channelrhodopsin (ChR2) stimulation of cortical glutamatergic and
nigrostriatal dopaminergic terminals in the dorsal striatum. Acute striatal brain slices were prepared from mice virally transfect one the light sensitive cation channel ChR2 in either dopaminergic terminals (targeted by viral injection in the midbrain in DAT-cre mice; ChR2-DA) (Figure 5c) or glutamatergic terminals of the striatum (targeted by viral injection in the frontal cortices, ChR2-GLU) (Figure 5a). Upon optical stimulation of ChR2-DA terminals with a 473 nm laser (5 pulses at 25 Hz, 1 mW/mm²) in the dorsal striatum, we observed significant \((\Delta F/F)_{\text{max}} = 0.1\) fluorescence modulation of nIRCat signal (Figure 5d). When optogenetic stimulation instead targeted glutamatergic terminals, we could observe excitatory postsynaptic currents (EPSCs) in an MSN neuron, but fluorescent nIRCat signals did not rise above baseline fluctuation (Figure 5b). We also examined the effects of nomifensine on nIRCat signals elicited by optogenetic stimulation of dopaminergic terminals. Optogenetic stimulation of dopaminergic terminals in the striatum of DAT-cre mice in the presence of 10 µM nomifensine led to significantly longer decay times in evoked nIRCat signal, consistent with the predicted slowing of dopamine clearance from the ECS (Figure S4).

![Diagram](image_url)

**Figure 5. Optogenetic release of dopamine imaged by nIRCats.** (a) Schematic of ChR2 expression in cortical glutamatergic terminals of the dorsal striatum. (b) No nIRCat fluorescence modulation was observed after stimulation of glutamate terminals. Inset: glutamate release was confirmed by excitatory postsynaptic current on medium spiny neuron. (c) Schematic of ChR2 expression in nigrostriatal dopaminergic terminals of the dorsal striatum. (d) Stimulation of dopaminergic terminals resulted in nIRCat fluorescence modulation. Stimulation protocol in (b) and (d) was 5 pulses at 25 Hz and power flux of 1 mW/mm².
Conclusion

To capture spatial information of neuromodulator kinetics and to integrate our understanding of neuromodulation with other measures of brain activity, there is need for new optical tools to measure the extracellular dynamics of neuromodulator release and reuptake. Here, we demonstrated the feasibility of using of a non-genetically encoded fluorescent sensor, nIRCat, which enables optical detection of catecholamine release and reuptake with sub-second temporal and with micrometer spatial resolution. We used electrical and optogenetic methods in ex vivo brain slices to demonstrate that nIRCat fluorescent signals can successfully report differences in evoked dopamine release and pharmacologically induced changes in dopamine reuptake kinetics in striatal tissue.

Here, we focused nIRCat imaging experiments within the dorsal striatum, a region that receives dense dopaminergic innervation and negligible norepinephrine innervation. Therefore, while nIRCats are not selective for dopamine over norepinephrine, they effectively function as a dopamine sensor within the context of the striatum. Given that striatal dopamine regulates fundamental processes including motor function, motivation and learning, nIRCats represent an important addition to the neuroscience investigative toolkit.

While other tools are emerging to optically report dopamine fluctuations via cell-surface engineered proteins, nIRCats are likely to fulfill a niche amongst currently available methods for detecting dopamine neurotransmission due to their unique near-infrared fluorescence, and because they do not rely on genetic delivery and expression. In addition, the functionality of nIRCats is not hindered by the presence of dopamine receptor ligands, in contrast to receptor-based fluorescent sensors which cannot report endogenous dopamine in the presence of ligands to the engineered receptor. nIRCats also offer spatial advantages over FSCV, and our initial experiments suggest that the temporal resolution of the nIRCat signal is comparable to that of FSCV. nIRCat decay profiles exhibit a wider range than that observed from FSCV data, and included a significant number of ROIs that showed seconds-long time constants. These results may capture the unique spatial property of nIRCats: unlike FSCV probes which sample catecholamine concentration at a single point in space, each distributed nIRCat molecule can act as a probe within the ECS and therefore yield a “higher resolution” picture of catecholamine dynamics. Future experiments will investigate how the heterogeneity of nIRCat signals (amplitude and kinetics) relates to structural and functional properties at dopamine terminals and within the ECS. We predict that new optical tools for measuring dopamine dynamics with high spatial resolution will be enable new sights into the regulation of dopamine release and reuptake.

In the future, we see potential for expansion of a larger family of SWNT based near-infrared nanosensors (nIRNS) similar to nIRCats for multiple neurochemical imaging applications. Several lines of evidence illustrate their future potential. First, nIRNS are easily functionalized with a wide range of synthetic molecular recognition moieties, affording fine control of their surface functional elements and their interactions with the local chemical environment. SWNT fluorescence can be finely tuned to
monochromatic emission in the near-infrared II (1000-1700 nm) window by controlling the SWNT chirality.\textsuperscript{40} This chirality-dependent fluorescence in the near-infrared II window provides further avenues for designing color-specific responses to multiple molecular analytes simultaneously, thereby affording synthesis of ratiometric and multiplexed analyte imaging platforms, as we have shown previously.\textsuperscript{41} Second, SWNT-based nanosensors rely on near-infrared fluorescence, which greatly reduces the impact of tissue scattering in the emission window and therefore may enable through-cranium imaging.\textsuperscript{42} nIRNS are compatible with multi-photon imaging with 1600 nm excitation\textsuperscript{43} and as such could permit nanoscale imaging of intact neuronal structures as has been shown with visible wavelength-emitting fluorophores.\textsuperscript{44} Third, nIRNS have robust photostability allowing for use in long-term imaging experiments.\textsuperscript{45} Fourth, because nIRNS are not genetically-encoded, they could enable use in species where gene delivery and protein expression is too difficult, time consuming, or undesirable. Finally, nanosecond-scale binding kinetics and the nanoscale dimensions of nIRNS\textsuperscript{48}, are likely to enable generation of nanosensors with improved temporal and spatial resolution. In sum, nIRCats are versatile catecholamine probes amenable to multiplexing with existing tools for concurrent investigation of dopaminergic neuromodulation with other core mechanisms of brain function.

**Methods**

*Nanosensor synthesis*

\((\text{GT})_6\) oligonucleotides were purchased from Integrated DNA Technologies (IDT, Standard Desalting). HiPCo SWNT were purchased from NanoIntegris (Batch \# HR27-104). \((\text{GT})_6\)-SWNT colloidal suspension (nIRCat) was prepared by mixing 1 mg of \((\text{GT})_6\) and 1 mg of SWNT in 1 mL of a 100 mM NaCl solution. The solution was bath sonicated (Branson Ultrasonic 1800) for 10 minutes and probe-tip sonicated for 10 minutes at 5 W power (Cole Parmer Ultrasonic Processor, 3 mm tip diameter) in an ice-bath. The sonicated solution was incubated at room temperature for 30 minutes. The product was subsequently centrifuged at 16,000 g (Eppendorf 5418) for 90 minutes to remove unsuspended SWNT bundles and amorphous carbon, and the supernatant was recovered for characterization and use. Each nanosensor suspension was stored at 4°C until use.

*Nanosensor characterization*

To characterize nIRCats post-synthesis, the full visible and near-infrared absorption spectrum was taken for each nanosensor batch (UV-VIS-NIR spectrophotometer, Shimadzu UV-3600 Plus) or UV-VIS (ThermoFisher Scientific Genesys 20). SWNT concentrations of as-made nanosensor batches were determined using absorbance at 632 nm (UV-VIS) with an extinction coefficient of \(\varepsilon = 0.036 \text{ (mg/L)}^{-1} \text{ cm}^{-1}\). Full spectrum absorbance measurements were made with UV-VIS-NIR after dilution to 5 mg/L SWNT concentration in 100 mM NaCl. For fluorescence spectroscopy, each sensor batch was diluted to a working concentration of 5 mg/L in 100 mM NaCl, and aliquots of 198 \(\mu\)L were placed in each well of a 96-well plate (Corning). Fluorescence measurements were obtained with a 20 X objective on an inverted Zeiss microscope (Axio Observer.D1) coupled to a Princeton Instruments spectrograph (SCT 320) and liquid nitrogen cooled...
Princeton Instruments InGaAs linear array detector (PyLoN-IR). A 721nm laser (OptoEngine LLC) was used as the excitation light source for all characterization experiments.

**Neurotransmitter screening**

For neurotransmitter response screens, we collected the near-infrared fluorescence spectrum from 198 μL aliquots of nanosensor (5 mg/L SWNT concentration) before and after addition of 2 μL of 10 mM solutions of each analyte neurotransmitter (for a 100 μM final analyte concentration in each well of a 96-well plate). All neurotransmitter analytes were purchased from Sigma-Aldrich. Neurotransmitter analytes were incubated for 5 minutes before taking post-analyte fluorescence measurements. Responses were calculated for the integrated fluorescence count as $\Delta F/F_0 = (F - F_0)/F_0$, where $F_0$ is total fluorescence before analyte addition and $F$ is total fluorescence after analyte addition or for peak fluorescence change corresponding to the (9,4) SWNT chirality (~1128 nm center wavelength). All measurements were made in triplicate. Reported results are mean and standard deviations of the triplicate measurements. All nIRCat nanosensor batches were tested for catecholamine responses prior to use for tissue catecholamine imaging.

**Nanosensor reversibility testing**

A #1.5 glass coverslip was functionalized with (3-Aminopropyl) triethoxysilane (APTES, Sigma Aldrich) by soaking in 10% APTES in ethanol for 5 min. The coverslip was then rinsed with DI water and left to dry. The coverslip was then fixed onto an ibidi sticky-Slide VI 0.4 forming 6 microfluidic channels. First, 100 μL of PBS was pipetted through a channel. Next, the channel was filled 50 μL of a 5 mg/L solution of nIRCats and left to incubate at room temperature for 5 min. The channel was rinsed using three 50 μL PBS washes, keeping the channel filled with solution at all times. The surface immobilized nIRCats in PBS were imaged on an epifluorescence microscope with 721 nm excitation and a Ninox VIS-SWIR 640 camera (Raptor). One end of the flow channel was connected to a syringe pump (Harvard Apparatus) using Luer lock fittings. Prior to the start of image acquisition, the opposite flow reservoir was filled with PBS and the pump was set to refill mode at a volumetric flow rate of 40 μL min$^{-1}$. Once the liquid in the reservoir was depleted, 40 μL of 10 μM dopamine in PBS was added. The process was repeated using alternating additions of 80 μL of PBS washes and 40 μL of dopamine solution.

**Acute slice preparation and nanosensor labeling**

Mice were C57 Bl/6 strain, 60 days old, and both male and female mice were used. Mice were group housed after weaning at P21 and kept with nesting material on a 12:12 light cycle. All animal procedures were approved by the UC Berkeley Animal Care and Use Committee. Acute brain slices were prepared using established protocols.$^{27}$ Briefly, mice were deeply anesthetized via intraperitoneal injection of ketamine/xyazine cocktail and transcardial perfusion was performed using ice-cold cutting buffer (119 mM NaCl, 26.2 mM NaHCO$_3$, 2.5 mM KCl, 1mM NaH$_2$PO$_4$, 3.5 mM MgCl$_2$, 10 mM Glucose, 0 mM CaCl$_2$), after which the brain was rapidly extracted. The cerebellum and other connective
tissues were trimmed using a razor blade and the brain was mounted onto the cutting stage of a vibratome (Leica VT 1200S). Coronal slices (300 µm in thickness) including the dorsal striatum were prepared. Slices were incubated at 37°C for 60 minutes in oxygen saturated ACSF (119 mM NaCl, 26.2 mM NaHCO₃, 2.5 mM KCl, 1mM NaH₂PO₄, 1.3 mM MgCl₂, 10 mM Glucose, 2 mM CaCl₂) before use. Slices were then transferred to room temperature for 30 minutes before starting imaging experiments and were maintained at room temperature for the remainder of experimentation.

For nanosensor labeling, slices were transferred into a small volume brain slice incubation chamber (Scientific Systems Design, Inc., AutoMate Scientific) and kept under oxygen saturated ACSF (total 5 mL volume). 100 µL of 100 mg/L nIRCat nanosensor was added to the 5mL volume and the slice was incubated in this solution for 15 minutes. The slice was subsequently recovered and rinsed in oxygen saturated ACSF to wash off nIRCats that did not localize into the brain tissue. The rinsing step was performed by transferring the slice through 3 wells of a 24 well plate (5 seconds in each well) followed by transfer to the recording chamber with ACSF perfusion for a 15-minute equilibration period before starting the imaging experimentation. All imaging experiments were performed at 32°C.

**Acute slice preparation for FSCV recording**

Acute slices were prepared as described previously. Extracellular dopamine concentration evoked by local electrical stimulation was monitored with FSCV at carbon-fiber microelectrodes (CFMs) using Millar voltammeter. CFMs were ~ 7 µm in diameter encased in glass capillary pulled to form a seal with the fiber and cut to final tip length of 70-120 µm. The CMF was positioned ~100 µm below the tissue surface at a 45-degree angle. A triangular waveform was applied to the CMF scanning from -0.7 V to +1.3 V and back, against Ag/AgCl reference electrode at a rate of 800 V/s. Evoked dopamine transients were sampled at 8 Hz, and data were acquired at 50 kHz using AxoScope 10.5 (Molecular Devices). Oxidation currents evoked by electrical stimulation were converted to dopamine concentration from post-experimental calibrations. Recorded FSCV signals were identified as dopamine by comparing oxidation (+0.6 V) and reduction (-0.2 V) potential peaks from experimental voltammograms with currents recorded during calibration with 2 µM dopamine dissolved in ACSF. For stimulation, a bipolar stimulation electrode (FHC CBAEC75) was positioned on top of the brain slice and approximately 100 µm away from the CFM. Following 30-minute slice equilibration in the recording chamber, dopamine release was evoked using a square pulse (0.3 mA pulse amplitude, 3 ms pulse duration) controlled by Isoflex stimulus isolator (A.M.P.I) and delivered out of phase with the voltammetric scans. Stimulation was repeated 3 times. To compare FSCV and nIRcat data, each signal was normalized against its peak value ([DA]max or (∆F/F)max) and co-aligned at stimulation time. Latency to peak were computed as tpeak - tstim where tpeak is the time at which peak signal is attained and tstim is time of stimulation. Decay time constants (τ) were computed from model fits to a first order decay process.
Microscope construction and slice imaging

*Ex vivo* slice imaging was performed with a modified upright epifluorescent microscope (Olympus, Sutter Instruments) mounted onto a motorized stage. Nanosensor excitation was supplied by a 785 nm, CW, DPSS laser with adjustable output power to a maximum of 300 mW and a near TEM00, top hat beam profile (OptoEngine LLC). The beam was expanded using a Keplerian beam expander comprised of two plano-convex lenses (Thorlabs, f=25 mm, f=75 mm, AR coating B) to a final beam diameter of approximately 1 cm. The beam was passed through a custom fluorescence filter cube (excitation: 800 nm shortpass (FESH0800), dichroic: 900 longpass (DMLP990R), emission: 900 longpass (FELH0900), Thorlabs) to a 60X Apo objective (1.0 NA, 2.8 mm WD, water dipping, high NIR transmission, Nikon CFI Apo 60XW NIR). Emission photons collected from the sample were passed through the filter cube and were focused onto a two-dimensional InGaAs array detector (500-600 nm: 40% quantum efficiency (QE); 1000-1500 nm: >85% QE; Ninox 640, Raptor Photonics) and recorded using Micro-Manager Open Source Microscopy Software. Laser power was adjusted to maximize collected photons and fill the pixel bit depth on the detector but did not exceed 70 mW at the objective back focal plane. YFP fluorescence was imaged by switching the filter cube (FESH 0899 for excitation, FELH 0900 for emission, Thorlabs) and using a mercury-vapor lamp (Olympus) for excitation.

Electrical and optical stimulation-evoked dopamine imaging with near-infrared microscopy

For electrical stimulation experiments, a bipolar stimulation electrode was positioned in field of view within the dorsomedial striatum identified using a 4X objective (Olympus xFluor 4x/340). Using 60X objective, the stimulation electrode was brought into contact with top surface of the brain slice and an imaging field of view was chosen at a nominal distance of 150 µm from the stimulation electrode within the dorsomedial striatum. All stimulation experiments were recorded at video frame rates of 9 frames per second (nominal) and single pulse electrical stimulations were applied after 200 frames of baseline were acquired. Each video acquisition lasted 600 frames. Stimulation amplitudes were staggered and each stimulation amplitude was repeated three times within a field of view. Slices were allowed to recover for 5 minutes between each stimulation with the excitation laser path shuttered. For optogenetic stimulation, a fiber-coupled 473 nm blue laser (OptoEngine LLC DPSS) was positioned in close proximity to the brain slice using a micromanipulator. Expression of ChR2 was confirmed via visible fluorescence imaging and an imaging field of view was chosen in dorsomedial striatum with robust expression level. Stimulation pulses (5 pulses, 5 ms duration per pulse, delivered at 25 Hz, 1 mW/mm²) were delivered after acquiring 200 baseline frames and the video acquisition lasted 600 frames at nominal 9 frames per second.

Viral transfection of mice for optogenetic stimulation

Adult male and female mice (>P60) were used for all surgeries. Bilateral viral injections were performed using previously described procedures at the following stereotaxic coordinates: dorsomedial prefrontal cortex (dmPFC): 1.94 mm from Bregma, 0.34 mm
lateral from midline, and 0.70 mm vertical from cortical surface; substantia nigra pars compacta (SNc): -3.08 mm from Bregma, 1.25 mm lateral from midline, and 4.0 mm vertical from cortical surface. For glutamatergic corticostralial axon stimulation experiments, mice were injected with 0.5 µL of CAG-ChR2-EYFP virus bilaterally into dmPFC. For nigrostriatal dopaminergic axon stimulation experiments, DAT-Cre mice were injected with 0.5 µL DIO-ChR2-EYFP virus bilaterally. For all optogenetic experiments, we waited at least three weeks from viral injection to experimental stimulation to allow for sufficient ChR2 gene expression.

Image processing and data analysis of nIRCat fluorescence response

Raw movie files were processed using a custom-built MATLAB program (available for download: https://github.com/jtdbod/Nanosensor-Brain-Imaging). Briefly, for each raw movie stack (600 frames), a per pixel \((F-F_0)/F_0\) was calculated using the average intensity for the first 5% of frames as \(F_0\). Regions of interest (ROIs) were identified by calculating a median filter convolution and then performing thresholding using Otsu’s method followed by a morphological dilation operation. \(\Delta F/F\) traces were then calculated for each generated ROI. ROI sizes were computed using the measured pixel area and approximating each as a circle to calculate an equivalent radius.

To compare responses across stimulation amplitudes and bath application of nomifensine, mean results were obtained as follows: First, all identified ROIs from a field of imaging were averaged. Mean traces were further averaged over different fields of view within the same slice and across slices (1-2 field of view per slice, 1-2 slices per animal) and then averaged over experimental animals. Decay time constants \(\tau\) were computed by fitting \(\Delta F/F\) time traces to a first order decay process. Latency to peak were computed as \(t_{peak} - t_{stim}\) where \(t_{peak}\) is the time at which peak signal is attained and \(t_{stim}\) is time of stimulation.

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Competing Interests

The authors declare no competing financial interests.

Supplementary Information
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