Combined Optogenetic Approaches Reveal Quantitative Dynamics of Endogenous Noradrenergic Transmission in the Brain

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HIGHLIGHTS
We report all-optogenetic stimulation and imaging of NA transmission in cortex
NA fiber stimulation elicits robust PKA activation in neurons, in slice and in vivo
PKA activation involves both β1 and α2 adrenoceptors but not dopamine receptors
Noradrenaline reuptake transporter activity greatly limits response intensity

Nomura et al., iScience 23, 101710
November 20, 2020 © 2020
The Author(s).
https://doi.org/10.1016/j.isci.2020.101710

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Combined Optogenetic Approaches Reveal Quantitative Dynamics of Endogenous Noradrenergic Transmission in the Brain

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SUMMARY
Little is known about the real-time cellular dynamics triggered by endogenous catecholamine release despite their importance in brain functions. To address this issue, we expressed channelrhodopsin in locus coeruleus neurons and protein kinase-A activity biosensors in cortical pyramidal neurons and combined two-photon imaging of biosensors with photostimulation of locus coeruleus cortical axons, in acute slices and in vivo. Burst photostimulation of axons for 5–10 s elicited robust, minutes-lasting kinase-A activation in individual neurons, indicating that a single burst firing episode of synchronized locus coeruleus neurons has rapid and lasting effects on cortical network. Responses were mediated by β1 adrenoceptors, dampened by co-activation of α2 adrenoceptors, and dramatically increased upon inhibition of noradrenaline reuptake transporter. Dopamine receptors were not involved, showing that kinase-A activation was due to noradrenaline release. Our study shows that noradrenergic transmission can be characterized with high spatiotemporal resolution in brain slices and in vivo using optogenetic tools.

INTRODUCTION
The catecholamines dopamine (DA) and noradrenaline (NA) are neurotransmitters that widely modulate brain circuits and behaviors. Their dysfunctions are associated with cognitive, emotional, and motor disorders, and many drugs target catecholaminergic transmission in therapy of human brain diseases. Little is known about the cellular dynamics of endogenous catecholaminergic transmission, because of the difficulty of selectively stimulating catecholaminergic fibers and because catecholamines modulate neuronal excitability but do not elicit fast and prominent membrane currents, thereby limiting the sensitivity of electrophysiological readout. However, recent reports demonstrate that optogenetic tools allow the detection of catecholamine release and of their effects at the cellular level with physiologically relevant temporal resolution in brain slices and in vivo (Goto et al., 2015; Ma et al., 2018; Sun et al., 2018; Feng et al., 2019; Oe et al., 2020).

NA is involved in arousal, attention, memory, and stress (Sara 2009; Benarroch 2009). It is synthesized in discrete brainstem nuclei from the amino acid tyrosine, transformed by tyrosine hydroxylase (TH) into DA, which is in turn converted into NA by dopamine beta hydroxylase (DBH). NA effects are mediated by G protein-coupled receptors, among which Gs-coupled β1-3 and Gi-coupled α2 receptors activate and inhibit adenylate cyclase, respectively, whereas Gq-coupled α1 receptors activate phospholipase C (Bjelke 1992; Cotecchia 2010; Evans et al., 2010). NA-releasing fibers are widely distributed in the brain and, in the cerebral cortex, stem from neurons of the locus coeruleus (LC), whose axons innervate all layers of the entire cortical mantle (Morrison et al. 1978, 1979; Latsari et al., 2002; Nomura et al., 2014). In pyramidal cells, the major cortical neuron type, exogenous NA elicits only modest variations of membrane potential but triggers intense cAMP-protein kinase A (PKA) signals through activation of β1 adrenoceptors (McCormick et al., 1991; Castro et al., 2010; Nomura et al., 2014). Likewise, imaging of genetically encoded optical sensors for cAMP-PKA activity provides a sensitive readout of NA effects and allows detecting endogenous NA transmission events (Nomura et al., 2014; Ma et al., 2018; Oe et al., 2020).

The use of channelrhodopsin2 (ChR2) photostimulation to trigger neurotransmitter release (Nagel et al., 2003; Zhang et al., 2007) is compatible with two-photon imaging of NA effects, owing to the virtual absence
of ChR2 activation upon two-photon laser scanning (Rickgauer and Tank 2009; Schoenenberger et al., 2011). We thus combined two-photon imaging of pyramidal cells expressing fluorescent PKA activity sensors with one-photon stimulation of ChR2-expressing LC fibers to characterize catecholaminergic transmission dynamics in the neocortex. We found that burst photostimulation of LC fibers resulted in transient and repeatable activation of PKA in pyramidal cells and used selective antagonists to characterize catecholamine receptors and reuptake transporters involved in this response, in brain slice and in vivo.

RESULTS
Expression of Channelrhodopsin in LC Fibers and of AKAR3EV in Cortical Neurons
In order to stimulate specifically LC fibers, we selectively expressed a fusion protein consisting of ChR2 and YFP in LC neurons using conditional viral transduction in DBH-Cre mice (see Methods and Nomura et al., 2014). Eight weeks after virus injection, immunohistochemistry revealed that ChR2-YFP was selectively expressed in a large proportion of DBH-positive neurons in LCs of both hemispheres (Figure 1A) but not in other catecholaminergic nuclei (not shown, but see Nomura et al., 2014). We found that 87.0 ± 2.7% of DBH-positive neurons in the LC expressed GFP (n = 214, N = 3 mice), consistent with earlier observations (Nomura et al., 2014). Conversely, only 4.9 ± 0.9% of GFP-positive cells were DBH-negative (n = 197, N = 3). ChR2-YFP was efficiently targeted to the plasma membrane of TH-positive somata (Figure 1A). In the parietal cortex, ChR2-YFP-positive LC fibers coursed throughout all layers (Figure 1B), consistent with their wide distribution in the cortical mantle (Nomura et al., 2014). These results show that the present protocol enables selective and efficient expression of ChR2-YFP in the soma and axonal processes of LC neurons.

To monitor cortical responses to photostimulation of ChR2-expressing LC fibers, we also injected recombinant viruses encoding PKA activity sensors in the parietal cortex after LC transduction (see Methods). For imaging in cortical slices, we used a Sindbis virus to express the AKAR3EV sensor (Komatsu et al., 2011). AKAR3EV is based on FRET between two GFP mutants, whose brightness facilitates the imaging in diffusive slices from myelinated adult tissue. Ten hours after virus injection, AKAR3EV fluorescence was observed throughout cortical layers II to VI, with AKAR3EV-expressing neurons observed as far as 300 μm away from the injection site (Figure 1C). For imaging cortical neurons in vivo, we used an AAV to express the single-chromophore, GFP-based GAKdYmut sensor (Bonnot et al., 2014), because its large dynamic range enables the detection of small changes in PKA activity and single-chromophore sensors allow two-photon imaging without the depth-dependent re-absorption problem of FRET sensors. Five weeks after AAV injection, GAKdYmut was widely expressed in layer V pyramidal neurons typically exhibiting a prominent apical dendrite (Figure 1D), allowing two-photon imaging of GAKdYmut-positive neurons in the depth of the cortex in vivo (Figure 1E). Co-injection of recombinant viruses encoding the red fluorophores mCherry or TurboRFP (see Methods and Figures 1C–1E) allowed visualization of viral transduction areas in cortical slices or in vivo, respectively, with minimal excitation of ChR2. These results show that the present approaches result in efficient cortical expression of both ChR2 in LC fibers and PKA sensors in neurons.

Photostimulation of LC Fibers Elicits PKA Activation in Pyramidal Neurons
The Sindbis virus preferentially transduces neocortical pyramidal neurons and allows overnight expression of biosensors without conspicuous alteration of neuronal physiology (Lendvai et al., 2000; Gervasi et al., 2007; Drobac et al., 2010; Hu et al., 2011). We imaged PKA dynamics in AKAR3EV-expressing layers II/III and V pyramidal cells from cortical slices, which both respond to exogenous catecholamines (Nomura et al., 2014), and focused primarily on layer V pyramidal neurons. In a series of pilot experiments, photostimulation patterns were tested for their ability to evoke PKA signals, based on the in vivo burst firing of LC neurons that discharge action potentials at a maximal frequency of 10 Hz (Aston-Jones and Bloom, 1981a; 1981b), which is compatible with ChR2 kinetic properties (Mattis et al., 2011). We thus applied trains of light pulses of variable length, at a maximal frequency of 10 Hz, for different overall durations. Most protocols elicited transient PKA activation (see examples in Figure S1). However, photostimulation consisting of 5-ms light pulses applied at 10 Hz during 5 s yielded robust and reproducible responses (Figure 2A) and was used in all subsequent experiments in brain slices. Unless otherwise stated, responses were measured at the soma of pyramidal neurons. At the end of photostimulation series, saturation of the AKAR3EV sensor was obtained by bath application of a maximally effective concentration of the adenylyl cyclase activator forskolin (FSK, 12.5 μM; Gervasi et al., 2007) (Figure 2A). Photostimulation induced clearly detectable peaks of PKA activity in 93 ± 7% of individual layer II/III (n = 34 of 37, N = 4 slices from four mice) and in 86 ± 14% of individual layer V (n = 18 out of 21, N = 3 slices from three mice) pyramidal neurons. Responses were
Figure 1. Expression of ChR2 in LC Fibers and of PKA Sensors in Cortical Neurons

(A) Upper panels: Immunolabelling and DAPI counterstaining of a hindbrain section shows Cre-dependent expression of ChR2-YFP in TH-positive neurons 8 weeks after bilateral AAV injection into the LC in a DBH-Cre mouse. V: lumen of the fourth cerebral ventricle. Lower panels: Higher magnification shows that ChR2-YFP is selectively expressed in a large...
Figure 1. Continued

...ranient and were consistently observed at each stimulus upon repetitive stimulation (interval between
photostimulation trains, ~10 min, see examples in Figures 2 A and S2). Despite individual variability (Fig-
ure S3), mean responses of layer V pyramidal cells to five successive photostimulus trains exhibited roughly
stable amplitudes that ranged between 86\% and 100\% of the first response (n = 18 cells, N = 3
slices from three mice; Figures 2 A and S3 and Table 1). Essentially similar results were obtained in the
dendrites of these neurons (Figure S4), although larger mean amplitudes were observed in dendrites than in
somata (Figure S4 and Table 1). The increase of the FRET ratio elicited by the first photostimulation in

Figure 2. Imaging NA Transmission Triggered by Photostimulation of LC Fibers in Cortical Slices

Two-photon imaging of layer V pyramidal neurons expressing the AKAR3EV sensor.

(A) Top: The grayscale image shows the F535 intensity and pseudocolor images show variations of the F535/F480 ratio
coded by pixel hue) and F535 intensity (encoded by pixel intensity), in pyramidal cells during the course of the
experiment. Middle: variations of the F535/F480 emission ratio measured at the soma of these individual pyramidal
neurons (n = 7, gray traces, mean trace in red), in response to burst photostimulation (arrowheads) and to bath application
of forskolin (FSK, 12.5 \mu M). Arrows indicate time points corresponding to pseudocolor images in the upper panel.
Bottom: Results obtained in n = 18 layer V pyramidal cells from N = 3 slices, from one mouse, each. Differences are not
significant. Values are mean ± SEM.

(B) Effects of \( \beta_1 \) (CGP20712, 100 nM) and \( \alpha_2 \) (yohimbine, 1 \mu M) adrenoceptor antagonists and of the NA transporter
inhibitor reboxetine (100 nM) on responses to photostimulation. Left: responses of individual pyramidal neurons (gray
traces, mean trace in red) in the presence or absence of drug. Right, from top to bottom: results obtained in N = 4 and n =
18, N = 3 and n = 25, N = 5 and n = 27. * Significantly different from the response to the first photostimulation. Values are
mean ± SEM. *p < 0.05.
### Table 1. Properties of PKA Responses Obtained in Pyramidal Neurons from Cortical Slices upon Five Consecutive Photostimulation Episodes

Each photostimulation episode consisted of 5-ms light pulses at 10 Hz for 5 s AKAR3EV imaging. R = F535/F480. Time to peak refers to the time between the beginning of the photostimulation and the maximum of the response. The value has been calculated for each response and the mean value has been replaced by a 30-s-long time interval to take in account the acquisition rate (see Methods). The decay time \( \tau \) was calculated by fitting the decay of the response with a single exponential. Time to peak and decay were calculated for responses in somata. Experiments were performed on four and three independent brain slices for layers II/III and V, respectively. Values are mean ± SEM. * and † indicate values significantly different (p < 0.05) from those obtained for the first and second stimulation, respectively. NM, not measured.

| Photostimulation | Cortical Layer | 1 | 2 | 3 | 4 | 5 | Forskolin |
|------------------|---------------|---|---|---|---|---|-----------|
|                  | II/III | V | II/III | V | II/III | V | II/III | V | II/III | V | II/III | V |
| Amplitude in somata (ΔR/R0%) | | | | | | | | | | | | | |
| 1st stim. | 7.3 ± 0.5 (n = 34) | 7.4 ± 1.2 (n = 18) | 6.0* ± 0.4 (n = 34) | 6.1 ± 0.9 (n = 18) | 4.9*† ± 0.3 (n = 34) | 5.6 ± 0.8 (n = 18) | 4.6*† ± 0.4 (n = 34) | 5.7 ± 0.9 (n = 18) | 4.5*† ± 0.3 (n = 34) | 6.4 ± 1.0 (n = 18) | 20.5 ± 0.9 (n = 34) | 25.7 ± 2.0 (n = 18) |
| Normalized amplitude in somata (% first stim.) | 100 ± 7.3 (n = 34) | 100 ± 9.8 (n = 18) | 77.6* ± 5.6 (n = 34) | 91.4 ± 11.5 (n = 18) | 58.36* ± 5.6 (n = 34) | 99.4 ± 19.9 (n = 18) | 59.7* ± 5.0 (n = 34) | 85.8 ± 13.6 (n = 18) | 53.1* ± 4.7 (n = 34) | 99.6 ± 12.9 (n = 34) | NM | NM |
| Amplitude in dendrites (ΔR/R0%) | NM | 10.2 ± 1.8 (n = 31) | NM | 8.5 ± 1.5 (n = 31) | NM | 8.1 ± 1.5 (n = 31) | NM | 8.2 ± 1.5 (n = 31) | NM | 7.9 ± 1.4 (n = 31) | NM | 28 ± 5 (n = 31) |
| Time to peak range (s) | [60–90] (n = 24) | [60–90] (n = 17) | [60–90] (n = 24) | [60–90] (n = 17) | [60–90] (n = 24) | [60–90] (n = 17) | [60–90] (n = 24) | [60–90] (n = 17) | [60–90] (n = 24) | [60–90] (n = 17) | NM | NM |
| Decay (σ in s) | 221.7 ± 17.1 (n = 20) | 178.4 ± 17.7 (n = 12) | 220.8 ± 26.4 (n = 20) | 147.4 ± 10.6 (n = 12) | 223.6 ± 27.4 (n = 19) | 172.2 ± 27.8 (n = 12) | 233.9 ± 24.4 (n = 20) | 159.7 ± 15.6 (n = 12) | 212.6 ± 28.3 (n = 20) | 140.5 ± 15.2 (n = 12) | NM | NM |
the soma of these neurons was 7.4 ± 1.2%, which represents roughly one-third of their response to FSK (25.7 ± 2.0% increase of FRET ratio, Table 1). For comparison, PKA responses of cortical pyramidal cells to bath application of a saturating dose of NA reach 60% of the response to FSK (Nomura et al., 2014), corresponding to only twice the response amplitude measured here upon LC fiber photostimulation. This suggests that a significant proportion of the catecholamine receptors expressed in pyramidal cells are activated upon burst photostimulation of LC fibers. Responses of layer V pyramidal cells peaked within 1.5 min after photostimulation and decayed with time constants ranging from 140 ± 15 to 178 ± 18 s (Table 1). Response kinetics did not significantly differ between the five photostimulation trains (Table 1). Essentially similar results were obtained in layer II/III pyramidal neurons, except that response amplitude gradually declined upon repetitive stimulation (Figure S2 and Table 1). We thus focused on somata of layer V pyramidal neurons in subsequent brain slice experiments. These results suggest that selective photostimulation of cortical LC fibers triggers the release of endogenous catecholamines, which robustly activate cAMP/PKA signaling in pyramidal neurons, thereby allowing the pharmacological characterization of their responses.

PKA Dynamics Evoked by LC Fibers Photostimulation Are Mediated by NA Receptors

LC fibers can release both NA and DA in the forebrain (Devoto et al. 2001, 2005, 2008, 2015; Kempadoo et al., 2016), and both NA and DA receptors are widely expressed in the neocortex, beyond the restricted distribution of dopaminergic fibers (Anano and Sibley 1994; Khan et al., 1998; Luedtke et al., 1999; Rivera et al., 2008; Oda et al., 2010; Nomura et al., 2014). In the parietal cortex, DA fibers are scarce and restricted to layer VI, but Gs-coupled D1-5 and Gi-coupled D2-like DA receptors are nonetheless functionally expressed and coupled to cAMP/PKA signaling in layers II-V pyramidal cells (Nomura et al., 2014). We thus characterized receptors involved in light-induced responses using specific antagonists of Gs- and Gi-coupled NA and DA receptors. Following two photostimulation trials in control condition to assess the stability of the PKA response, the third photostimulation was performed in the presence of antagonist pre-applied for at least 10 min. Response amplitudes thereafter are expressed as percentage of the first response amplitude. Perfusion of the specific β1-adrenergic receptor antagonist CGP20712 (100 nM) almost completely prevented PKA activation (Figure 2B and Table 2). The amplitude of the residual response was 12 ± 3% of the first response (N = 4, n = 18). Conversely, application of the D1/D5 receptor antagonist SCH23390 (1 µM) did not significantly modify the response (117 ± 10%, N = 3, n = 25; Table 2). These results indicate that light-induced PKA activation in pyramidal cells was essentially mediated by β1-adrenergic receptors, without detectable contribution of the D1/D5 receptors functionally expressed in these cells (Nomura et al., 2014). We next tested whether responses to photostimulation also involve Gi-coupled receptors by applying the a2 adrenoceptor antagonist yohimbine (1 µM) and the D2-like dopaminergic receptor antagonist haloperidol (10 µM). Application of yohimbine resulted in an increase of response amplitude (143 ± 15%, N = 3, n = 21; Figure 2B and Table 2), whereas haloperidol did not significantly alter the response (87 ± 10%, N = 3, n = 21; Table 2). Hence, PKA activation upon photostimulation of LC fibers is mediated by Gs-coupled β1 adrenoceptors and is dampened by co-activation of Gi-coupled a2 adrenoceptors. This suggests that NA released by LC fibers activates both receptor types in pyramidal cells, as also observed upon application of exogenous NA (Nomura et al., 2014). Conversely, these results rule out the involvement of DA receptors in PKA responses of pyramidal cells to burst photostimulation of LC fibers.

Cortical NA Transmission Is Critically Controlled by Catecholamine Reuptake

The clearance of released catecholamines involves their recapture into catecholaminergic axons or neighboring cells by various transporters showing low selectivity for NA versus DA (Grundemann et al., 1998; Torres et al., 2003). We tested the role of the NA transporter (NET), DA transporter (DAT), and extraneuronal transporters of the organic cation transporter (OCT) family in light-induced PKA responses. We found that application of the specific NET inhibitor reboxetine (100 nM) largely increased the response to photostimulation, whose amplitude was more than twice that of the response to the first photostimulation (251 ± 23%, N = 5, n = 27; Figure 2B). This was accompanied by an increase of the response time to peak, without significant modification of decay kinetics (Table 2). In contrast, neither the specific DAT inhibitor GBR12783 (100 nM) nor the OCT inhibitor corticosterone (100 µM, Grundemann et al., 1998) significantly modified the amplitude of the PKA response (GBR12783: 86 ± 8%, N = 3, n = 20; corticosterone: 90 ± 8%, N = 3, n = 24; Table 2). These results indicate that the effects of NA released upon burst photostimulation of LC fibers are strongly reduced by reuptake through NET transporters. Conversely, DAT and OCT were not involved in shaping cortical catecholaminergic transmission from LC fibers.
Photostimulation consisted of 5-ms light pulses at 10 Hz for 5 s, and PKA activity was measured in somata of layer V pyramidal cells from three to five independent brain slices. Values are mean SEM. * and n correspond to values significantly different (p < 0.05) from the values obtained for the first and second stimulation, respectively.

Table 2. Pharmacological Properties of Responses to Photostimulation of LC Fibers in Cortical Slices
Photostimulation consisted of 5-ms light pulses at 10 Hz for 5 s, and PKA activity was measured in somata of layer V pyramidal cells from three to five independent brain slices. Values are mean ± SEM. * and n correspond to values significantly different (p < 0.05) from the values obtained for the first and second stimulation, respectively.

Cortical NA Transmission In Vivo
In order to validate, in the intact neocortex, the main results obtained in brain slice, we used in vivo photostimulation of LC fibers and imaging of pyramidal neurons expressing the GAKDYmut PKA sensor and TurboRFP (see Methods and Figure 1). We monitored GAKDYmut fluorescence intensity in dendrites and somata during time windows corresponding to three consecutive bursts of photostimulation (interval between bursts, 5 min, Figure 3A). The photostimulation pattern used in slices (5 ms pulse at 10 Hz for 5 s) failed to evoke consistent responses in vivo (Figure S5), presumably because side illumination through the cranial window (see Methods) and tissue light scattering properties resulted in suboptimal ChR2 activation in depth of the cortex. We thus increased photostimulation (5 ms at 20 Hz for 10 s) and observed clear responses, which consisted in transient GAkdYmut fluorescence increases in dendrites (see example in Figure 3A). Mean responses of dendritic ROIs reached 5.7 ± 0.7% of baseline GAkdYmut fluorescence (n = 960 ROIs, 12 stimulation trials performed in N = 6 mice). No change in dendritic TurboRFP fluorescence was observed in the same experiments (Figure S6). In order to establish a criterion to select regions of interest (ROIs) for further quantitative analyses of responses, fluorescence changes of each dendritic ROI were submitted to a statistical test to determine their individual responsiveness to photostimulation (see Methods). Only ROIs whose post-stimulus fluorescence significantly differed from baseline were considered as responsive. Among the 90 dendritic ROIs shown in Figure 3A, we found that 60, 67, and 58 ROIs significantly responded to the first, second, and third successive stimulation trials, respectively, with 42.2% of these 90 ROIs responding to all three photostimulation trials (Figure S7). Responses of individual dendritic ROIs showed ROI-to-ROI and stimulus-to-stimulus variability (Figure S7), but averaged responses of pooled dendritic ROIs had stable amplitude upon repetitive photostimulation (see example in Figure 3A). Although responses to photostimulation were also detectable in somatic ROIs (Figure 3A), they were much less frequent compared with dendrites. Indeed, using the same statistical criterion as above, we found that 10, 7, and 8 of 35 somatic ROIs significantly responded to the first, second, and third successive stimulation.
trials, respectively, with only 5.7% of these 35 somatic ROIs responding to all three photostimulation trials (Figure S8). Furthermore, somatic fluorescence changes were of much smaller amplitude than dendritic ones (p < 0.001, Mann Whitney test, Figure S8), consistent with the differential responsiveness of these subcellular domains to β-adrenoceptor-induced cAMP/PKA signaling (Castro et al., 2010). When considering all responsive dendritic ROIs over the three trials shown in Figures 3A and S7, fluorescence increases were larger than 2% of baseline in 99.5% of these ROIs (n = 59 + 67 + 58 = 184 of 185 traces, see above). Conversely, among fluorescence responses higher than 2% (225 traces), 81.8% of traces passed our statistical test. Thus, in the rest of this in vivo study, we only considered dendritic fluorescence changes larger
than 2% of baseline for analyses of light-induced responses. Among ROIs imaged in control conditions (n = 960 ROIs, 12 stimulation trials performed in N = 6 mice), those showing fluorescence changes >2% (76 ± 3% of ROIs) were selected to calculate mean response kinetics, yielding onset (τ = 14.7 ± 4.5 s) and decay (τ = 154.1 ± 31.0 s) time constants. We then imaged GAkdYmut fluorescence in dendrites to characterize the pharmacological properties of responses to effective photostimulation (5 ms pulse, 20 Hz, 10 s) of LC fibers in vivo. In each experiment, photostimulation was performed first in control condition, then 15 min after i.p. injection of saline, and finally after i.p. injection of drug. Response amplitudes remained stable after saline injection (103 ± 4% of control, n = 520 ROIs, N = 6) but markedly changed after drug administration (Figures 3B and 3C). Indeed, the response was virtually abolished 15 min after injection of the β-adrenoceptor antagonist propranolol (4 mg/kg, - 11 ± 5% of control, n = 330 ROIs, N = 3) and only partially recovered 30 min after propranolol injection (24 ± 5% of control, n = 330 ROIs, N = 3, Figure 3B). Conversely, response amplitude was largely enhanced 15 min after injection of reboxetine (10 mg/kg, 182 ± 14% of control, n = 190, N = 3), reaching a maximum 20 min after injection (250 ± 20% of control, n = 125, N = 2) before declining 30 min after reboxetine treatment (214 ± 13% of control, n = 190, N = 3, Figure 3C). These results confirm, in the cortex in vivo, that burst photostimulation of LC fibers triggers the release of NA, which activates PKA signaling in pyramidal neurons via β-adrenoceptors, and whose effects are limited by reuptake through NET transporters.

**DISCUSSION**

We combined imaging of PKA sensors expressed in pyramidal neurons with photostimulation of ChR2-expressing LC fibers in the neocortex, in acute slices and in vivo. We found that burst photostimulation of LC fibers elicited fluorescence changes of the sensors, indicative of PKA activation in somata and dendrites of individual pyramidal cells. These responses were transient, and their amplitude was stable upon repetitive stimulation. The use of selective antagonists showed that responses were mediated by β1 adrenoceptors and dampened by co-activation of α2 adrenoceptors, but did not involve DA receptors, showing that NA was the major catecholamine released from LC fibers. Responses to photostimulation were largely increased upon NET inhibition, indicating that the clearance of NA released from LC fibers is primarily achieved by reuptake through this transporter. Our results demonstrate that the present optogenetic tools provide sensitive and reliable means to analyze quantitatively the dynamics of NA transmission in the brain at cellular and subcellular levels.

**Burst Photostimulation of LC Fibers Triggers PKA Activation**

Biosensors of the AKAR series, which include AKAR3EV and GAkdYmut, are based on conformational changes upon phosphorylation of a PKA substrate domain and selectively report PKA activity in cellulo (Zhang et al. 2001, 2005; Allen and Zhang 2006; Gervasi et al., 2007; Hu et al., 2011; Bonnot et al., 2014). Our results thus indicate that burst photostimulation of LC fibers evoked robust cAMP/PKA signaling transients, clearly detectable in real time at the level of single pyramidal neurons and their dendrites, in slices and in vivo. Light-induced responses in slices reached one-third the amplitude of responses to FSK, indicating that NA released by LC fibers is able to activate a significant proportion of somatodendritic β1 adrenoceptors present at the membrane of pyramidal neurons, despite reuptake by NET. For comparison, the maximal effect of exogenous NA is about two-thirds of the FSK effect on cortical pyramidal cells (Nomura et al., 2014), suggesting that NA released upon burst photostimulation in slices reached the membrane of these neurons at a mean concentration close to its EC50 (25 nM, Nomura et al., 2014). Smaller light-induced responses were observed in vivo, reaching only ~8% of the FSK effect previously measured in slice (74% increase of GAkdYmut fluorescence, Bonnot et al., 2014), consistent with less efficient photostimulation of LC fibers in vivo. Indeed, in these latter conditions (see Methods), illumination angle, light reflection/refraction at cranial glass window interfaces, as well as light scattering and absorption through blood-perfused tissue likely resulted in low illumination intensity of LC fibers in the imaging field in layer II/III. Dendritic responses were larger than somatic responses both in slices and in vivo. The differential responsiveness of these cellular domains to β-adrenoceptor-induced cAMP/PKA signaling in cortical pyramidal cells is documented and attributed to the effect of phosphodiesterase type 4 that prevents diffusion of cAMP from the submembrane space to the cytosol, thereby favoring domains with a high surface to volume ratio, such as dendrites (Castro et al., 2010). Responses were time-locked on the photostimulus, and imaging at high temporal resolution in vivo (6 Hz) revealed rapid onset (τ = 15 s) and slower decay (τ = 154 s) kinetics, in good correspondence with our results in brain slices. Prior observations indicate that the time course of light-induced responses of the biosensors followed that of the PKA signal in pyramidal cells: (1) responses of AKAR probes to cAMP photo-uncaging peak within 30 s (Zhang et al., 2005; Gervasi et al., 2007), (2)
responses of AKAR probes and of cAMP biosensors exhibit similar onset kinetics upon activation of Gs-coupled receptors (Castro et al., 2010), (3) relaxation kinetics of AKAR probes (i.e., their dephosphorylation by cellular phosphatases) are faster than the decay of the present responses to photostimulation (Allen and Zhang 2006; Dunn et al., 2006; Gervasi et al., 2007). Hence, our results indicate that burst activation of LC fibers for a few seconds rapidly triggers robust, minutes-lasting cAMP/PKA signals in cortical pyramidal cells.

PKA Dynamics Evoked by Burst Photostimulation Rely on NA but Not DA Transmission

The activation of PKA in pyramidal cells was essentially mediated by β1 adrenoceptors, consistent with NA release from LC fibers and with the effect of exogenous NA (Nomura et al., 2014). The α2 antagonist yohimbine increased the amplitude of light-induced responses by 140%, an effect larger than that observed on responses to exogenous NA (117%, Nomura et al., 2014). This suggests that recruitment of α2 adrenoceptors dampens the response both by inhibiting adenylate cyclase in pyramidal cells and by acting at LC fibers to decrease NA release, as classically described (Starke 2001). The NET inhibitor reboxetine dramatically enhanced the amplitude of light-induced responses but did not change their decay kinetics. This indicates that NA reuptake by NET limits the number of receptors activated upon NA release by minimizing the spatial extent of NA diffusion but is not involved in limiting the duration of NA transmission events. We exclude a role of DAT and OCT in shaping light-induced responses but cannot rule out a contribution of the corticosterone-insensitive uptake2 transporter PMAT (Engel et al., 2004; Engel and Wang 2003) in NA clearance, since the fluorescence of its antagonist decynium-22 (Inyushin et al., 2010) precluded its use in our imaging study.

Devoto et al. (2001, 2005, 2008, 2015) have shown that electrical stimulation of the LC in vivo results in cortical release of both NA and its biosynthetic precursor DA. However, although both D1/D5 and D2-like receptors are functionally coupled to adenylate cyclase in pyramidal cells (Nomura et al., 2014), we failed to detect their involvement in PKA responses to photostimulation in cortical slices. This discrepancy presumably relates to the low potency of DA (notably a ~20-fold higher EC50 than NA) at stimulating PKA in cortical pyramidal cells (Nomura et al., 2014), which suggests in turn two hypotheses regarding DA transmission from cortical LC fibers. (1) DA transmission may occur through DA receptors preferentially coupled to phospholipase C, as documented for D1, D2, and D5 receptors (reviewed by Beaulieu and Gainetdinov 2011), thereby escaping our present detection using PKA biosensors. (2) Alternatively, burst photostimulation in cortical slices may elicit only minimal DA release, whereas intense and prolonged activation of LC fibers in vivo (as performed by Devoto et al.) may favor DA release and thus lead to the recruitment of DA receptors coupled to adenylate cyclase. Indeed, in these latter conditions, DA may accumulate in vesicles pending depletion of vesicular DBH that is co-released with NA (Weinshilboum et al., 1971) or may be released by reverse transport from the axoplasm upon inversion of ion gradients (Levi and Raiteri 1993; Sitte and Freissmuth 2010). It is noteworthy that our in vivo data, i.e., the complete block of light-induced responses by the β-adrenergic antagonist propranolol, do not rule out DA receptor activation, since propranolol also prevents D1/D5 receptor-mediated effects of exogenous DA on pyramidal cells (Nomura et al., 2014). Hence, further work is required to assess and characterize NA and DA co-transmission from cortical LC fibers.

Phasic Noradrenergic Transmission and Burst Firing of LC Neurons

The firing of LC neurons consists either of single spikes emitted at low frequency (tonic mode) or of bursts of action potentials (phasic mode) that involve synchronization of the population of LC neurons (Aston-Jones and Bloom, 1981a; 1981b). The burst firing of LC neurons is associated with a peak of NA release and has important behavioral correlates (Florin-Lechner et al., 1996; Aston-Jones and Cohen 2005). Our results obtained upon bulk photostimulation indicate that a single episode of synchronized burst firing in LC neurons has rapid and long-lasting effects on the cortical network, and that phasic NA transmission operates with higher intensity and temporal precision than generally thought of a neuromodulatory system relying on volume transmission. Light-induced responses exhibited cell-to-cell and dendrite-to-dendrite variability, as observed upon exogenous NA application (Nomura et al., 2014), presumably reflecting variations in distance to NA release sites, involvement of α2 adrenoceptors, and basal level of cAMP/PKA activity. Nonetheless, population responses were robust and reproducible. Gs-coupled receptors enhance neuronal excitability by decreasing PKA-sensitive membrane potassium currents with onset kinetics similar to those of AKAR probes responses (Gervasi et al., 2007; Hu et al., 2011). Based on these latter studies, the present light-induced PKA responses (~30% of FSK effect) would translate into a ~50% reduction of PKA-sensitive
potassium currents in pyramidal neurons. Hence, a single episode of synchronized burst firing in LC neurons likely induces a rapid and substantial increase of the excitability of pyramidal neuron populations with potentially important acute and long-term effects on the function of the cortical network.

**Imaging the Cellular Dynamics of Catecholaminergic Transmission**

The detectability of endogenous catecholaminergic transmission in vivo has been recently demonstrated using fluorescent cAMP or PKA activity sensors expressed in neurons or glia (Goto et al., 2015; Ma et al., 2018; Oe et al., 2020). Here we show that two-photon imaging of FRET or single-wavelength PKA sensors allows the quantitative analysis of key properties (intensity, kinetics, receptors, and reuptake transporters) of NA transmission events in cortical neurons and their dendrites, in brain slice and in vivo. Our results confirm that any optical sensor of cAMP/PKA signaling is suitable for the imaging of endogenous NA transmission, pending sufficient dynamic range. In this regard, the bright AKAR3EV FRET sensor proved useful for the imaging in diffractive adult slice, despite smaller dynamic range than GAKdYmut (ΔR/R0: ~25% versus ΔF/F0: 74%, respectively, in response to FSK in similar experimental conditions; see Table 1 and Bonnot et al., 2014). Conversely, GAKdYmut enabled fast acquisition (375 μm frame at 6.2 Hz) to analyze PKA response kinetics with high resolution in vivo. This single-wavelength green sensor is well adapted to two-photon imaging, as fluorescence can be collected in its entire emission spectrum for increased sensitivity, and is easily compatible with other sensors for multicolor imaging, as exemplified here by the dual monitoring of GAKdYmut and TurboRFP in vivo. It is thus likely that GAKdYmut can be combined with single-wavelength cAMP or Ca2+ red sensors with large dynamic range (e.g., Dana et al., 2016; Ohta et al., 2018; Oe et al., 2020) for future examination of endogenous catecholaminergic transmission dynamics and their impact on network electrical activity.

**Limitations of the Study**

The amplitude and kinetics of cortical NA transmission events occurring upon spontaneous burst firing of LC neurons in awake mice may largely differ from those presently reported using optogenetic stimulation in conditions of low network activity found in slices and anesthetized animals. Another limitation concerns the use of biosensors for PKA activity, whose readout of receptor binding/unbinding is presumably delayed as compared with sensors for upstream signaling events (e.g., cAMP changes) and which cannot detect the possible recruitment of Gq-coupled α1 adrenoceptors (Cotecchia 2010) and DA receptors (Beaulieu and Gainetdinov 2011) elicited by LC fibers stimulation.

**Resource Availability**

**Lead Contact**

Further Information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Bertrand Lambolez (bertrand.lambolez@upmc.fr).

**Material Availability**

This study did not generate new unique reagents.

**Data and Code Availability**

The data and codes reported in this study are available from the Lead Contact on request.

**METHODS**

All methods can be found in the accompanying Transparent Methods supplemental file.

**SUPPLEMENTAL INFORMATION**

Supplemental Information can be found online at https://doi.org/10.1016/j.isci.2020.101710.

**ACKNOWLEDGMENTS**

The authors thank Katia Boutourlinski, Bruno Giros, Hirokazu Tanimoto, the IBPS Cell Imaging and Animal Facilities, and the OIST Animal Resources and Imaging sections for their valuable help. This work was supported by grants from Ecole des Neurosciences de Paris (“Network for Viral Transfer”), Fondation pour la Recherche sur le Cerveau/Rotary Club de France, Agence Nationale de la Recherche (“IHU Institut de Neurosciences Translationelles de Paris,” ANR-10-IAIHU-06), and the OIST Graduate University. S.N. was
recipient of a Nakajima Foundation fellowship. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

AUTHOR CONTRIBUTIONS
S.N., L.T., I.C., and R.H. performed experiments. S.N., L.T., B.K., and R.H. analyzed data. S.N., B.K., R.H., and B.L. designed the study. S.N., B.K., R.H., and B.L. supervised the work and wrote the manuscript, with input from the other authors.

DECLARATION OF INTERESTS
The authors declare no competing interests.

Received: March 18, 2020
Revised: July 29, 2020
Accepted: October 16, 2020
Published: November 20, 2020

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Supplemental Information

Combined Optogenetic Approaches Reveal
Quantitative Dynamics of Endogenous Noradrenergic Transmission in the Brain

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Supplementary Figure 1. Test of different patterns of LC fibers photostimulation in cortical slices (related to figure 2).

Two-photon imaging of layer V pyramidal neurons expressing the AKAR3EV sensor. Traces show the F535/F480 emission ratio measured at the soma of individual cells (grey traces, mean trace in red) upon burst photostimulation (arrowheads) of indicated patterns, and upon bath application of forskolin (FSK, 12.5 µM). (A) Data obtained in n=8 pyramidal neurons from N=1 slice. Note the decrease of response amplitude upon repetitive stimulation. * significantly different from the response to the first photostimulation. (B) Data obtained in n=9 cells, N=1 slice. Photostimulation elicited a 3.4 ± 0.4 % increase in ΔR/R0. (C) Data obtained in n=19 cells, N=2 slices. Note the absence of response, presumably related to ChR2 desensitization during 100 ms light pulses.
Supplementary Figure 2. Responses of layer II/III pyramidal neurons to photostimulation of LC fibers in cortical slices (related to figure 2 and table 1).

Two-photon imaging of layer II/III pyramidal neurons expressing the AKAR3EV sensor. Traces show variations of the F535/F480 emission ratio measured at the soma of individual pyramidal neurons (n=5, grey traces, mean trace in red), in response to burst photostimulation (arrowheads) and to bath application of forskolin (FSK, 12.5 µM). The bar graph summarizes results obtained in n=34 cells from N=4 slices, with differences from the response to the first photostimulation being statistically significant.

Supplementary Figure 3. Variability of layer V pyramidal neurons responses to photostimulation of LC fibers in cortical slices (related to figure 2).

Two-photon imaging of layer V pyramidal neurons expressing the AKAR3EV sensor. Traces show the F535/F480 emission ratio measured at the soma of individual cells (n=18, from 3 experiments as indicated, mean trace in black). Green traces correspond to the experiment shown in Fig. 2A). To compensate for variable baseline drifts during the course of the experiments, individual traces were realigned by normalizing to the baseline value measured before each stimulation trial.
Supplementary Figure 4. Comparison between dendritic and somatic responses triggered by photostimulation of LC fibers in cortical slices (related to figure 2).

Left: Two-photon imaging of neuronal somata and dendrites expressing the AKAR3EV sensor in layer V. The greyscale image shows the F535 intensity and dendritic (red) or somatic (blue) ROIs delineated for measurements of F535/F480 emission ratio. Right: Traces show mean variations of the F535/F480 emission ratio measured at these somatic (n=8, black) and dendritic (n=14, red) ROIs in response to burst photostimulation (arrowheads) and to bath application of forskolin (FSK, 12.5 µM).
Supplementary Figure 5. Test of different patterns of LC fibers photostimulation in vivo (related to figure 3).

Two-photon imaging of cortical neurons expressing the GAkdYmut sensor. Traces show averaged GAkdYmut fluorescence intensity of 50 dendritic ROIs before and after photostimulation of indicated patterns delivered during the recording shut off intervals. Note the difference with clear responses shown in Fig. 3.
Supplementary Figure 6. LC fibers photostimulation in vivo does not trigger TurboRFP fluorescence changes (related to figure 3).

Two-photon imaging of cortical neurons co-expressing TurboRFP and the GAkdYmut sensor. **Upper:** Raster plots show TurboRFP fluorescence intensity (calibration bar: ΔF/F) of dendritic and somatic ROIs corresponding to GAkdYmut recordings shown in Fig. 3A, before and after each of three consecutive photostimulation trials (darker vertical bar, 5 ms pulses at 20 Hz for 10 s). **Lower:** Traces show averaged fluorescence intensity of the same ROIs.
Supplementary Figure 7. Responses of individual dendritic ROIs to photostimulation of LC fibers in vivo (related to figure 3). (A) Rasterplot showing responsiveness of 90 dendritic ROIs arranged from top to bottom by increasing response amplitude upon three successive photostimulation trials (stim). ROI responsiveness was determined by comparing the fluorescence signals before and after stimulation using a Kolmogorov-Smirnov statistical test (see Methods). (B) Distribution of the same individual ROIs as a function of their number of responses to the three stimulation trials. (C) Distribution of amplitudes of all statistically significant responses of the same ROIs across the three stimulation trials. 99.5% of these significant responses corresponded to fluorescence increases larger than 2% of baseline. (D) Fluorescence traces of the four ROIs exhibiting the largest responses to the three stimulation trials (t=0, end of the stimulus). The signal was smoothed by median filtering using a 20 datapoint sliding window.
Supplementary Figure 8. Responses of individual somatic ROIs to photostimulation of LC fibers in vivo (related to figure 3). (A) Rasterplot showing responsiveness of 35 somatic ROIs arranged from top to bottom by increasing response amplitude upon three successive photostimulation trials (stim). ROI responsiveness was determined by comparing the fluorescence signals before and after stimulation using a Kolmogorov-Smirnov statistical test (see Methods). (B) Distribution of the same individual ROIs as a function of their number of responses to the three stimulation trials. (C) Fluorescence changes normalized to baseline in individual somatic and dendritic ROIs upon three successive photostimulation trials. Dendritic fluorescence changes were larger than somatic ones. (D) Fluorescence traces of the two somatic ROIs showing statistically significant responses to the three stimulation trials (t=0, end of the stimulus). The signal was smoothed by median filtering using a 20 datapoint sliding window.
Transparent Methods

Animals

Experiments were in accordance with the European Communities Council Directive 86/609/062, approved by the Charles Darwin ethics committee (project 01907.1) and the OIST Institutional Animal Care and Use Committee, and performed in accredited facilities. Transgenic DBH-Cre mice were a gift from Bruno Giros [MMRRC line: Tg(Dbh-cre)KH212Gsat/Mmucd, stock number 032081-UCD (Gong et al. 2007)]. Genotyping of DBH-Cre mice was done by PCR with primers: forward AATGGCAGAGTGGGGTTGGG, reverse CGGCAAACGGACAGAAGCATT (225 bp). Animals were maintained in a 12 h light–12 h dark cycle, in stable conditions of temperature (22°C), with food and water available ad libitum. Data were collected from 29 mice aged 5-7-weeks at the time of first viral injection, including 9 females and 13 males for brain slice experiments, and 7 females for in vivo experiments.

Recombinant viruses

The adeno-associated virus (AAV) driving Cre-dependent expression of a fusion protein containing channelrhodopsin 2 (ChR2) and the yellow fluorophore YFP (AAV2/1-EF1a-DIO-hChR2(H134R)-EYFP-WPRE-HGHpA) was produced from Addgene plasmid #20298 (a gift from Karl Deisseroth) at vector core facilities of Nantes University (France, titer: 3x10^{11} gc/ml) and Pennsylvania University (USA, titer: 8x10^{12} gc/ml). The Sindbis virus encoding the AKAR3EV PKA activity sensor (Komatsu et al. 2011) was produced as described (Hepp et al. 2007, SIN-AKAR3EV titer: 1x10^{9} ip/ml) and mixed with a Sindbis virus encoding the red fluorophore mCherry (SIN-mCherry, gift from Katia Boutourlinski, UMR8246 Paris) at a ratio 40:1. The AAV encoding the GAkdYmut PKA activity sensor (Bonnot et al. 2014) was custom made and produced by the vector core facility of Pennsylvania University (AAV2/1-hSyn-GAkdYmut-hGH, titer: 4x10^{14} gc/ml), and mixed with AAV2/1-hSyn-TurboRFP-WPRE encoding the red fluorophore TurboRFP (titer: 4x10^{13} gc/ml, same supplier) at a ratio 1:1.

Expression of ChR2 and AKAR3EV for optogenetic experiments in cortical slices

Site- and cell-type-specific expression of ChR2 in NA neurons was achieved by viral transfer into the LC of DBH-Cre mice, exactly as described (Nomura et al. 2014). Mice (1-month-old) were anesthetized with isoflurane and the AAV-ChR2-YFP suspension (titer: 3x10^{11} gc/ml) injected bilaterally at coordinates from bregma: anteroposterior (AP) -5.45 mm, mediolateral (ML) ± 1 mm, dorsoventral (DV) -3.65 mm, through an internal canula at a rate of 0.1 µl/min for 10 min (1 µl/site). For postoperative care, ropivacaine (10 mg/ml) was added dropwise on the skin suture, and mice received buprenorphine (0.1 mg/kg). Eight to ten weeks after AAV-ChR2-YFP injection, mice were
anesthetized with 2 % isoflurane and the SIN-AKAR3EV/SIN-mCherry viral suspension mix was injected bilaterally in the primary sensory area of the parietal cortex at coordinates from bregma: AP -1.5 mm, ML ± 2 mm, DV -0.4-0.5 mm, through a glass pipette at a rate of 0.1 µl/min (2 µl/site). Mice received postoperative care as described above. Ten to twelve hours after Sindbis virus injection, mice were deeply anesthetized with 10 mg/kg ketamine and 0.1 mg/kg xylazine before transcardiac perfusion with 20 ml ice cold high-sucrose/low-calcium artificial cerebrospinal fluid (ACSF) containing: 73 sucrose, 85 NaCl, 2.5 KCl, 1.25 NaHPO4, 0.4 CaCl2, 7 MgCl2, 26 NaHCO3, 20 D-glucose, 5 Na pyruvate, 2 kynurenic acid (mM), 50 nM minocycline hydrochloride (Tikka et al. 2001), saturated with 5% CO2/95% O2. The brain was dissected and coronal slices (300 µm thick) of neocortex were cut in high-sucrose/low-calcium ACSF using a VT1000S Vibratome (Leica). Slices were kept at 33°C for 30 min in ACSF containing 126 NaCl, 2.5 KCl, 1.25 NaHPO4, 2 CaCl2, 1 MgCl2, 26 NaHCO3, 20 D-glucose, 5 Na pyruvate (mM), supplemented with 2 mM kynurenic acid and 50 nM minocycline hydrochloride, and bubbled with 5% CO2/95% O2. Slices were then allowed to recover at room temperature for 1 hour. Slices were transferred into a recording chamber perfused with ACSF (2 ml/min, 30 °C, pH 7.4) bubbled with 95% O2 and 5% CO2. Slices were inspected for mCherry expression using wide-field fluorescence to localize viral transduction area with minimal excitation of ChR2.

Expression of ChR2 and GAkdYmut for optogenetic experiments in vivo

ChR2 was expressed in NA neurons essentially as described above. The AAV-ChR2-YFP suspension (liter: 8x10^{12} gc/ml) was injected bilaterally into the LC of 5 to 7-week-old DBH-Cre mice at a rate of 10 nl/min (350 nl/site). Three weeks after AAV-ChR2-YFP injection, anesthesia was induced with a mixture of medetomidine (0.3 mg/kg), midazolam (4 mg/kg) and butorphanol (5 mg/kg), and a 5 mm craniotomy was performed under 2% isoflurane anesthesia. Then, 70-140 nl of the AAV-GAkdYmut/AAV-TurboRFP suspension mix was injected in layer V of the parietal cortex at coordinates from bregma: AP -1.5 mm, ML 1.7 mm, DV -0.6-0.7 mm, at a rate of 10 nl/min. A chronic cranial window (5 mm glass coverslip) was next mounted as described (Roome and Kuhn 2014). At the end of the surgery, mice received atipamezole (0.3 mg/kg) for recovery from anesthesia, and postoperative care as described above. Five to eight weeks after the last AAV injection, mice were head-fixed for imaging experiments performed under anesthesia with 1% isoflurane.

Optical stimulation and imaging in cortical slices

Two-photon images were obtained with a custom built two-photon laser scanning microscope described earlier (Castro et al. 2010, Bonnot et al. 2014), and using a 60X (0.9 NA, Olympus) water-immersion objective. Sampling rate was at 1.4 frame / s. For imaging of AKAR3EV, based
on fluorescence resonance energy transfer (FRET) between a blue donor (ECFP) and a yellow acceptor (YPet), 2-photon excitation of ECFP was performed at 850 nm with a power of 30 mW. ECFP and YPet fluorescence signals (peaks 480 and 535 nm, respectively) were filtered (Chroma Technology: E700 SP, Semrock: FF01-479/40, FF01-542/50), separated with a dichroic beamsplitter (FF506-Di02-25x36 Semrock) and simultaneously detected in two channels (H9305 photomultipliers, Hamamatsu). Images were acquired every minute prior to ChR2 photostimulation, and then every 30 s following photostimulation.

ChR2 photostimulation consisted in 470 nm light pulses delivered by a LED (LEDD1B, Thorlabs) through a 400 µm diameter fiber optic cable (ThorLabs) controlled with micromanipulators (ROE-200, Sutter Instrument). The light power at the tip of the optical fiber was 10 mW. The fiber tip was placed just above the slice surface and positioned 400 µm away from the center of the imaging field with an angle of 30° from the slice surface. Light pulses and a shutter (DSS25 and VDM1000, UNIBLITZ) protecting photomultipliers from light interference during photostimulation were controlled by custom software written in LabVIEW (National Instruments).

The 512x512-pixel images were exported using MATLAB (The Mathworks), and analyzed with custom macros derived from ImageJ (U.S. National Institute of Health). Fluorescence intensity of regions of interest (ROIs) was calculated for each time point from average intensity z-projection of 3 frames by averaging pixel intensity. Fluorescence changes were measured as the ratio R=F535/F480. Pseudocolor hue saturation value (HSV) encoding of fluorescence intensity and ratio was performed using MATLAB custom procedures written by Hirokazu Tanimoto (Yokohama City University). Decay time-constants were obtained using the exp-XOffset fit function of IgorPro6 (WaveMetrics) using the equation: $y_0 + A exp \left( -\frac{x-x_0}{\tau} \right)$

**Optical stimulation and imaging in the cortex *in vivo***

A combined wide-field/two-photon microscope (MOM, Sutter Instruments) with a femtosecond-pulsed Ti:sapphire laser (Vision II, Coherent) was used. To increase the point spread function of excitation the back aperture of the 25X water immersion objective (Olympus) was underfilled (spatial resolution 1 µm x 1 µm x 4 µm). The collar of the objective was adjusted to correct for the window glass thickness (170 µm). Simultaneous excitation of GAkdYmut (GFP-based single fluorophore sensor, Bonnot et al. 2014) and TurboRFP was performed at 950 nm with a typical power of 5-11 mW. Fluorescence was detected in two channels by GaAsP photomultipliers (Hamamatsu) in spectral windows 490-550 nm (GAkdYmut) and 600-700 nm (TurboRFP), separated by a 560 nm dichroic mirror (all Semrock). The microscope was controlled by a commercial software (MScan, Sutter Instruments). Sampling rate was 30.9 frame/s.

ChR2 photostimulation was done with a collimated blue LED (450-495 nm, BDX, X-Cite XLED1,
Lumen Dynamics) illuminating the area of the chronic cranial window with an angle of 30° from the glass surface. The power at the window glass surface was 0.35 mW/mm². Photomultipliers were manually shut off during photostimulation.

Movies (512 x 512 pixel per frame, corresponding to a field of view of 375 µm x 375 µm) were analyzed with custom made MATLAB code and ImageJ. The 30.9 Hz imaging data was temporally binned to 6.2 Hz. ROIs were selected from the GAKdYmut channel and relative fluorescence change \[ \frac{\Delta F}{F}(t) \] was calculated as: \[ \frac{\Delta F}{F}(t) = \frac{F(t) - F_{\text{baseline}}}{F_{\text{baseline}}} \], where \( F_{\text{baseline}} \) is the average fluorescence intensity during a 30 s interval ending 5 s before photostimulation. Onset and decay time-constants were obtained by fitting responses with the function: \[ A \left[ \exp\left(-\frac{t}{\tau_{\text{onset}}}\right) - \exp\left(-\frac{t}{\tau_{\text{decay}}}\right) \right] + B \]

**Histology**

The expression of ChR2-YFP and dopamine-β-hydroxylase was examined on fixed brain slices using a polyclonal rabbit anti DBH (1/1000, AB1538, Merck-Millipore) and a polyclonal chicken anti-GFP antibody (1/2000, GFP-1020, Aves Labs), as described (Nomura et al. 2014). Secondary antibodies were Alexa Fluor 488 goat anti chicken (1/1000, Life Technologies A11039) and goat anti Rabbit-Alexa 555 (1/1000, Life Technologies A-21428). Slices were mounted in fluoromount-G (Clinisciences). Images were acquired using a confocal laser scanning microscope (SP5, Leica) or using wide-field epifluorescence (DMR microscope, Leica) and processed with ImageJ. Following in vivo imaging experiments, brains were fixed, sliced in 60 µM sections, and images were acquired using a wide-field fluorescence microscope (Nikon Eclipse, Nikon).

**Drugs**

Drugs were purchased from Tocris or Sigma-Aldrich. Drugs were bath applied on brain slices: CGP20712 hydrochloride (100 nM), SCH23390 hydrochloride (1µM); Yohimbine (1 µM), Haloperidol (10 µM), Forskolin (12.5 µM), Reboxetine mesylate (100 nM), GBR12783 dihydrochloride (100 nM), Corticosterone (100 µM). For in vivo pharmacology, saline, Propranolol (4 mg/kg), Reboxetine mesylate (10 mg/kg) were injected intraperitoneally.

**Statistical analyses**

In this report, N represents the number of brain slices or mice tested while n represents the number of cells or ROIs on which measurements were performed. Data are presented as mean ± standard error of the mean (SEM). Statistical analyses were performed with MATLAB. Unless otherwise stated, one-way ANOVA followed by Tukey’s test was used to determine statistical
significance, except for analyses of single ROI responsiveness in vivo. To evaluate the responsiveness of individual ROIs in vivo, the fluorescence signal was binned 5 times to reach a 6.2 Hz sampling rate, and then normalized to the baseline calculated as the mean fluorescence intensity over the time window [-68.9; -20.4 s] (t=0 refers to the end of the photostimulation). Response amplitude measured over the interval [20.0; 52.4 s] (n=201 data points) was then compared to baseline signal (301 data points) using a Kolmogorov-Smirnov test. Throughout this manuscript, a p-value below 0.05 was considered statistically significant.

Supplemental references

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