Somatostatin interneurons activated by 5-HT$_{2A}$ receptor suppress slow oscillations in medial entorhinal cortex

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Abstract

Serotonin (5-HT) is one of the major neuromodulators present in the mammalian brain and has been shown to play a role in multiple physiological processes. The mechanisms by which 5-HT modulates cortical network activity, however, are not yet fully understood. We investigated the effects of 5-HT on slow oscillations (SOs), a synchronized cortical network activity universally present across species. SOs are observed during anesthesia and are considered to be the default cortical activity pattern. We discovered that (±)3,4-methylenedioxymethamphetamine (MDMA) and fenfluramine, two potent 5-HT releasers, inhibit SOs within the entorhinal cortex (EC) in anesthetized mice. Combining opto- and pharmacogenetic manipulations with in vitro electrophysiological recordings, we uncovered that somatostatin-expressing (Sst) interneurons activated by the 5-HT$_{2A}$ receptor (5-HT$_{2A}$R)
play an important role in the suppression of SOs. Since 5-HT2A signaling is involved in the etiology of different psychiatric disorders and mediates the psychological effects of many psychoactive serotonergic drugs, we propose that the newly discovered link between Sst interneurons and 5-HT will contribute to our understanding of these complex topics.
Introduction

5-HT is one of the most important neuromodulators in the central nervous system. Projections originating from the Raphe nuclei, the brain-stem structure that comprises the majority of 5-HT releasing neurons in the brain, innervate all cortical and sub-cortical areas (Descarries et al., 2010). 5-HT levels in the brain are closely linked to the sleep-wake cycle: the activity of serotonergic raphe neurons is increased during wakefulness, decreased during slow-wave sleep (SWS) and virtually silent during REM sleep (McGinty and Harper, 1976, Oikonomou et al., 2019, Unger et al., 2020). Cortical activity is also influenced by the behavioral state of the animal: SWS is generally associated with “synchronized” patterns of activity, characterized by low-frequency global fluctuations, whereas active wakefulness and REM sleep feature “desynchronized” network activity, in which low-frequency fluctuations are absent. The shifting of cortical networks between different patterns of activity is controlled, at least in part, by neuromodulators (Tukker et al., 2020, Lee and Dan, 2012). For instance, Acetylcholine (ACh) can profoundly alter cortical network activity by inducing desynchronization via activation of Sst interneurons (Chen et al., 2015). However, ACh is not solely responsible for suppressing cortical synchronized activity, as lesions of cholinergic neurons are not sufficient to abolish desynchronization (Kaur et al., 2008). On the other hand, blocking ACh and 5-HT transmission at the same time causes a continuous "synchronized" cortical state, even during active behavior, thus suggesting that 5-HT plays an important role in mediating transitions between different network states (Vanderwolf and Baker, 1986). In agreement with this line of thought, electrical and optogenetic stimulation of the Raphe nuclei reduce low frequency (1-10 Hz) power in the cortex, implying a reduction in neuronal synchronization at these frequencies (Puig et al., 2010, Grandjean et al., 2019). Moreover, optogenetic stimulation of serotonergic neurons is sufficient to awaken mice from SWS (Oikonomou et al., 2019). These data suggest a relationship between 5-HT levels and patterns of cortical activity (Lee and Dan, 2012, Harris and Thiele, 2011). The exact mechanism by which 5-HT modulates network activity in the cortex however, is still not fully understood.

Here, we used electrophysiological techniques together with pharmacology, optogenetics and pharmacogenetics to investigate the effect of 5-HT on SOs, a network oscillation characterized by synchronized transitions (< 1 Hz) between periods of high activity (upstates) and relative quiescence (downstates) (Steriade et al., 1993, Neske, 2015, Isomura et al., 2006). SOs are a global phenomenon observed throughout the cerebral cortex and are considered to be the default emergent activity of cortical networks during SWS and anesthesia (Neske, 2015, Sanchez-Vives et al., 2017, Wolansky et al., 2006). We performed our experiments in
the medial entorhinal cortex (mEC), a region where SOs can be recorded both under anesthesia and in acutely prepared cortical slices (Tahvildari et al., 2012, Beed et al., 2020). Pyramidal neurons located in L3 of mEC provide the excitatory drive underlying each upstate (Namiki et al., 2013, Beed et al., 2020). Similarly to other mammalian cortical areas, mEC comprises different types of inhibitory GABAergic neurons that can be grouped into three main classes according to immunoreactivity: parvalbumin (PV), somatostatin (Sst) and 5-HT₃ (Miao et al., 2017, Rudy et al., 2011). Most PV neurons target the soma and the spike initiation zone, have low input resistance and minimal spike frequency adaptation. Sst neurons are divided into two groups, one showing features similar to PV interneurons and a second one (i.e. Martinotti cells) that, in contrast, tend to form synapses onto the dendritic trees of their target cells, have high input resistance and show a considerable adaptation. 5-HT₃ neurons are usually located in the superficial layers, have high input resistance and mixed adaptation. While these different classes of interneurons are all depolarized during upstates, PV interneurons receive decidedly the strongest excitation (Tahvildari et al., 2012, Neske et al., 2015a). Recurrent excitation and temporal summation of inputs contribute to the transition from downstate to upstate (Sanchez-Vives et al., 2017, Tukker et al., 2020). This excitation propagates to L5 and differentially entrains L5a and L5b excitatory neurons. L5a neurons do not participate in SOs whereas L5b neurons are steadily synchronized. Both intrinsic and synaptic mechanisms have been implicated in upstate termination (Neske, 2015, Tukker et al., 2020). Activity-dependent K⁺ channels decrease excitability of neurons over time causing a generalized reduction of facilitation (Neske, 2015, Harris and Thiele, 2011). At the same time, blockage of GABAᵦ receptors significantly extends upstate duration (Craig et al., 2013) and inhibitory drive has been observed to increase during upstate termination (Lemieux et al., 2015). Sst interneurons, and in particular Martinotti cells, characterized by strongly facilitating synapses (Beierlein et al., 2003, Gibson et al., 1999) have been proposed as an important source of inhibition in the termination of upstates (Melamed et al., 2008, Krishnamurthy et al., 2012).

Our results show that MDMA and fenfluramine (Fen), two substitute amphetamines that induce robust 5-HT release, suppress SOs in anesthetized mice and, concurrently, activate a small group of neurons characterized by an intermediate waveform shape. Using cortical slices in vitro, we demonstrate that 5-HT inhibits SOs in the mEC and that Sst interneurons, activated by 5-HT₂₅R, are involved in the modulatory effect of 5-HT. While previous studies have shown that parvalbumin (PV) interneurons are excited by 5-HT₂₅R (Puig et al., 2010,
Athilingam et al., 2017), our results identify cortical Sst interneurons as novel targets of the 5-HT neuromodulatory system via 5-HT$_{2A}$R.

**Results**

**Pharmacological release of 5-HT inhibits SOs in anesthetized mice**

We investigated the effect of 5-HT on network activity in anesthetized mice using multisite silicon microelectrodes placed in the mEC L3, a region located in the medial temporal lobe interconnected to a variety of cortical and subcortical areas, including the Raphe nuclei (Figure 1A) (van Strien et al., 2009, Muzerelle et al., 2016). Under urethane anesthesia, EC, like the rest of the cortex, displays SOs (Figure 1B). As expected, we found that upstates were present synchronously in the local field potential (LFP) of all the recording channels (Figure 1–figure supplement 1), and that every upstate coincided with large increases in population spiking activity.

![Figure 1](https://bioxiv.org/figure.php?doi=10.1101/2020.05.26.113373&version=1&file=figure-1.png)

**Figure 1** | MDMA/Fen inhibit SOs in anesthetized mice. (A) Immunocytochemical analysis of an ePet-YFP mouse showing serotonergic fibers in medial entorhinal cortex, horizontal slice (M = medial, L = lateral, P= posterior, A = anterior). Scale bar: 20 µm. (B) LFP (top) and instantaneous population activity (bottom) of a representative *in vivo* recording during SOs (spikes/s units in thousands), cyan rectangles represent detected upstates. (C) 3D visualization of the microelectrode location of the recording shown in E. EC represented in grey. (D) Left: microelectrode tracks (red) of the recording shown in (E). Right: EC position represented in green. (E) Top: Cyan lines represent

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detected upstates. Bottom: LFP (black) and average upstate incidence per minute (cyan). Pink dotted line represents MDMA application time. (F) Fourier transformation and (G) instantaneous population activity for the recording shown in E. (H) Mean upstate incidence after saline (control), Fen or MDMA application (control: n = 5, Fen: n = 6, MDMA: n = 7; p < 10^{-4}, unpaired t test with Holm-Šidák correction). (I) Mean normalized spike rate after saline (control), Fen or MDMA application (control: n = 5, Fen: n = 6, MDMA: n = 7; p < 10^{-4}, unpaired t test with Holm-Šidák correction).

(Figure 1B). 5-HT does not cross the blood brain barrier (Hardebo and Owman, 1980), therefore, to understand the effect of 5-HT on SOs we used MDMA, a potent presynaptic 5-HT releaser and popular recreational drug (Green et al., 2003) that has shown promising results in the treatment of post-traumatic stress disorder (PTSD) (Inserra et al., 2021).

Intraperitoneal injections of MDMA (1.25 mg/kg) caused a strong suppression of upstate incidence (Figure 1E-H), a decrease in power of low frequencies (1-20 Hz) (Figure 1F), and a reduction of population spiking activity (Figure 1G-I). In addition to 5-HT, MDMA has been shown to cause the release of other neuromodulators, such as dopamine and noradrenaline (NE), although to a much lesser extent (Green et al., 2003). To test whether the effect of MDMA was mediated specifically by 5-HT, we repeated the experiment using Fen (5 mg/kg), a more selective 5-HT releaser (Rothman and Baumann, 2002, Heifets et al., 2019, Baumann et al., 2000). Intraperitoneal injection of Fen had a comparably strong suppressive effect on both the occurrence of upstates and population spiking activity (Figure 1H-I). Given the similarity of the observed effects, we grouped the results of Fen and MDMA application and found that both of these drugs significantly decreased LFP power in the delta and theta frequency ranges (Figure 1–figure supplement 2). Furthermore, the duration and area of upstates were also significantly reduced (Figure 1–figure supplement 3). These results point to a likely involvement of 5-HT in modulating ongoing oscillatory activity and suppressing low-frequency fluctuations.

**MDMA and Fen activate a subgroup of cortical neurons**

In addition to the LFP signal, we recorded the activity of 355 single units within the EC. Because of the similar effect of MDMA and Fen on spike rates (Figure 1I), we pooled all units recorded in both types of experiments. We found that drug injections differentially affected spiking rates (Figure 2A) of recorded units: while spiking decreased in most units (‘non-activated’), a small group of units (‘activated’) responded in the opposite fashion (n=31/355, 8.7%).

We then calculated the trough-to-peak (TP) latency of the spike waveforms, which has been
consistently used as a metric to classify units. In accordance with previous studies (Senzai et al., 2019, Roux et al., 2014), we found a clear bimodal distribution of TP latencies distinguishing putative excitatory (Exc) and fast-spiking inhibitory (Int) groups. Analysis of cross-correlograms confirmed the inhibitory nature of a subset of putative FS units (Figure 2–figure supplement 1) (Barthó et al., 2004). The cumulative distribution of TP latencies of ‘activated’ units was significantly different from both Exc and Int groups (Figure 2B). Specifically, the average TP latency of ‘activated’ units was situated in between the Int and Exc groups (Figure 2B-D, Figure 2–figure supplement 2), possibly suggesting a non-FS interneuron identity (Trainito et al., 2019, Kvitsiani et al., 2013). Notably the intermediate TP latency of “activated” units is not the result of an equal distribution between high and low values. Units with intermediate TP latency (between 0.4 and 0.6 ms) are unique in showing a 40% to 80% likelihood of being “activated” by drug application (Figure 2B).

Figure 2| Divergent unit responses to MDMA/Fen application. (A) Spike rate of the activated units versus all the other units during MDMA/Fen application (activated: n = 31, Non-activated: n = 324). (B) Top: TP latencies color-coded by group. Middle: cumulative distribution of TP latencies (Kolmogorov-Smirnov test, p Activated vs Int < 10^-4, p Activated vs Exc < 10^-4). Bottom: bar plot representing probability distribution of TP latencies, on the right y axis dashed line representing the percentage of ‘activated’ units per TP latency bin. (C) Distribution of units according to trough-to-peak (TP) latencies and repolarization time. Units were classified as putative interneurons (Int, blue) and putative excitatory neurons (Exc,
dark gray) according to a threshold at 0.55 ms; activated units (red) could belong to either
group but were mostly intermediate as shown by the covariance (2 STD) of each group
(Ellipses). Units recorded during control experiments are represented by empty circles. (D)
Waveforms of recorded units (n= 355). Units were divided into “putative excitatory” (black)
and “putative inhibitory” (blue) neurons according to TP latencies. Units activated by either
MDMA or Fen application are represented in red. Inset shows the average waveform for each
group. Scale bars: 0.5 ms.

5-HT suppresses SOs and activates Sst interneurons via 5-HT2AR in vitro
To understand the mechanism underlying the suppression of SOs by MDMA/Fen, we
performed in vitro experiments combining electrophysiology and pharmacology. First, we
recorded simultaneously from up to four neurons in the superficial layers of the EC (Figure
3A). Brain slices were perfused with an extracellular solution containing Mg2+ and Ca2+ in
concentrations similar to physiological conditions. With this method we could reliably detect
SOs reminiscent of the in vivo network activity (Tahvildari et al., 2012). Release of 5-HT in
vitro, induced by Fen application (200 µM), caused a suppression of SOs similar to what we
observed in vivo (Figure 3–figure supplement 1). Likewise, application of low concentrations
of 5-HT (5 µM) resulted in the suppression of SOs (Figure 3B, C). This effect was highly
consistent across different slices and was readily reversible (Figure 3D). Similarly to
spontaneous upstates, electrically evoked upstates (Neske et al., 2015b) were also
suppressed by 5-HT (Figure 3–figure supplement 2). Increasing the stimulation intensity did
not rescue upstate generation, indicating that a lack of excitation alone cannot explain the
suppressive effect of 5-HT on SO. In accordance with these results, we decided to focus
solely on 5-HT, however, we acknowledge that other neurotransmitters might be involved in
the effect of MDMA/Fen we observed in vivo.

Suppression of activity can have either an intrinsic or synaptic origin (Turrigiano, 2011). A
substantial subset of EC excitatory neurons is known to express 5-HT1A receptor (5-HT1A-R)
and hyperpolarize upon 5-HT application via activation of G protein-coupled inwardly
rectifying potassium (GIRK) channels (Schmitz et al., 1998, Chalmers and Watson, 1991). The
suppression of SOs by 5-HT, however, was not influenced by blocking 5-HT1A-R (WAY 100635,
100 nM) (Figure 3–figure supplement 3A, E). Blocking 5-HT3-R (tropisetron, 1 µM), the receptor
that characterizes one of the three main groups of interneurons, also did not have any impact
on SOs suppression induced by 5-HT. Similarly, application of the 5-HT3-R agonist m-CPBG
(50 µM) did not have any effect on SOs (Figure 3–figure supplement 4). In contrast, blocking
5-HT2A-R with the selective antagonist ketanserin (1 µM) (Preller et al., 2018) strongly reduced
the suppression power of 5-HT on SOs from 95 ± 4% to 57 ± 10.1% (Figure 4–figure
supplement 3B, E). The remaining suppression can be possibly explained by the activation of
5-HT$_{1A}$R on excitatory cells, as is reflected by the reduced spiking activity of putative excitatory cells (Figure 3–figure supplement 5).

Selective activation of 5-HT$_{2A}$R by α-methyl-5-HT (5 µM) mimicked the suppression of SOs observed after 5-HT wash-in (Figure 3–figure supplement 3C, E). Together, these results point to the importance of 5-HT$_{2A}$R in the suppression of SOs. 5-HT$_{2A}$R activation is known to cause an increase in intracellular calcium and consequent depolarization of the resting potential (RP) (Nichols and Nichols, 2008). Accordingly, after 5-HT application, we found that a small group of neurons was depolarized (n= 6/48, 12.5%) (Figure 3–figure supplement 6B–C). Using a soft

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**Figure 3** 5-HT suppresses SOs and activates Sst interneurons. (A) Biocytin staining of four simultaneously recorded cells shown in (B) WFS1 expression (in red) delimits L2/3 border. (B) Intracellular recordings showing synchronous upstate events in 4 simultaneously recorded cells before (left) and after (right) 5-HT application. Scale bars: 1: 7.5 mV, 2: 25 mV, 3: 25 mV, 4: 10 mV; 10 s. (C) Upstate raster plot before and after 5-HT application, orange box represents 5-HT application (n = 17, p < 10$^{-4}$, Wilcoxon signed rank test). (D) Representative recording showing the temporary inhibitory effect of 5-HT on SOs in two simultaneously recorded cells. Scale bars: 5 min, 20 mV. (E) PCA projection plot of all the cells recorded (n = 48). Cells are color-coded according to group identity: Exc (black), FS (light blue) or LTS (green). Typical voltage responses to current injection (-150 and +250 pA) are plotted for each group. Inset shows the average spike waveform for each group. (F) Representative recording of an excitatory (black) and a low-threshold (green) neuron simultaneously recorded during 5-HT application. Scale bars: 10 mV, 30 s. (G) Average change of RP before and after 5-HT application, across excitatory, fast-spiking and low-threshold neurons (Exc: n = 34, FS: n = 6; LTS: n = 9; p < 10$^{-4}$, unpaired t test with Holm–Šidák correction). (H) Biocytin staining of cells recorded in Sst-tdTomato mouse. Biocytin in green,
tdTomato in red. Scale bar: 50 µm. (I) Representative recording of a Sst interneuron during 5-HT application. Scale bars: 10 mV, 30 s. (L) Average RP of Sst interneurons during 5-HT (red) and ketanserin + 5-HT (blue) application, orange bar represents 5-HT (5-HT: n = 19, ketanserin + 5-HT: n = 22).

Clustering approach with six electrophysiological parameters (see “Materials and Methods”) we divided the recorded cells into 3 groups: Excitatory (Exc), fast-spiking (FS) and low-threshold spiking (LTS). Strikingly, the cells excited by 5-HT belonged exclusively to the LTS group (Figure 3G, Figure 3–figure supplement 6). A substantial proportion of LTS neurons express Sst (Tremblay et al., 2016, Gibson et al., 1999), therefore we performed targeted patch-clamp recordings using a mouse line expressing tdTomato specifically in Sst-expressing interneurons (Figure 3H). A subgroup of Sst interneurons are FS cells (Urban-Ciecko et al., 2018), only LTS Sst interneurons were considered in the following analysis. Sst interneurons depolarized upon 5-HT application (n=19, ΔRP: 7.5 ± 1.23 mV) (Figure 3I-L) and in some cases continued to spike while SOs were suppressed (n= 8/17, 47.05%, mean spiking rate = 3.03 ± 0.39 spikes/s). This effect was blocked by ketanserin (n = 22) (Figure 3L). We confirmed the presence of 5-HT2A R in Sst interneurons using immunohistochemistry in mice expressing EGFP under the 5-HT2A R promoter. In accordance with a previous study we found the majority of 5-HT2A R positive cells in the deep layers of EC, with a peak in L6 (Weber and Andrade, 2010). We observed that 11.8 ± 2.9 % of the 5-HT2A R positive cells in EC colocalized with Sst (n = 7 mice) (Figure 3–figure supplement 9). These results suggest that Sst interneurons may provide the synaptic inhibition required for the suppression of SOs.

**Sst interneurons mediate the suppression of SOs by 5-HT in vitro**

To evaluate the contribution of Sst interneurons to the 5-HT-mediated silencing of SOs we used an opto- and pharmacogenetic approach. First, we transgenically expressed channelrhodopsin-2 (ChR2) in Sst interneurons (Figure 4A). Light-stimulation of ChR2-expressing Sst interneurons in vitro suppressed SOs consistently (Figure 4D-F). Expectedly, upstate associated spiking was also diminished (Figure 4C, E, F). At the end of the light stimulation, spontaneous upstates immediately reoccurred (Figure 4G-H), in line with a critical role of Sst interneurons in the modulation of SOs (Fanselow et al., 2008, Funk et al., 2017, Niethard et al., 2018). We acknowledge that activation of PV interneurons can cause a similar suppression (Zucca et al., 2017). While this experiment establishes the ability of Sst interneurons to suppress SOs, it does not causally link Sst interneuron activation to the suppression of SOs induced by 5-HT. Therefore, we generated a transgenic mouse line
Figure 4 | Sst interneurons activation suppresses SO. (A) Experimental protocol: Sst interneurons expressing ChR2 are activated by light during intracellular recording of L3 neurons in EC. (B) Representative recordings from a L3 neuron during Sst interneuron activation. Scale bars: 10 mV, 0.5 s. (C) Spikes raster (top) and density plot (bottom) during light stimulation. (D) Upstate raster (top) and density plot (bottom) during light stimulation. (E) Left: spike frequency during baseline light stimulation (n=14; p < 0.001, Wilcoxon signed rank test). Right: upstate incidence during baseline and light stimulation (n=14; p < 0.001, Wilcoxon signed rank test). Patches represent 95% confidence interval, lines represent standard deviation. (F) Left: spike probability polar plot during Sst interneurons light activation. Right: upstate probability polar plot during Sst interneurons light activation. Note the absence of both spiking activity and upstates during Sst interneurons activation.

carrying a Cre-conditional expression cassette of the pharmacogenetic silencer hM4Di (Figure 4—figure supplement 1) (Armbruster et al., 2007). Homozygous Cre-conditional hM4Di transgenic mice and Sst-Cre mice were bred to obtain heterozygous Sst-Cre/hM4Di offspring, which allow specific inhibition of Sst interneuron activity using Clozapine-N-Oxide (CNO). Following application of 5-HT we observed a strong reduction of upstate incidence, which was partially restored by subsequent application of CNO (Figure 5 A-B).
Activation of 5-HT$_{1A}$R on excitatory cells by 5-HT and the resulting decreased network excitation drive might account for the fact that the incidence of upstates did not completely return to the baseline level upon CNO wash-in. Additionally, in further experiments in which CNO was applied before 5-HT, the significant reduction in upstate incidence typically seen after 5-HT wash-in was not observed (Figure 5–figure supplement 1). CNO did not show any significant effect in both wild-type littermates and hM4Di-PV mice (Figure 5–figure supplement 2), indicating that the results observed are due to its specific effect on the activity of Sst interneurons. In summary, while activation of Sst interneurons either via 5-HT or directly by ChR2 suppresses SOs, the pharmacogenetic inactivation of Sst interneurons weakens the effect of 5-HT on SOs.

**Discussion**

In this study we show that the substitute amphetamines MDMA and Fen suppress default cortical network oscillations *in vivo* in the mEC. Furthermore, using an opto- and pharmacogenetic approach *in vitro*, we demonstrate that Sst interneurons, activated by 5-HT$_{2A}$R, contribute to this suppression.

Organization of cortical activity is brain state-dependent, ranging continuously from “synchronized” to “desynchronized” states (Harris and Thiele, 2011). SOs are on one end of
this continuum, representing the prototypical synchronized state. Our results, in line with previous studies (Puig et al., 2010, Grandjean et al., 2019), show that 5-HT can suppress synchronized cortical activity; in addition, we identify Sst interneurons as contributing to this suppression.

Sst interneurons have been previously proposed to provide the inhibition necessary for the termination of upstates, due to their strongly facilitating synapses (Krishnamurthy et al., 2012, Melamed et al., 2008). Additionally, increased inhibition, as shown both in computational models and experimental data, can counteract temporal summation of inputs and reduce correlation due to tracking of shared inputs between inhibitory and excitatory populations (Sippy and Yuste, 2013, Renart et al., 2010, Stringer et al., 2016). In the case of Sst interneurons, this would especially impact summation in distal compartments of pyramidal cells, due to the axonic targeting of Sst cells onto pyramids. Sst interneurons are specifically known to be sufficient to cause desynchronization in V1 (Chen et al., 2015). While it is well known that Sst interneurons are potently excited by ACh in various cortical areas (Chen et al., 2015, Obermayer et al., 2018, Fanselow et al., 2008), including mEC (Desikan et al., 2018), and NE in frontal cortex (Kawaguchi and Kubota, 1998), our work identifies them as a novel target of 5-HT regulation via 5-HT$_{2A}$R. Recently it has been proposed that inhibitory interneurons play a key role in mediating the effect of ACh and NE on cortical state transitions (Cardin, 2019). Our results add a new level of complexity to this picture.

The excitation of Sst interneurons by 5-HT possibly contributes to the net inhibitory effect of 5-HT release observed in many cortical areas (Grandjean et al., 2019, Seillier et al., 2017, Azimi et al., 2020), and could explain why the inhibition strength is linearly correlated to 5-HT$_{2A}$R expression (Grandjean et al., 2019). Giving further support to this idea, Sst interneurons in the somatosensory cortex show increased cFos levels following 5-HT$_{2A}$R activation (Martin and Nichols, 2016). Previous studies have reported either 5-HT$_{2A}$R dependent inhibition or 5-HT$_{2A}$R dependent activation of interneurons in the prefrontal cortex (PFC) (Abi-Saab et al., 1999, Ashby et al., 1990, Athilingam et al., 2017), piriform cortex (Marek and Aghajanian, 1994, Sheldon and Aghajanian, 1990), cingulate cortex (Zhou and Hablitz, 1999), cochlear nucleus (Tang and Trussell, 2017), amygdala (Sengupta et al., 2017), olfactory bulb (Petzold et al., 2009, Hardy et al., 2005), visual cortex (Michaieil et al., 2019, Azimi et al., 2020) and hippocampus (Wyskiel and Andrade, 2016). However, none of these studies identified interneurons using molecular markers, so we do not exclude that different interneuron classes in other cortical areas might mediate the inhibitory downstream effects of 5-HT$_{2A}$R. For example, in the PFC a subgroup of PV interneurons has been reported to be activated by this
receptor (Athilingam et al., 2017, Puig et al., 2010). Furthermore, we know that PV neurons can reliably induce up to down state transitions (Zucca et al., 2017). 5-HT modulation is also involved in gain regulation. In the olfactory cortex, 5-HT has a selective subtractive effect on stimulus evoked firing (Lottem et al., 2016), and a recent study has shown in the visual cortex that the reduced gain of evoked responses is dependent on 5-HT\textsubscript{2A}R activation (Azimi et al., 2020). Intriguingly, Sst interneurons have been shown to regulate subtractive inhibition (Sturgill and Isaacson, 2015, Wilson et al., 2012).

5-HT levels in the brain, similarly to other neuromodulators, are synchronized to the sleep wake cycle, with higher levels present during the waking state (Oikonomou et al., 2019). How does this notion relate to activation of Sst interneurons by 5-HT\textsubscript{2A}R? A limitation of our study is the absence of data in naturalistic conditions, with anesthesia potentially being a confounding factor (Adesnik et al., 2012). We can nonetheless speculate that Sst interneurons should be more active during states with higher 5-HT levels (wake>SWS>REM). A previous study directly measuring the activity of various neuronal classes across different states seems to support this idea. Sst interneurons in the dorsal cortical surface display their highest activity during waking states, lower activity during SWS and lowest activity during REM (Niethard et al., 2016). Following pharmacological release of 5-HT in vivo we observed increased spiking activity in a subgroup of neurons whose waveform features are compatible with Sst interneurons. Our in vitro data using 5-HT at a concentration commonly used in the field (Gorinski et al., 2019, Wang et al., 2016, Huang et al., 2009) shows that Sst interneurons can be activated by 5-HT\textsubscript{2A}R, however, recordings during natural sleep and wake conditions in combination with optotagging will be needed to demonstrate conclusively that Sst interneurons are activated by 5-HT in physiological conditions in vivo. Potential differences between physiological and pharmacological 5-HT release are especially of interest considering the reported efficacy of MDMA in the treatment of PTSD and its recently approved status as breakthrough therapy (Inserra et al., 2021, Mithoefer et al., 2019).

While our data relative to MDMA application supports a role of 5-HT in modulating SOs in vivo we cannot exclude that other neuromodulators might play a role. For example, it is known that dopamine can suppress SOs in vitro (Mayne et al., 2013). Fen, however, has been reported to selectively increase 5-HT concentration in the brain (Rothman and Baumann, 2002) and has been used previously to specifically disentangle the effect of MDMA on different neuromodulatory systems (Heifets et al., 2019). This, together with our in vitro results, suggests that 5-HT is most likely mediating the effect induced by MDMA/Fen.
Besides its involvement in various physiological brain processes, 5-HT is also associated with the etiology of various psychiatric disorders, as are Sst interneurons (Pantazopoulos et al., 2017, Lin and Sibille, 2015). Furthermore, 5-HT is linked to the psychological effect of many psychotropic drugs; specifically, 5-HT_2A R activation has been reported to be essential for the psychological effects induced by various psychedelics (Nichols, 2016), and in the case of MDMA, has been linked to perceptual and emotional alterations (Liechti et al., 2000, Kuypers et al., 2018). Broadband reduction in oscillatory power, triggered by 5-HT_2A R, seems to be linked to the subjective effect of serotonergic drugs (Carhart-Harris et al., 2016, Carhart-Harris and Friston, 2019) and has been consistently observed in humans and rodents following administration of MDMA (Frei et al., 2001, Lansbergen et al., 2011) or various other 5-HT_2A R agonists (Kometer et al., 2015, Muthukumaraswamy et al., 2013, Carhart-Harris et al., 2016, Wood et al., 2012). The link between 5-HT_2A R and perception is further supported by the fact that several routinely used antipsychotic drugs are potent 5-HT_2A R antagonists (Marek et al., 2003, Meltzer, 1999). While the most recent attempts to explain the psychological effects of 5-HT_2A R activation focus on the increased spiking of cortical pyramidal neurons in the deep layers (Carhart-Harris and Friston, 2019, Nichols, 2016), our study suggests that Sst interneurons may also play a role. Sst interneurons, in contrast to PV interneurons, form synapses on the dendrites of their target cell (Tremblay et al., 2016). A wealth of evidence suggests that active dendritic processing in cortical pyramidal neurons has a critical influence on sensory perception (Takahashi et al., 2016, Murayama et al., 2009, Smith et al., 2013, Ranganathan et al., 2018) and, in accordance with their unique anatomical properties, Sst interneurons strongly influence dendritic computations and directly modulate perceptual thresholds (Takahashi et al., 2016).

We propose that the novel link between 5-HT_2A R and Sst interneurons might help elucidate the mechanisms underlying a host of psychiatric disorders and contribute to our understanding of how serotonergic drugs exert their psychological effects.
Materials and Methods

All experiments were conducted according to regulations of the Landesamt für Gesundheit und Soziales (Berlin [T 0100/03], Berlin [G0298/18]) and the European legislation (European Directive 2010/63/EU).

Animals

Data for the in vivo part of the study was collected from C57Bl/6 mice (aged 6 to 10 weeks). Data for the in vitro part was collected from C57Bl/6 (P10-P17), Sst-tdTomato (P10-P30), Sst-Chr2-EYFP (P10-P16), hM4Di-Sst (P10-P15), hM4Di-Sst (+/-) (P10-P15) and hM4Di-PV (P10-P15) mice. Immunostainings to localize 5-HT2aR were performed on 5-HT2aR-EGFP mice (P20-P90) and immunostainings to localize 5-HT fibers were performed on an ePet-YFP mouse (P35). Sst-Cre mice (RRID:IMSR_JAX:013044) have Cre recombinase targeted to the Sst locus; they were obtained from Jackson Laboratory (ME, USA). PV-Cre mice (RRID:IMSR_JAX:008069) have Cre recombinase targeted to the Pvalb locus; they were obtained from Charles River Laboratory (MA, USA). tdTomato mice (Ai9, RRID:IMSR_JAX:013044) were obtained from Jackson Laboratory. Chr2-EYFP mice (Ai32, RRID:IMSR_JAX:012569) were obtained from Jackson Laboratory. ePet-Cre mice (RRID:IMSR_JAX:012712) have Cre recombinase targeted to the Fev locus; they were obtained from Jackson Laboratory. 5-HT2aR-EGFP mice (RRID:MMRRC_010915-UCD) express EGFP reporter protein under the control of the Htr2a gene, they were obtained from the Mutant Mouse Resource & Research Centers (MMRRC, CA, USA). Generation of hM4Di mice is described in the paragraph “Generation of Cre-conditional hM4Di mice”. The animals were housed in a 12:12 hour light-dark cycle in singularly ventilated cages with ad libitum access to food and water. SOs in vitro recordings were performed on P12-P16 mice.

Generation of Cre-conditional hM4Di mice

We produced a transgenic mouse line carrying a Cre-conditional hM4Di expression cassette in the Rosa26 locus. The transgene construct was inserted by recombination-mediated cassette exchange (RMCE). RMCE relies on recombination events between attB and attP recognition sites of the RMCE plasmid and genetically modified acceptor embryonic stem (ES) cells, mediated by the integrase of phage phiC31 (Hitz et al., 2007). The RMCE construct is thereby shuttled into the Rosa26 locus of the ES cells, along with a Neomycin resistance cassette (fig. S12A). The acceptor cell line IDG3.2-R26.10-3 (I3) was kindly provided by Ralf...
Kühn (GSF National Research Centre for Environment and Health, Institute of Developmental Genetics, Neuherberg, Germany).

We subcloned a Cre-conditional FLEX (flip-excision) cassette (Schnutgen et al., 2003) into pRMCE, and inserted a strong CAG promoter (CMV immediate early enhancer/modified chicken β-actin promoter, from Addgene Plasmid #1378) in front of the FLEX-cassette to create pRMCE-CAG-Flex. The coding sequence of hM4Di-mKateT was inserted into the FLEX cassette in reverse orientation to the promoter (fig. S12A). Finally, a rabbit globulin polyA cassette including stop codons in every reading frame was placed downstream of the FLEX cassette, in the same direction as hM4Di, in order to prevent unintended transcriptional read-through from potential endogenous promoters. The construct was completely sequenced before ES cell electroporation.

Electroporation of the RMCE construct together with a plasmid encoding C31int was performed by the transgene facility of the ‘Research Institute for Experimental Medicine’ (FEM, Charité, Berlin) according to published protocols (Hitz et al., 2009, Hitz et al., 2007). Recombinant clones were selected by incubation with 140 µg/ml G418 for at least 7 days. To activate hM4Di expression by recombination of the FLEX switch, selected clones were further transfected transiently with pCAG-Cre-EGFP using Roti-Fect (Carl Roth, Karlsruhe, Germany). G418-resistant clones were analyzed by PCR for successful integration and recombination of the construct (fig. S12B), using the following primers:

- **GT001** PGK3’-fw: CACGCTTCAAAAGCGCAGTCTG;
- **GT002** Neo5’-rev: GTTGTGCCAGTCATAGCCGAATAG;
- **GT005** PolyA-fw: TTCCTCCTCTCTGACTACTCC;
- **GT006** Rosa3’-rev: TAAGCCTGCCAGAAGACTC;
- **GT013** hM4Di3’rec-rev: CAGATACTGCCACCTCCCTA

After verification of correct integration and functional FLEX-switch recombination, we generated chimeras by blastocyst injection of I3 ES cells. Heterozygous offsprings were mated with a Flpe deleter mouse line in order to remove the neomycin resistance cassette by Flp-mediated recombination.

Mice homozygous for the Rosa-CAG-FLEX-hM4Di-mKateT allele are viable and fertile and show no obvious phenotype. Importantly, application of CNO to these mice does not induce any behavioral effects. Homozygous Cre-conditional hM4Di transgenic mice and Sst-Cre mice (Taniguchi et al., 2011) were maintained on a C57Bl/6 genetic background and were bred to obtain heterozygous Sst-Cre / hM4Di offsprings.

Drugs
Urethane (U2500, Merck), Fenfluramine (+(+)-Fenfluramine hydrochloride, F112, Merck), 5-HT (Serotonin creatinine sulfate monohydrate, H7752, Merck), m-CPBG (1-(3-Chlorophenyl)biguanide hydrochloride, C144, Merck), tropisetron (Tropisetron hydrochloride, Y0000616, Sigma), WAY-100635 (W108, Merck), α-Methylserotonin (α-Methylserotonin maleate salt, M110, Merck) MDMA ((±)3,4-methylenedioxymethamphetamine, 64057-70-1, Merck), CNO (Clozapine N-oxide dihydrochloride, 6329, Tocris) were dissolved in water for in vitro application and in 0.9% normal saline for in vivo application. Ketanserin (Ketanserin (+)-tartrate salt, S006, Merck) was dissolved in Dimethyl sulfoxide (DMSO).

**Surgery and in vivo recording**

Before recordings mice were briefly anaesthetized with isoflurane (2%) and then injected intraperitoneally with urethane (1.2 g/kg, Sigma Aldrich, Munich, Germany). The level of anesthesia was maintained so that hindlimb pinching produced no reflex movement and supplemental doses of urethane (0.2 g/kg) were delivered as needed. Upon cessation of reflexes the animals were mounted on a stereotaxic frame (Kopf Instruments, Tujunga, California), and body temperature was maintained at 38°C. The scalp was removed, and the skull was cleaned with saline solution. A craniotomy was performed at +3 mm ML, -3 mm AP, +3.25 mm DV to reach mEC.

Extracellular recordings from EC (Figure 1-2) were performed using a Cambridge Neurotech (Cambridge, United Kingdom) silicon probe (64-channels (n = 15) or 32-channels (n = 3). The recording electrode was painted with the fluorescent dye Dil (Thermo Fisher Scientific, Schwerte, Germany) and then slowly lowered into the craniotomy using micromanipulators (Luigs&Neumann, Ratingen, Germany) at a 25° angle AP (toward the posterior side of the brain). The exposed brain was kept moist using saline solution. A ground wire connected to the amplifier was placed in the saline solution covering the skull to eliminate noise. Brain signals were recorded using a RHD2000 data acquisition system (Intan Technologies, Los Angeles, California) and sampled at 20kHz. Recording quality was inspected on-line using the open-source RHD2000 Interface Software. A supplementary dose of urethane (0.2 g/kg) was injected right before the start of the recording to standardize anesthesia level across different experiments and avoid the arise of theta activity in the first 30 minutes of recording.

Recordings began after a 10-minute waiting period in which clear upstates could consistently be seen at a regular frequency.
**In vivo analysis**

We selected the channel to use for upstate detection based on the standard deviation (STD) of the trace during baseline (first 5 minutes of recording): the channel with the highest STD was selected, as larger voltage deflection increases detection algorithm accuracy. Given the highly synchronous nature of SOs (Supplementary Fig. 3) the spatial location of the channel selected was not considered. Upstates were detected comparing threshold crossing points in two signals: the delta-band filtered signal (0.5-4 Hz) and the population spike activity. Candidate upstates were identified in the delta-band filtered signal using two dynamic thresholds ‘a’ and ‘b’:

\[
\begin{align*}
a &= m + \frac{\sigma}{1.5} \\
b &= m + \frac{\sigma}{0.8}
\end{align*}
\]

Where \(\sigma\) is the standard deviation of the signal during the first five minutes of recording (baseline) and \(m\) is the centered moving median calculated using 60 s windows (Matlab function `movmedian`). The median was used instead of the mean to account for non-stationarities in the data. A candidate upstate was identified at first using the threshold crossings of the signal compared to ‘a’: candidates shorter than 200 ms were deleted and multiple candidates occurring within a window of 300 ms were joined together. Subsequently the threshold ‘b’ was used to separate upstates occurring in close proximity: if the signal within one candidate crossed the threshold ‘b’ in more than one point then the candidate upstate was split in two at the midpoint between the two threshold crossings. Candidate upstates were finally confirmed if the population spike activity (calculated in 100 ms windows) within the candidate crossed a threshold of 1 \(\sigma\) (calculated during the baseline).

**Units detection and classification**

Spike detection was performed offline using the template-based algorithm Kilosort2 ([https://github.com/MouseLand/Kilosort2](https://github.com/MouseLand/Kilosort2)), with the following parameters:

- \(\text{ops.fshigh} = 300\)
- \(\text{ops.fsslow} = 8000\)
- \(\text{ops.minfr_goodchannels} = 0\)
- \(\text{ops.Th} = [8 4]\)
- \(\text{ops.lam} = 10\)
- ops.AUCsplit = 0.9
- ops.minFR = 1/1000
- ops.momentum = [20 400]
- ops.sigmaMask = 30
- ops.ThPre = 8
- ops.spkTh = -6
- ops.nfilt_factor = 8
- ops.loc_range = [3 1]
- ops.criterionNoiseChannel = 0.2
- ops.whiteningrange = 32
- ops.ntbuff = 64

Spike sorting in recordings performed with Neuropixel probes was performed using ecephys_spike_sorting ([https://github.com/AllenInstitute/ecephys_spike_sorting](https://github.com/AllenInstitute/ecephys_spike_sorting)). Manual curation of the results was performed using Phy ([https://github.com/cortex-lab/phy](https://github.com/cortex-lab/phy)). Each isolated unit satisfied the following two criteria: Refractory period (2 ms) violations < 5%, fraction of spikes below detection threshold (as estimated by a gaussian fit to the distribution of the spike amplitudes) < 15%. Units with negative maximal waveform amplitude were further classified as putative excitatory if the latency (TP latency) was > 0.55 ms or putative inhibitory when TP latency < 0.55 ms. The value 0.55 was chosen in accordance with previous studies (Senzai et al., 2019, Antoine et al., 2019). In pharmacological classification, units were classified as 'activated' if their firing rate in the 25 minutes following drug injection was 2 σ (standard deviation) above the baseline rate for at least 5 minutes. Remaining units were pulled together in the category 'non-activated'.

**Cross-correlogram analysis**

Cross-correlogram based connectivity analysis was performed for every unit to identify inhibitory connections. Units with a spiking rate smaller than 0.3 spikes/s were discarded from the analysis. We used total spiking probability edges (TPSE) algorithm ([https://github.com/biomemsLAB/TSPE](https://github.com/biomemsLAB/TSPE)) (De Blasi et al., 2019) to identify in a computationally efficient manner putative inhibitory connections between units and all clusters recorded. The parameters used were:
\[
\begin{align*}
  & d = 0, \\
  & \text{neg}_\text{wins} = [2, 3, 4, 5, 6, 7, 8], \\
  & \text{co}_\text{wins} = 0, \\
  & \text{pos}_\text{wins} = [2, 3, 4, 5, 6], \\
  & \text{FLAG}_\text{NORM} = 1.
\end{align*}
\]

The connectivity vectors of each unit resulting from TSPE were sorted by inhibition strength. The top 20 connections were further analyzed using custom Matlab (RRID:SCR_001622) code. A connection was classified as inhibitory if the cross correlogram values \((x)\) were smaller than the mean of \(x\) by more than one standard deviation \((x < \text{mean}(x) - \text{std}(x))\) in at least 4 consecutive bins (bin size = 1 ms) in a window 4 to 9 ms after the center of the cross-correlogram.

**Slice preparation**

We prepared acute near horizontal slices (~15° off the horizontal plane) of the medial entorhinal cortex (mEC) from C57Bl/6 mice. Animals were decapitated following isoflurane anesthesia. The brains were quickly removed and placed in ice-cold (~4°C) ACSF (pH 7.4) containing (in mM) 85 NaCl, 25 NaHCO3, 75 Sucrose, 10 Glucose, 2.5 KCl, 1.25 NaH2PO4, 3.5MgSO4, 0.5 CaCl2, and aerated with 95% O2, 5% CO2. Tissue blocks containing the brain region of interest were mounted on a vibratome (Leica VT 1200, Leica Microsystems), cut at 400 μm thickness, and incubated at 35 °C for 30 min. The slices were then transferred to ACSF containing (in mM) 85 NaCl, 25 NaHCO3, 75 Sucrose, 10 Glucose, 2.5 KCl, 1.25 NaH2PO4, 3.5 MgSO4, 0.5 CaCl2. The slices were stored at room temperature in a submerged chamber for 1-5 hr before being transferred to the recording chamber.

**In vitro recording**

In order to perform whole-cell recordings slices were transferred to a submersion style recording chamber located on the stage of an upright, fixed-stage microscope (BX51WI, Olympus) equipped with a water immersion objective (x60, Olympus) and a near-infrared charge-coupled device (CCD) camera. The slices were perfused with ACSF (~35 °C bubbled with 95 % O2-5 % CO2) at 3-5 ml/ min to maintain neuronal health throughout the slice. The ACSF had the same composition as the incubation solution except for the concentrations of calcium and magnesium, which were reduced to 1.2 and 1.0 mM, respectively. Recording
electrodes with impedance of 3-5 MΩ were pulled from borosilicate glass capillaries (Harvard Apparatus, Kent, UK; 1.5 mm OD) using a micropipette electrode puller (DMZ Universal Puller). The intracellular solution contained the following (in mM): 135 K-gluconate, 6 KCl, 2 MgCl₂, 0.2 EGTA, 5 Na₂-phosphocreatine, 2 Na₂-ATP, 0.5 Na₂-GTP, 10 HEPES buffer, and 0.2% biocytin. The pH was adjusted to 7.2 with KOH. Recordings were performed using Multiclamp 700A/B amplifiers (Molecular Devices, San Jose, California). The seal resistance was >1 GΩ. Capacitance compensation was maximal and bridge balance adjusted. Access resistance was constantly monitored. Signals were filtered at 6 kHz, sampled at 20 kHz, and digitized using the Digidata 1550 and pClamp 10 (Molecular Devices, San Jose, California). Activation light was delivered by a 460 nm laser (DPSS lasers, Santa Clara, California) using a 460–480 nm bandpass excitation filter. Stimulation consisted of 500 ms pulses at 1 Hz.

Stimulation experiments were performed using a bipolar micro-electrode (glass pipette filled with ACSF solution, wrapped by a fine grounding wire) connected to an isolated voltage stimulator (ISO-Flex, A.M.P.I., Israel). A 4x objective (Olympus) was used to visually guide the stimulating electrode into the mEC. Stimulation power was adjusted to achieve consistent upstate generation during baseline (> 95%). Each stimulus had a duration of 50 μs and the inter-stimulus interval was 8-10 seconds.

**In vitro analysis**

*In vitro* upstates were detected in Matlab using an algorithm similar to the one described in the *in vivo* analysis method section. We used a coincident detection in two signals. In multicellular recordings we used the membrane potential of 2 cells, in single cell recordings we used membrane potential and the envelope of the gamma filtered trace (50-250 Hz), as upstates are characterized by an increase in gamma activity (Neske, 2015).

The baseline condition was defined as the last 120 s before drug application, while the post-drug application condition was defined as the 120 s of recording after drug application (Total recording duration: 600 s).

Excitatory (Exc), fast-spiking (FS) and low-threshold spiking (LTS) neurons were classified using Gaussian mixture models (GMM) with a soft clustering approach in Matlab. Input resistance (Rᵢ), Δafter-hyperpolarization (ΔAHP), sag, rheobase, spike width and resting potential (RP) were extracted from each neuron and used in the classification. The first two components of the principal component analysis (PCA) were used to fit the data to a Gaussian mixture model distribution. Initial values were set according to the k-means algorithm with
centroid positioned at x and y position: 5, 0; -15, -15; -15, 10. This centroid were placed according to the loadings of the PCA to identify 3 clusters with the following main features:

- Cluster 1 (putative Exc): high spike width, low AHP, low rheobase.
- Cluster 2 (putative FS): low spike width, low SAG, high rheobase, low R_{in}.
- Cluster 3 (putative LTS): low spike width, high SAG, high AHP, high R_{in}.

Covariance matrices were diagonal and not shared. Neurons with a posterior probability of belonging to any of the three clusters of < 90% were discarded from further analysis (1/49).

While the majority of Sst-interneurons display LTS features, a minority (~10%) belong to the FS group (Urban-Ciecko et al., 2015). To distinguish FS and LTS interneurons in the Sst-Td Tomato mice, we employed the GMM with posterior probability threshold of 90%. Only LTS Sst neurons were considered for further analysis.

### Histological analysis

For the post-mortem electrode track reconstructions of the in vivo recordings, mice were not perfused; rather, brains were extracted from the skull, post-fixed in 4% PFA overnight at 4°C and afterwards cut with a vibratome (Leica Microsystems, Wetzlar Germany) in 100 μM thick sequential sagittal slices. Images were taken using a 1.25x objective and stitched together using the microscope software (BX61, Olympus). Afterwards we used AllenCCF code (https://github.com/cortex-lab/allenCCF) to identify electrode shank location (Shamash et al., 2018).

For the anatomical reconstructions of recorded cells in vitro, brain slices were fixed with 4% paraformaldehyde in 0.1 M phosphate buffer (PB) for at least 24 hours at 4°C. After being washed three times in 0.1 M PBS, slices were then incubated in PBS containing 1% Triton X-100 and 5% normal goat serum for 4 hr at room temperature (RT). To visualize biocytin-filled cells we used Streptavidin Alexa 488 conjugate (1:500, Invitrogen Corporation, Carlsbad, CA, RRID:AB_2315383). WFS1 (1:1000, Rabbit, Proteintech, IL, USA, RRID:AB_2880717) was used in a subset of experiments to visualize the L2/L3 border, and Sst (1:1000, Rat, Bachem, Switzerland, RRID:AB_2890072) was used in the 5-HT_{3A}R localization analysis. Slices were incubated with primary antibodies for 48 hours at RT. After rinsing two times in PBS, sections were incubated in the PBS solution containing 0.5% Triton X-100, Alexa fluor 488, Alexa fluor 555 and Alexa fluor 647 (Invitrogen Corporation, Carlsbad, CA) according to the number of
antibodies used. Slices were mounted in Fluoroshield (Sigma-Aldrich) under coverslips 2-3 hr after incubation with the secondary antibodies and stored at 4 °C.

Labeled cells were visualized using 20x or 40x objectives on a confocal microscope system (SP8, Leica, RRID:SCR_018169). For the 5-HT₂AR localization analysis, images of the whole EC were acquired and stitched together using the auto stitching method, without smoothing. Z stacks were acquired every 30 µM. The image stacks obtained were registered and combined in Fiji (RRID:SCR_002285) to form a montage of the sections. Cell counting was executed using the Fiji multi-point tool. X-Y-Z coordinates of each 5-HT₂AR-EGFP positive cell were exported to Matlab and subsequently, using custom written code in Matlab, we semi-automatically inspected each cell for colocalization between EGFP(5-HT₂AR) and Sst.

**Statistical Analysis**

All datasets were tested to determine normality of the distribution either using D'Agostino-Pearson omnibus normality test or Shapiro-Wilk normality test. Student’s t-test and one-way ANOVA were used for testing mean differences in normally distributed data. Wilcoxon matched-pairs signed rank test and Kruskal-Wallis were used for non-normally distributed datasets. Dunn-Sidak multiple comparison test was used to compare datasets with 3 or more groups. Kolmogorov-Smirnov test was used to compare cumulative distributions. Statistical analysis was performed using Prism (6.01, RRID:SCR_002798) and Matlab. All data are expressed as mean ± SEM. Asterisks in figures represent p-values smaller than 0.05, unless stated otherwise in the legend.
**Supplementary Figures**

![Supplementary Figures](image)

**Figure 1—figure supplement 1. In vivo upstate spatial features.**

(A) Microelectrode implant location and microelectrode features. 64 channels ($n_{\text{animals}} = 15, n_{\text{shanks}} = 4$) and 32 channels ($n_{\text{animals}} = 3, n_{\text{shanks}} = 2$) microelectrodes were used in this study, analysis shown in this figure excludes data recorded with 32 channels probe due to the different spatial configuration of the channels. (B) Average upstate voltage deflection for each channel of the microelectrode. For experiments with drug application (either MDMA or Fen), only baseline upstates were taken in account. (C) Left: Average upstate voltage deflection grouped by shank. Right: Average upstate voltage deflection grouped by depths (right). Insets show the normalized correlation between averages in the two different groups.
Figure 1–figure supplement 2. LFP power analysis for saline and MDMA/Fen injection.

(A) Average normalized LFP power during baseline and saline injection. (B) Average LFP power during baseline and MDMA/Fen injection. (C) Box plots of normalized LFP power per frequency band: delta (Control: 0.98 ± 0.01, MDMA/Fen: 0.82 ± 0.04, p = 0.019, Mann-Whitney test), theta (Control: 0.90 ± 0.01 µV, MDMA/Fen: 0.76 ± 0.04, p = 0.046, Mann-Whitney test), alpha (Control: 0.77 ± 0.01, MDMA/ Fen: 0.45 ± 0.03, p > 0.05, Mann-Whitney test), beta(Control: 0.49 ± 0.02, MDMA/ Fen: 0.30 ± 0.03, p > 0.05, Mann-Whitney test), and gamma (Control: 0.28 ± 0.02, MDMA/ Fen: 0.25 ± 0.02, p > 0.5, Mann-Whitney test).
**Figure 1—figure supplement 3. In vivo upstate metrics for saline and MDMA/Fen injection.**

**(A)** Average upstate voltage deflection for control and MDMA/Fen, patches represent 1 standard deviation. Scale bars: 0.5 seconds, 200 µV. **(B)** Violin plots of duration (control: 0.98 ± 0.02 s, MDMA/Fen: 0.69 ± 0.04 s, p < 0.001, Mann-Whitney test). **(C)** Violin plots of peak 1 amplitude (control: 434.3 ± 13.09 µV, MDMA/Fen: 248.1 ± 10.49 µV, p < 0.001, Mann-Whitney test). **(D)** Violin plots of peak 2 amplitude (control: 232.4 ± 3.53 µV, MDMA/Fen: 187.3 ± 7.40 µV, p < 0.001, Mann-Whitney test). **(E)** Violin plots of area (control: 185.6 ± 4.14 µV•s, MDMA/Fen: 187.3 ± 7.40 µV, p < 0.001, Mann-Whitney test). **(F)** Violin plots of inter-event intervals (control: 5.55 ± 0.26 s, MDMA/Fen: 12.02 ± 1.75 s, p < 0.001, Mann-Whitney test).
Figure 2–figure supplement 1. Cross-correlogram (CCG) based connectivity analysis.

(A) Units are plotted according to TP latencies and repolarization time and color-coded according to the number of inhibitory connections detected. Units displaying a TP latency < 0.55 ms were classified as putative inhibitory interneurons (‘Putative int’), the remaining units were classified as putative excitatory neurons (‘Putative exc’). Inhibitory connections were detected using Total Spiking Probability Edges (TSPE) (See Supplementary materials). Putative interneurons had a 40.0 % chance of displaying at least one inhibitory connection in the CCGs with an average number of 3.38 ± 0.68 inhibitory connections while putative excitatory cells had a 0.33 % chance of displaying inhibitory connections. Units with a spiking...
rate lower than 0.3 spikes/s were discarded from the analysis. Units from control experiments are included in the analysis. **(B)** Connectivity scheme of one putative inhibitory unit (source unit, black circle) displaying 10 inhibitory connections with surrounding clusters. Grey rhombi represent recording channels on the probe with the tip of the shanks pointing north. Each colored circle represents the location of an inhibited unit. Waveforms of the inhibited units are plotted nearby the location using the same color. Inset shows the location of the source unit on the probe. **(C)** CCGs of the connections displayed in (B) using the same color code. Solid lines represent mean, dashed lines represent 1 standard deviation.
Figure 2—figure supplement 2. TP latency density distributions.

(A) Dashed lines represent kernel density estimations of probability density functions of the TP latencies of putative inhibitory (blue), putative excitatory (black) and ‘activated’ units. Solid lines represent gaussian fitting curves for each group. Peak inhibitory gaussian: 0.31 ms, peak excitatory gaussian: 0.81 ms and peak ‘activated’ gaussian: 0.56 ms. (B) Goodness of fit metrics for each gaussian fit. Sse = Sum of squares due to error, rsquare = R-squared (coefficient of determination), dfe = Degrees of freedom in the error, adjrsquare = Degree-of-freedom adjusted coefficient of determination.

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<th>Activated</th>
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**Figure 3–figure supplement 1. Effect of Fen on SOs in vitro.**

(A) Top: upstate raster plot during Fen application. Bottom: Histogram of upstate incidence before and after Fen application (n= 6, baseline: 1.13 ± 0.17 upstates/10s, Fen: 0.04 ± 0.03 upstates/10s).

(B) Top: upstate raster plot during ketanserin + Fen application. Bottom: Histogram of upstate incidence before and after ketanserin + Fen application (n= 5, baseline: 1.18 ± 0.07 upstates/10s, Fen: 1.03 ± 0.17 upstates/10s).

(C) Upstate metrics during baseline and ketanserin + Fen. Left: violin plots of inter-event interval (baseline = 9.46 ± 0.28 s, ketanserin + Fen = 9.63 ± 0.74 s). Middle: violin plots of inter-event upstate duration (baseline = 2.13 ± 0.08 s, ketanserin + Fen = 2.21 ± 0.05 s). Right: violin plots of upstates area (baseline = 45.55 ± 2.45 mV•s, ketanserin + Fen = 39.45 ± 2.92 mV•s).
Figure 3—figure supplement 2. 5-HT suppresses evoked upstates in vitro.

(A) Experimental protocol: recording and stimulation electrode were placed in mEC layer 3, stimulation electrode was located towards the lateral side of the slice. (B) Effect of electrical stimulation before (black) and after 5-HT application (orange). 5-HT consistently suppressed spiking. Increasing the stimulation power up to 10x (n = 40/80 in 4 slices) had no rescue effect. Top: voltage responses to electrical stimulation of a representative neuron. Middle: summary spike raster plot before and after 5-HT application. Bottom: spike rate line histogram (C) Scatter plot showing area (top, n = 8 neurons, mean \(_{\text{control}} = 16.22 \pm 0.80\) \(\mu\)V, mean \(_{5-\text{HT}} = 1.24 \pm 0.97\), \(p < 10^{-4}\), Wilcoxon matched-pairs signed rank test) and duration (bottom, n = 8 neurons, mean \(_{\text{control}} = 1.92 \pm 0.07\) \(\mu\)s, mean \(_{5-\text{HT}} = 0.24 \pm 0.01\), \(p < 10^{-4}\), Wilcoxon matched-pairs signed rank test) of evoked upstates before (black) and after 5-HT application (orange). (D) Representative voltage responses to 1 second 4 Hz stimulation following wash-in (left) and wash-out (right) of 5-HT. 5-HT prevents spiking from input summation.
5-HT2ARs are involved in 5-HT mediated SOs suppression in vitro.

(A) Top: upstate raster plot during application of WAY 100635 (5-HT1A antagonist) + 5-HT. Bottom: change in RP in putative excitatory cells after application of WAY 100635 (5-HT1A antagonist) + 5-HT (n = 25 cells). (B) Top: upstate raster plot during application of ketanserin (5-HT2A antagonist) + 5-HT. Bottom: change in RP in putative excitatory cells after application of ketanserin (5-HT2A antagonist) + 5-HT (n = 21 cells). (C) Top: upstate raster plot during application of α-methyl-5-HT (5-HT2 agonist). Bottom: change in RP in putative excitatory cells after application of α-methyl-5-HT (5-HT2 agonist) (n = 11 cells). (D) Dot plot showing change in RP for each pharmacological condition (5-HT: -4.52 ± 0.64 mV, WAY + 5-HT: -2.09 ± 0.47 mV, ketanserin + 5-HT: -3.68 ± 0.60 mV and α-methyl-5-HT: -1.67 ± 1.13 mV; p = 0.0329).
Kruskal-Wallis with Dunn’s multiple comparisons test). (E) Dot plot showing percentage reduction of upstate incidence for each pharmacological condition (5-HT: 95 ± 4 %, WAY + 5-HT: 100 ± 0 %, ketanserin + 5-HT: 57 ± 10.1 % and α-methyl-5-HT: 100 ± 0 %; p < 10^-4, Kruskal-Wallis with Dunn’s multiple comparisons test).
Figure 3–figure supplement 4. 5-HT₃ receptor is not involved in 5-HT mediated SOs suppression.

(A) Top: upstate raster plot during application of m-CPBG (5-HT3 agonist). Bottom: histogram of upstate incidence before and after m-CPBG application (n = 6, baseline: 1.04 ± 0.19 upstates/10s, m-CPBG: 1.11 ± 0.21 upstates/10s). Patches represent 95% confidence interval, lines represent standard deviation.

(B) Top: upstate raster plot during application of tropisetron (5-HT3 antagonist) + 5-HT. Bottom: histogram of upstate incidence before and after tropisetron + 5-HT application (n = 6, baseline: 1.23 ± 0.18 upstates/10s, tropisetron + 5-HT: 0.01 ± 0.01 upstates/10s). Patches represent 95% confidence interval, lines represent standard deviation.
Figure 3—figure supplement 5. *In vitro* upstates metrics during baseline and ketanserin + 5-HT application.

(A) Box plot showing spiking rate before and after ketanserin + 5-HT (baseline: 0.38 ± 0.14 spikes/s, ketanserin + 5-HT: 0.15 ± 0.06 spikes/s, p= 0.011, Mann-Whitney test). Patches represent 95% confidence interval, lines represent standard deviation. (B) Violin plots of upstate inter-event interval (baseline = 10.92 ± 0.34 s, ketanserin+5-HT = 15.28 ± 1.34 s, p = 0.071, Mann-Whitney test). (C) Violin plots of upstate duration (baseline = 3.11 ± 0.16 s, ketanserin +5-HT = 1.91 ± 0.15 s, p < 0.001, Mann-Whitney test). (D) Violin plots of upstate area (baseline = 31.55 ± 1.13 mV•s, ketanserin+5-HT = 28.16 ± 1.91 mV•s, p = 0.0825, Mann-Whitney test).
Figure 3–figure supplement 6. LTS neurons are depolarized by 5-HT.

(A) PCA projection plot of all the cells recorded. Cells are color-coded according to group identity (posterior probability > 0.9): excitatory (black), fast-spiking (blue), low-threshold spiking (green). Red circles represent PCA loadings. (B) PCA projection plot color-coded according to ΔRP after 5-HT application. Inset shows a recording from one LTS neuron during 5-HT application. Scale bars: 20 mV, 25 s. Dotted line showing -70 mV. (C) PCA projection plot color-coded according to Δspikes/s after 5-HT application. (D) Posterior probability of being classified as Exc, FS or LTS.
Figure 3–figure supplement 7. Excitatory, fast-spiking and LTS cells have unique sets of electrophysiological features.
(A) Box plot showing the values of input resistance ($R_{in}$), delta after-hyperpolarization (ΔAHP), SAG, RP, rheobase and spike width of excitatory (Exc, black), fast-spiking (FS, blue) and low-threshold spiking (LTS, green) cells ($n_{Exc} = 33, n_{FS} = 6, n_{LTS} = 9$; asterisk means $p < 0.05$, double
asterisk means p < 0.01). patches represent 95% confidence interval, lines represent standard deviation. (B) Table showing values plotted in (A).
Figure 3–figure supplement 8. Classification of cells recorded in Sst-tdTomato mice.

(A) PCA projection plot of cells recorded in Sst-tdTomato mice. Cells are color-coded according to group identity (posterior probability > 0.9): fast-spiking (blue) or low-threshold spiking (green). Red circles represent PCA loadings. (B) Posterior probability of being classified as FS. (C) Posterior probability of being classified as LTS.
Figure 3–figure supplement 9. Spatial localization of 5-HT$_2A$R positive cells in EC.

(A) 3D visualization of EC (purple). (B) 3D localization of all the 5-HT$_2A$R positive cells detected in EC using same perspective as (A). (C and D) Co-localization of 5-HT$_2A$R and Sst, arrows point to colocalized cells (scale bar: 100 µm, $n_{\text{animals}}$ = 7, total number of 5-HT$_2A$R positive cells = 3570, average number of 5-HT$_2A$R positive cells per animal = 510 ± 80.32). (E) Spatial distribution of 5-HT$_2A$R positive cells and colocalized cells along the 3 dimensions depicted in (A) (Z dimension centered on the midline).
Figure 4–figure supplement 1. Vector construction and RMCE for the generation of a transgenic mouse line with Cre-conditional hM4Di expression.

(A) The coding sequence of hM4Di-mKate flanked by two opposing loxP and lox2272 sites was placed in reverse orientation to the CAG- promoter in the pRMCE. In the acceptor ES cells the ROSA26 allele harbours a PGK promoter driving the hygromycin selection marker, flanked by two attP sites. RMCE by C31int replaces the hygromycin resistance by the neomycin resistance of the donor vector. Location of primer binding sites in the Rosa26-hM4Di locus is indicated by green triangles. (B) Identification of successful genomic integration events and Cre-mediated inversion of the hM4Di coding sequence by PCRs. PCR 1 and PCR 2 test for correct integration of the 5’ and 3’ end of the construct into the ROSA26 locus. The lower band in PCR 1 results from the Neomycin resistance cassette of the feeder cells in the ES cell culture. PCR 3 tests for successful recombination of the FLEX site by Cre. A successful Cre-mediated recombination of the FLEX cassette was observed for clone 1, resulting in an 826 bp product in PCR 3. C: control cells (not electroporated), H: H2O input.
Figure 5–figure supplement 1. CNO application prevents 5-HT mediated upstates suppression.

(A) Upstate raster plot during CNO and subsequent 5-HT application. Orange box represents 5-HT, purple boxes represent CNO. CNO significantly prevents 5-HT-induced suppression of upstates. (B) Upstate incidence during 5-HT, CNO and 5-HT+CNO application (n=5, baseline: 1.13 ± 0.13 upstates/10s, CNO: 1.13 ± 0.10 upstates/10s, 5-HT: 0.93 ± 0.24 upstates/10s). Patches represent 95% confidence intervals, lines represent standard deviation.
Figure 5–figure supplement 2. CNO application in wild-type littermates and PV-hM4Di mice.

(A) CNO application in wild-type littermates. Top: Experimental protocol. Orange box represents 5-HT and purple boxes represent CNO application. Bottom: upstate incidence during 5-HT and 5-HT+CNO application, patches represent 95% confidence interval, lines represent standard deviation (n = 11 cells in 4 mice, baseline = 0.91 ± 0.10, 5-HT = 0.09 ± 0.04, CNO = 0.01 ± 0.01, p baseline vs 5-HT = 0.0004, p baseline vs CNO < 10^{-4}, Kruskal-Wallis test).

(B) Same as (A) but in PV-hM4Di mice (n = 10 cells in 4 mice, baseline = 1.02 ± 0.08, 5-HT = 0.04 ± 0.03, CNO = 0.08 ± 0.07, p baseline vs 5-HT < 10^{-4}, p baseline vs CNO = 0.0003, Kruskal-Wallis test).
References


GORINSKI, N., BIJATA, M., PRASAD, S., WIRTH, A., ABDEL GALIL, D., ZEUG, A.,
BAZOVKINA, D., KONDAUROVA, E., KULIKOVA, E., ILCHIBAEVA, T., ZAREBA-
KOZIEL, M., PAPALEO, F., SCHEGGIA, D., KOCHLAMAZASHVILI, G., DITYATEV, A.,
SMYTH, I., KRZYSTYNIAK, A., WLODARCZYK, J., RICHTER, D. W., STREKALOVA, T.,
SIGRIST, S., BANG, C., HOBUS, L., FIEDLER, J., THUM, T., NAUMENKO, V. S.,
PANDEY, G. & PONIMASKIN, E. 2019. Attenuated palmitoylation of serotonin receptor
5-HT1A affects receptor function and contributes to depression-like behaviors. Nature
Communications, 10, 3924.

GRANDJEAN, J., CORCOBA, A., KAHN, M. C., UPTON, A. L., DENERIS, E. S., SEIFRITZ, E.,

Pharmacology and Clinical Pharmacology of 3,4-Methylenedioxymethamphetamine

HARDEBO, J. E. & OWMAN, C. 1980. Barrier mechanisms for neurotransmitter monoamines

HARDY, A., PALOUZIER-PAULIGNAN, B., DUCHAMP, A., ROYET, J. P. & DUCHAMP-VIRET,
P. 2005. 5-Hydroxytryptamine action in the rat olfactory bulb: in vitro
electrophysiological patch Clamp recordings of juxtaglomerular and mitral cells.
Neuroscience, 131, 717-31.


HEIFETS, B. D., SALGADO, J. S., TAYLOR, M. D., HOERBELT, P., CARDozo PINTO, D. F.,
mechanisms for the prosocial and rewarding properties of MDMA. Sci Transl Med, 11.


HUANG, C.-C., LIANG, Y.-C., LEE, C.-C., WU, M.-Y. & HSU, K.-S. 2009. Repeated Cocaine
Administration Decreases 5-HT2A Receptor-Mediated Serotonergic Enhancement of
Synaptic Activity in Rat Medial Prefrontal Cortex. Neuropsychopharmacology, 34,

INSERRA, A., DE GREGORIO, D. & GOBBI, G. 2021. Psychedelics in Psychiatry: Neuroplastic,

ISOMURA, Y., SIROTA, A., OZEN, S., MONTGOMERY, S., Mizuseki, K., Henze, D. A. &
Buzsaki, G. 2006. Integration and segregation of activity in entorhinal-hippocampal
subregions by neocortical slow oscillations. Neuron, 52, 871-82.

JUN, J. J., STEINMETZ, N. A., SIGLE, J. H., DENMAN, D. J., BAUZA, M., BARBARITS, B.,
LEE, A. K., ANASTASSIOU, C. A., ANDREI, A., AYMIDN, Ç., BARBIC, M., BLANCHE, T.
J., BONIN, V., COUTO, J., DUTTA, B., GRATIY, S. L., GUTNISKY, D. A., HAUSSER,


MARTIN, D. A. & NICHOLS, C. D. 2016. Psychedelics Recruit Multiple Cellular Types and Produce Complex Transcriptional Responses Within the Brain. EBioMedicine, 11, 262-277.


OHMURA, Y., IZUMI, T., YAMAGUCHI, T., TSUTSUI-KIMURA, I., YOSHIDA, T. & YOSHIOKA, M. 2010. The serotonergic projection from the median raphe nucleus to the ventral hippocampus is involved in the retrieval of fear memory through the corticotropin-releasing factor type 2 receptor. Neuropsychopharmacology: official publication of the American College of Neuropsychopharmacology, 35, 1271-1278.


