Dynamic coupling of whole-brain neuronal and neurotransmitter systems

Morten L. Kringlebøch, Josephine Cruzat, Joana Cabral, Gitte Moos Knudsen, and Gustavo Deco

Remarkable progress has come from whole-brain models linking anatomy and function. Paradoxically, it is not clear how a neuronal dynamical system running in the fixed human anatomical connectome can give rise to the rich changes in the functional repertoire associated with human brain function, which is impossible to explain through long-term plasticity. Neuroradiation evolved to allow for such flexibility by dynamically updating the effectiveness of the fixed anatomical connectivity. Here, we introduce a theoretical framework modeling the dynamical mutual coupling between the neuronal and neurotransmitter systems. We demonstrate that this framework is crucial to advance our understanding of whole-brain dynamics by bidirectional coupling of the two systems through combining multimodal neuroimaging data (diffusion magnetic resonance imaging [dMRI], functional magnetic resonance imaging [fMRI], and positron electron tomography [PET]) to explain the functional effects of specific serotoninergic receptor (5-HT2AR) stimulation with psilocybin in healthy humans. This advance provides an understanding of why psilocybin is showing considerable promise as a therapeutic intervention for neuropsychiatric disorders including depression, anxiety, and addiction. Overall, these insights demonstrate that the whole-brain mutual coupling between the neuronal and the neurotransmission systems is essential for understanding the remarkable flexibility of human brain function despite having to rely on fixed anatomical connectivity.

H
duman connectomics has been very successful in revealing how function arises from structure (1, 2), and by showing how anatomy can give rise to a complex dynamical neuronal system as measured with multimodal neuroimaging (3–5). Despite the attractiveness of the idea that function is shaped by anatomy, it is also clear that the single fixed structure of the anatomical connectome should not be able to give rise to the full palette and complexity of brain function. However, evolution has found a solution to this apparent paradox by dynamically modulating the connectome over time through neurelmutilatory systems, enabling the richness of behaviors needed for survival. Indeed, the necessary dynamics of human brain function can be obtained by modulating the effective connectivity of the coupling over time, as proposed by Friston and many others (6, 7). Still, a principled and mechanistic description of the dynamic connectome must bring together the anatomical, neuronal, and neurotransmitter systems at the whole-brain level (8).

Here, we show how the mutual coupling between the neuronal and neurotransmitter systems is fundamental to understanding the dynamic connectome. This can be achieved through whole-brain modeling of multimodal neuroimaging data using a mutually coupled neuronal–neurotransmitter whole-brain model where the structural anatomical connectivity can be measured using diffusion magnetic resonance imaging (dMRI), the functional connectivity with functional magnetic resonance imaging (fMRI), and neurotransmission (receptor density) with positron electron tomography (PET). In this model, the synaptic/neuronal activity and the neurotransmitter diffusive system are portrayed in a realistic biophysical way by a set of separate dynamical equations, which are mutually coupled through the receptor maps and synaptic dynamics (neurotransmitters to neuronal), and the excitation of the projections from the brain regions producing the neurotransmitters (neuronal to neurotransmitters). The explicit linkage between these dual-coupled dynamical systems yields a deeper understanding of the crucial mutual coupling between neuronal and neurotransmitters systems at the whole-brain level. We perceive this as a significant improvement compared to our previous unique whole-brain study, which had only one neuronal dynamical system influenced by static neurotransmitter concentrations modulating the neuronal gain (9). Specifically, we demonstrate the explanatory and predictive power of this mutually coupled whole-brain model by investigating the effects of psychedelics on brain activity.

Significance

In a technical tour de force, we have created a framework demonstrating the underlying fundamental principles of bidirectional coupling of neuronal and neurotransmitter dynamical systems. Specifically, in the present study, we combined multimodal neuroimaging data to causally explain the functional effects of specific serotoninergic receptor (5-HT2AR) stimulation with psilocybin in healthy humans. Longer term, this could provide a better understanding of why psilocybin is showing considerable promise as a therapeutic intervention for neuropsychiatric disorders including depression, anxiety, and addiction.


This open access article is distributed under a Creative Commons Attribution-NonCommercial-NoDerivatives License 4.0 (CC BY-NC-ND). Database deposition: The code to run the analysis and the multimodal neuroimaging data (dMRI, fMRI, and PET) from the experiment are available on GitHub (https://github.com/decolab/pnas-neuromod). See online for related content such as Comments.

1To whom correspondence may be addressed. Email: morten.kringlebøch@psych.ox.ac.uk, niklmpg@stanford.edu, or gustavo.deco@upf.edu.

First published April 13, 2020.
Psilocybin, the prodrug of psilocin (4-OH-dimethyltryptamine), is a good model system to demonstrate the power of a mutually coupled whole-brain model, since it has been shown to act mainly through the serotonin 2A receptor (5-HT2A,R) (10), rather than more complex interactions between many receptors. The serotonin system works through the projections of the raphe nucleus. In addition, conveniently, the 5-HT receptor density maps have recently been mapped with PET (11). Here, we were interested in revealing the effects of mutual coupling of both neuronal and neurotransmitter systems on brain repertoire and specifically the effects of psilocybin on resting-state activity on healthy human participants.

Specifically, the bidirectional coupling of the neuronal and neurotransmitter systems was modeled in the following way: For the placebo condition, we used a standard whole-brain model to simulate the neuronal system, i.e., modeling spontaneous brain activity at the whole-brain level (measured with blood oxygen level-dependent [BOLD] [MR1), where each node represents a brain area and the links between them are represented by white matter connections (measured with dMRI). For the psilocybin condition, we mutually coupled the whole-brain neuronal and neurotransmitter systems by including an explicit description of the neurotransmitter dynamical system and the mutual coupling with the neuronal system. This was done by modeling the dynamics of the neurotransmitter system through simulating the release-and-reuptake dynamics, where the serotonin receptor density of each brain area is measured with PET. The neurotransmitter dynamics are then in turn coupled with the neuronal activity through the firing rate activity of the raphe nucleus, source of the serotonin neurotransmitter.

The relevance of the whole-brain modeling developed here is strongly supported by recent studies that have started to demonstrate the functional neuroanatomy underlying the experience of unconstrained cognition and enhanced mind-wandering reported following psilocybin (12–15). Due to its therapeutic action for the treatment of neuropsychiatric disorders such as depression, anxiety, and addiction (16–18), psilocybin has begun to elicit significant interest from the neuropsychiatric community as a potential treatment (19). Long term, the in silico framework presented here for investigating the dynamical connectome has the potential to bring insights and help design interventions for brain disease including neuropsychiatric disorders, which are otherwise difficult to study with traditional animal models.

Results

The main aim of the mutually coupled neuronal–neurotransmitter whole-brain model (shown in Fig. 1) is to investigate the tight interactions between these two different but mutually coupled dynamical whole-brain systems (Fig. 1A), which are fitted (Fig. 1B) to the empirical neuroimaging BOLD data (Fig. 1C). In other words, neuronal-only models that fit neuroimaging data using local node dynamics constrained by anatomical structural connectivity (20–23) are now described by a balanced dynamic mean field model that expresses consistently the time evolution of the ensemble activity of the different neural populations up the spiking network (24, 25). The spontaneous activity of each single brain region is described by a network consisting of two pools of excitatory and inhibitory neurons (Fig. 1D and Eqs. 2–7 in Materials and Methods). The neurotransmitter system, on the other hand, uses a separate set of differential equations describing the dynamics of the neurotransmitter concentration level, given by the well-known Michaelis–Menten release-and-reuptake dynamics (Eq. 13 in Materials and Methods and shown in Fig. 1E) (26–28). We couple the neurotransmitter and neuronal dynamical systems by means of the anatomical connectivity between the raphe nucleus and the rest of the brain, estimated using Human Connectome Project (HCP) dMRI (Materials and Methods). The explicit coupling between the neurotransmitter and the neuronal system is shown in Fig. 1F (and in Eqs. 14 and 15 in Materials and Methods). As can be seen, the neurotransmitter currents are applied to each region’s excitatory and inhibitory pools of neurons using the effectivity/conductivity parameters (\(W_e\) and \(W_i\), respectively). In each region, the neurotransmitter currents are also scaled by each region’s receptor density (measured in vivo using PET). The reverse coupling from the neuronal to the neurotransmitter system is given by inserting in the Michaelis–Menten release-and-reuptake equation the neuronal population firing rate of the source brain region of the neurotransmitter spread from the raphe nucleus.

As a demonstration of the power of this general framework, we show how it can be used to explain the specific case of modulation of the serotonin system by psilocybin by analyzing the neuroimaging data from a group of healthy participants being administered psilocybin i.v. (see details in Materials and Methods). This allowed us to study the relevant role of the coupling between the neuronal and neurotransmitter systems to understand the specificity of the 5-HT2A receptor densities [mapped with PET in vivo (11)]. Specifically, the main source of serotonin neurotransmission is the raphe nucleus, which is essential for the correct coupling between both dynamical systems. We therefore used the structural connectivity from this region estimated using dMRI tractography (Materials and Methods).

We fitted the mutually coupled whole-brain model using our framework of describing a brain state as an ensemble or probabilistic “cloud” in a given state space (29). This cloud is of course not clustered into discrete states (30), but it has been shown that clustering can nevertheless be useful in providing so-called “metastable substates” that can significantly distinguish between brain states (31, 32). A brain state is determined by a collection of metastable substates, i.e., of time-varying pseudo-states resulting from the clustering process (33–35).

We extracted this probabilistic metastable substate (PMS) space for the empirical psilocybin data (placebo and active condition) using the leading eigenvector dynamics analysis (LEiDA) method (described in detail in Materials and Methods and schematized in Fig. 2) (31). Briefly, we define a time-resolved dynamic FC (dFC) matrix by using BOLD phase coherence connectivity. In order to reduce the dimensionality of the problem, we compute the corresponding time-resolved leading eigenvector, which captures the dominant connectivity pattern of dFC(t) at time \(t\). Then, we determine a discrete number of metastable substates by clustering the dFC(t) across time points and subjects. The obtained \(k\) cluster centroids define the PMS space, for which we compute the probabilities, lifetimes, and transition probability between them for both the placebo and active conditions of psilocybin. The placebo and the active conditions of the psilocybin can be significantly distinguished by three substates. Fig. 3A shows three different substates of the PMS methodology with associated probabilities and transition probabilities between them for the placebo and active conditions of psilocybin. As can be seen in the subplots, two substates (1 and 3) are significantly different between the two conditions (\(P < 10^{-4}\) in terms of probability and substate 3 is significantly different for lifetimes (\(P < 10^{-3}\)). This demonstrates that the clustering approach is indeed useful for distinguishing brain states.

We first fitted the whole-brain model to the PMS space of the placebo condition of psilocybin using only the neuronal system (and thus without coupling the neurotransmitter system). We did this by fitting the minimum of the symmetrized Kullback–Leibler distance (KLD) between the empirical placebo condition PMS and modeled PMS (Materials and Methods). This yielded a global neuronal coupling parameter \(G = 1.6\). Equally, we measured the Markov entropy distance (ME) (Materials and Methods). The optimal values were found to be KLD = 0.002 and ME = 0.05. Second, we proceeded to study the role of the neurotransmitter dynamical system by coupling both systems in the model.
This allowed us to predict the changes in brain dynamics under the effects of psilocybin. Specifically, the mutually coupled whole-brain model used the same centroids found in the neuroimaging empirical data and a two-dimensional (2D) matrix of coupling parameters $W_{E}^{F}$ and $W_{I}^{F}$ for generating the modeled PMS (Fig. 3B). We searched this matrix for the optimal fit (global minimum) to the empirical neuroimaging data, which is described by probabilistic metastable substate (PMS) space (C), which is extracted from the empirical BOLD data. (D) We achieve this by adding a coupled neurotransmitter system (blue) to modulate and interact with the neuronal system (green), which was modeled using a balanced dynamic mean field model that expresses the steady state is achieved, and then we take the average of the firing rates of the ensemble activity of the different neuronal populations building up the spiking network (24, 25). (E) The neurotransmitter system (blue) is modeled by a set of differential equations describing the spontaneous activity of each single brain region consisting of two pools of excitatory and inhibitory neurons and the rest of the brain, estimated using dMRI from the HCP (Materials and Methods). We couple the neurotransmitter and neuronal dynamical systems through the anatomical connectivity between the raphe nucleus and the rest of the brain, estimated using dMRI from the HCP (Materials and Methods). The explicit coupling between the neurotransmitter and the neuronal system is given in Eqs. 16–17 (shown here and described in Materials and Methods). As can be clearly seen, the neurotransmitter currents are applied to each region’s excitatory and inhibitory pools of neurons using the effectiveness/conductivity parameters ($W_{E}$ and $W_{I}$, respectively). In each region, the neurotransmitter currents are also scaled by the region’s receptor density (measured in vivo using PET). The reverse coupling from the neuronal to the neurotransmitter system is given by inserting in the Michaelis–Menten release-and-reuptake dynamics (Materials and Methods) (26–28). (F) The neuronal coupling dynamics (green) is modeled by another set of differential equations describing the spontaneous activity of each single brain region consisting of two pools of excitatory and inhibitory neurons (Materials and Methods). We couple the neurotransmitter and neuronal dynamical systems through the anatomical connectivity between the raphe nucleus and the rest of the brain, estimated using dMRI from the HCP (Materials and Methods). The explicit coupling between the neurotransmitter and the neuronal system is given in Eqs. 16–17 (shown here and described in Materials and Methods). As can be clearly seen, the neurotransmitter currents are applied to each region’s excitatory and inhibitory pools of neurons using the effectiveness/conductivity parameters ($W_{E}$ and $W_{I}$, respectively). In each region, the neurotransmitter currents are also scaled by the region’s receptor density (measured in vivo using PET). The reverse coupling from the neuronal to the neurotransmitter system is given by inserting in the Michaelis–Menten release-and-reuptake equation the neuronal population firing rate of the source brain region of the neurotransmitter spread from the raphe nucleus.
unable to influence the neurotransmitter dynamics that are frozen.

Fig. 4C shows the significant difference between using the empirical 5-HT₂A receptor densities across the regions at the optimal fit compared with the results of randomly shuffling the receptor densities ($P < 10^{-4}$). Similarly, as shown in Fig. 5A, we also tested the specificity of the receptor binding maps by comparing (at the optimal fit) the 5-HT₂A receptor with the other serotonin receptors, namely 5-HT₁A, 5-HT₁B, and 5-HT₄, which showed significant differences between 5-HT₂A and 5-HT₁A ($P < 10^{-6}$), 5-HT₁B ($P < 10^{-6}$), and 5-HT₄ ($P < 10^{-6}$). This strongly confirms the main role this receptor plays in the effects of psilocybin (10). Finally, in Fig. 5B, we show the comparison to the 5-HTT, which is also significant ($P < 0.02$). 5-HTT is not a receptor but has been shown to play an important role for the treatment of depression.

**Discussion**

Here, we have shown how a dynamic mutually coupled whole-brain model can address the major challenge in neuroscience, which is to explain the paradoxical flexibility of human brain function despite the reliance on a fixed anatomical connectome. One of the most important elements in this flexibility is the way that the fixed connectome can be modulated by neuromodulators to selectively change the balance of the excitation and inhibition of individual brain regions. Here, we modeled the dynamical mutual coupling between neuronal and neuromodulator systems at the whole-brain level. In particular, we implemented a mutually coupled dynamical system, which couples at the whole-brain level the neuronal system, which was modeled using a balanced dynamic mean field model (24, 25) and the neuromodulator system describing the dynamics of the neurotransmitter concentration levels (measured in vivo using

**Fig. 2.** Computing the probabilistic metastable substate (PMS) space for whole-brain activity. For illustrative purposes in the following, we sketch the full process of computing the PMS space. For all parcellated brain regions of each participant (A), we extract the original BOLD signal (orange line), bandpass filter between 0.02 and 0.1 Hz (blue line), and then use the Hilbert transform to compute the time-varying amplitude $A$ and its phase $\theta$ (with real and imaginary components) of the BOLD signal (black dotted lines). The red arrows represent the BOLD phase at each TR, and, as can be seen, much of the original BOLD signal is captured by the BOLD phase, $\cos(\theta)$. (B) We compute the BOLD phase coherence matrix between brain regions (Lower) and extract its leading eigenvector $\mathbf{V_1}(t)$ for every timepoint $t$ (Top, colored red and blue according to their sign). (C) Finally, to determine the PMS space, we gather all leading eigenvectors $\mathbf{V_1}(t)$ for all participants for every time point as a low-resolution representation of the BOLD phase coherence patterns (leftmost panel). We apply a clustering algorithm ($k$-means), which divides the data into a predefined number of clusters $k$ (here $k = 3$). Each cluster is represented by a central vector (green, red, and blue), which represents a recurrent pattern of phase coherence, or substate, which occurs with a given probability (rightmost panel). Any given brain state can thus be represented by this PMS space.
PET) and modeled by the well-known Michaelis–Menten release-and-reuptake dynamics (26–28). Here, as proof of principle, we consider the effects of psilocybin on the serotonin system and therefore use anatomical connectivity between the raphe nucleus and the rest of the brain to couple the two systems.

Overall, the results show that the interaction between these dynamical systems is fundamental for explaining the empirical data. In other words, the dynamic mutual interaction between neuronal and neuromodulator systems at the whole-brain level is important to fully explain the functional modulation of brain activity by psilocybin, a powerful psychedelic drug, acting on the serotonin system. This is especially important given the demonstrated ability of psilocybin to rebalance the human brain in treatment-resistant depression. The results provide evidence for how the integration of dMRI (anatomy), fMRI (functional neuronal activity), and PET (neurotransmitter system) at the whole-brain level is necessary for predicting properly brain dynamics as a result of the mutual coupling between a dual dynamical system. This expands on the rich existing experimental and theoretical research on the local effects of neurotransmitters (e.g., refs. 36–40).

Specifically, the results provide insights into the underlying dynamics of neurotransmission involved in psilocybin. In terms of dynamics, we first uncoupled the neuromodulators from the neuronal systems and found that this produced a highly significant breakdown in fitting the empirical data (Fig. 4A). We then ran further simulations to investigate the role of specific parts of the dynamic coupling. The full mutually coupled whole-brain model ran until a steady state was achieved, upon which we then froze the feedback dynamics from neuromodulators to neuronal system by removing coupling through the raphe nucleus. This again produced a highly significant breakdown in fitting the empirical data (Fig. 4B).

In further sets of experiments designed to investigate the importance of the receptor distribution, we changed the distribution of regional receptor densities by 1) randomly shuffling the 5-HT2A (Fig. 4C and 2) replacing them with those of other serotonin receptors, known to be much less sensitive to psilocybin, namely 5-HT1A, 5-HT1B, and 5-HT3. Again, this produced a highly significant breakdown in the ability to explain the empirical data (Fig. 4B). This result confirms the crucial, causative role of the precise anatomical location and density of 5-HT2A.

Interestingly, as mentioned above, psilocybin has been demonstrated to play a role in rebalancing the human brain in treatment-resistant depression (19). The therapeutic action of psilocybin in depression is thought to depend on activation of serotonin 2A receptors, thereby initiating a multilevel plasticity (41). This is different from selective serotonin reuptake inhibitors, which are the most frequently prescribed antidepressant drug class and whose therapeutic action is thought to begin with reuptake blockade at the 5-HTT. We were, therefore, interested in further investigating this by replacing the receptor densities with the 5-HTT densities. We found that the fit with 5-HTT was significantly worse than 5-HT2A (Fig. 5C) and supports the potential for psilocybin in rebalancing brain function in treatment-resistant depression (19).

More broadly, the mutually coupled whole-brain model and results shed important light on our understanding of human brain function. Over the last decade, the discovery of whole-brain modeling has led to important new insights for explaining the principles of human brain function based on human neuroimaging data. The initial breakthrough was the recognition that the anatomical connectivity of the human brain shapes function at the whole-brain level (1, 2). However, it also became clear relatively quickly that this was a by-product of the close link between anatomy and function, given that it is possible to describe static spontaneous activity (i.e., grand average functional spatial connectivity over longer periods) even in the absence of neuronal dynamics, i.e., combining the underlying anatomy with noise (42). Still, it was shown by further investigations that capturing the dynamics of the spatiotemporal whole-brain activity requires more sophisticated dynamical modeling (3, 35, 43). All of these whole-brain studies were initially solely based on neuronal modeling, but recently it was shown that even better results can be obtained when the dynamics of the neuronal system (through the neuronal gain of excitatory neurons) is influenced by static neuromodulation (9).
Here, we have demonstrated the importance of having a biophysically realistic dynamic neuromodulator system, and more importantly the need for full mutual coupling between the full dynamics of both neuronal and neuromodulation systems. In other words, this framework for mutually coupled whole-brain modeling involves the underlying anatomical connectivity and two mutually interacting dynamical neuronal and neuromodulator systems.

Overall, the framework put forward here is likely to be essential for accurately modeling and explaining mechanisms of human brain function in health and disease. The framework combines multimodal neuroimaging from dMRI (anatomy), fMRI (functional neuronal activity), and PET (neurotransmitter system) at the whole-brain level. This offers unique opportunities for investigating how to rebalance human brain activity in disease through pharmacological manipulation that can be discovered and tested in silico in the mutually coupled whole-brain model proposed here. This whole-brain modeling can thus be used to make predictions that can be tested causally with electromagnetic stimulation (deep brain stimulation [DBS] and transcranial magnetic stimulation) or pharmacological interventions. In the future, this framework could further extended by measuring even faster whole-brain dynamics (measured with magnetoencephalography) (44) and incorporating gene expression at the whole-brain level, offering even more exciting avenues for discovering new and effective ways of rebalancing the human brain in disease.

Fig. 4. Optimal dynamical coupled neuronal–neurotransmission (CNN) whole-brain model is significantly better than alternative models. Further insights into the underlying dynamics of neuromodulation involved in psilocybin were obtained by comparing the Kullback–Leibler distance (KLD) for the PMS and the error lifetimes between the empirical data and the whole-brain model undergoing various manipulations. (A) For the optimal fit of the mutually coupled whole-brain model, we found a very significant difference between the optimal fit and the uncoupled system (i.e., without neuromodulation) ($P < 10^{-6}$). (B) We also found a significant difference when removing the feedback dynamics ($P < 10^{-6}$). We tested this by allowing optimal model coupling until steady state, at which point we kept just the average of the neurotransmitter variables while cancelling all feedback dynamics. (C) We found a significant difference between using the empirical 5-HT$_{2A}$ receptor densities across the regions at the optimal fit compared with randomly shuffling the receptor densities ($P < 10^{-4}$). This demonstrates the causal importance of the 5-HT$_{2A}$ receptor densities.
**Materials and Methods**

We provide here the details on the pipeline used to integrate structural and functional connectivity (dMRI and fMRI) with neurotransmission (PET) in a model of the placebo and psilocybin response in healthy participants (summarized in Fig. 1). Please note that all structural, functional and neurotransmission data are integrated into the Automated Anatomical Labeling (AAL) parcellation:

1. Structural connectivity: probabilistic tractography derived from the dMRI;
2. Functional dynamics: functional dynamics described as probability meta-stable states were estimated from the fMRI data obtained in the placebo and psilocybin condition;
3. Neurotransmitter receptor density: estimation of the density of the 5HT-2A receptors that has been obtained using PET;
4. Whole-brain neuronal model: description of the (uncoupled) neuronal dynamic mean field model used to fit the placebo condition;
5. Mutually coupled neuronal and neurotransmission whole-brain model: dynamic coupling of the neuronal and neurotransmission models to dynamically fit psilocybin condition;
6. Empirical fitting of mutually coupled whole-brain model to neuroimaging data: measuring the fit of model to the empirical neuroimaging data.

**Parcellation.** Based on our previous whole-brain studies, we used the AAL atlas but considered only the 90 cortical and subcortical noncerebellar brain regions (45). All structural, functional, and neuro-modulation data were integrated using this atlas.

**Structural connectivity.**

**Ethics.** The study was approved by the Research Ethics Committee of the Central Denmark Region (De Videnskabsetiske Komitee for Region Mid-tjylland). Written informed consent was obtained from all participants prior to participation.

**Participants and acquisition.** The dMRI data used in the study was collected from 16 healthy right-handed participants at Aarhus University, Denmark (11 men; mean age, 24.8 ± 2.5) (46, 47). We recorded the imaging data in a single session on a 3-T Siemens Skyra scanner at Center of Functionally Integrative Neuroscience, Aarhus University, Denmark. The structural MRI TI scan used the following parameters: reconstructed matrix size, 256 x 256; voxel size of 1 mm³; echo time (TE) of 3.8 ms; and repetition time (TR) of 2,300 ms. We collected dMRI using the following parameters of TE = 84 ms, TR = 9,000 ms, flip angle = 90°, reconstructed matrix size of 106 x 106, voxel size of 1.98 x 1.98 mm with slice thickness of 2 mm and a bandwidth of 1,745 Hz/Px. The data were recorded with 62 optimal nonlinear diffusion gradient directions at b = 1,500 s/mm². Approximately one nondiffusion weighted image (b = 0) per 10 diffusion-weighted images was acquired. We collected two dMRI datasets: one with anterior-to-posterior phase encoding direction and the other acquired in the opposite direction.

**Tractography.** We used the structural connectivity between 90 AAL regions in the dMRI dataset described above. First, we used the linear registration tool from the FSL toolbox (https://fsl.fmrib.ox.ac.uk/fsl/fslwiki; Functional Magnetic Resonance Imaging of the Brain [FMRIB], Oxford, UK) (48) to coregister the echo-planar imaging (EPI) image to the T1-weighted structural image. In turn, the T1-weighted image was co-registered to the T1 template of ICBM152 in Montreal Neurological Institute (MNI) space (49). We concatenated and invered the resulting transformations, which were further applied to warp the AAL template (45) from MNI space to the EPI native space; ensured discrete labeling values were preserved by interpolation using nearest-neighbor method. The brain parcellations were thus conducted in each individual’s native space. The structural connectivity maps were generated for each participant using the dMRI data acquired. We combined the two datasets acquired with different phase encoding to optimize signal in difficult regions. Constructing these structural connectivity maps or structural brain networks involved a three-step process: 1) Regions of the whole-brain network were defined using the AAL template as used in the functional MRI data. 2) Probabilistic tractography was used to estimate the connections between nodes in the whole-brain network (i.e., edges). 3) Data were averaged across participants.

**The FSL diffusion toolbox (Fdt) in FSL was used to carry out the various processing stages of the dMRI dataset. The default parameters of this imaging preprocessing pipeline were used for all participants. We then estimated the local probability distribution of fiber direction at each voxel (50). The probtrackx tool in Fdt was used to provide automatic estimation of crossing fibers within each voxel, which significantly improves the tracking sensitivity of nondominant fiber populations in the human brain (51).**

We defined the connectivity probability from a given seed voxel i to another voxel j by the proportion of fibers passing through voxel i that reach voxel j when using a sampling of 5,000 streamlines per voxel (51). We then extended the voxel level to the regional level, i.e., to a parcel in the AAL consisting of n voxels, i.e., 5,000 x n fibers were sampled. This allowed us to compute the connectivity probability $P_{ij}$ from region $n$ to region $p$ as
the number of sampled fibers in region \( n \) that connect the two regions divided by \( 5,000 \times v \), where \( v \) is the number of voxels in a given AAL region \( i \).

We computed the connectivity probability from a given brain region to each of the other 89 regions within the AAL. It is important to note that the dependence of tractography on the seeding location, i.e., the probability from voxel \( i \) to voxel \( j \) is not necessarily equivalent to that from \( j \) to \( i \). Still, these two probabilities are highly correlated across the brain for all participants (the least Pearson \( r = 0.70 \), \( P < 10^{-26} \)). We defined the unidirectional connectivity probability \( P_{ij} \) between regions \( n \) and \( p \) by averaging these two connectivity probabilities since directionality of connections cannot be determined based on dMRI. This unidirectional connectivity can be thought of as a measure of the structural connectivity between the two areas, with \( C_{np} = C_{pn} \). The computational of the regional connectivity probability was implemented using in-house Perl scripts. For both phase encoding directions, we constructed 90 \( \times \) 90 symmetric weighted networks, based on the AAL90 parcellation, and normalized by the number of voxels in each AAL region. This represents the structural connectivity network organization of the brain.

In order to consider the serotonin source region, we included in our model the connectivity between the raphe nucleus region and the rest of the brain. In this way, the neuronal activity across the whole brain excites through the existing fibers the raphe neurons, and this information is used in the neuromotor system (see below). In brief, we used the estimate of the raphe nucleus location from the Harvard Ascending Arousal Network Atlas (52). We then used LeadDBS Mapper (https://www.lead-dbs.org/) within Matlab to estimate the whole-brain tractography from this mask of the raphe nucleus to each AAL region, represented as a vector of 90 values. This represents the structural connectivity network organization of the brain.

The fMRI data were preprocessed using MELODIC 3.14 (60), part of FSL (FMRIB’s Software Library; https://fsl.fmrib.ox.ac.uk/fsl/fslwiki) with standard parameters and without discarding any ICA components. For each participant and for each brain state (i.e., placebo and psilocybin), we used FSL tools to extract and average the BOLD time courses from all voxels within each region defined in the AAL atlas (considering only the 90 cortical and subcortical noncerebellar brain regions) (45).

**Participants in psilocybin study.** fMRI data from nine healthy subjects were included in this study (seven men; age, 32 \( \pm \) 2.60 years). Using in-house Perl scripts. For both phase encoding directions, we constructed 90 \( \times \) 90 symmetric weighted networks, based on the AAL90 parcellation, and normalized by the number of voxels in each AAL region. This represents the structural connectivity network organization of the brain.

In order to consider the serotonin source region, we included in our model the connectivity between the raphe nucleus region and the rest of the brain. In this way, the neuronal activity across the whole brain excites through the existing fibers the raphe neurons, and this information is used in the neuromotor system (see below). In brief, we used the estimate of the raphe nucleus location from the Harvard Ascending Arousal Network Atlas (52). We then used LeadDBS Mapper (https://www.lead-dbs.org/) within Matlab to estimate the whole-brain tractography from this mask of the raphe nucleus to each AAL region, represented as a vector of 90 values. This represents the structural connectivity network organization of the brain.
Neurotransmitter receptor density. A previous study has carefully described the methods used to obtain the 5-HT_{2A} receptor density distribution (11), but here we briefly summarize the main methods.

Ethics. The study was approved by the Ethics Committee of Copenhagen and Frederiksborg, Denmark. Written informed consent was obtained from all participants prior to participation.

Participants. The participants were healthy male and female controls from the freely available Cimbi database (63). The present data analysis was restricted to include individuals aged between 18 and 45 y. Acquisition of PET data and structural MRI scans were taken from 210 individual participants, yielding a total of 232 PET scans; of which 189 participants had only one scan, 20 participants had two scans, and a single had three scans.

PET and structural MRI. Full details on PET and structural MRI acquisition parameters can be found in the original study (63) and in abbreviated form in ref. 9.

Extracting receptor density maps. Similar to Deco et al. (9), we extracted the average receptor density for each individual AAL region S-HT_{2A}, S-HT_{1B}, S-HT_{2A}, and S-HT_{A}, as well as S-HTT using standard FSL tools on the freely available receptor density maps in MNI space.

Whole-brain model neuronal system. Whole-brain neuronal model (anatomy-plus-neuronal activity). First, we modeled the system without any coupling between systems, i.e., a pure neuronal-dynamical system (9). For this, we simulated the spontaneous brain activity at the level of the whole brain in a network model where each node represents a brain region and the links between nodes represent white matter connections. As proposed by Deco et al. (25), a dynamic mean field (DMF) model is used to simulate the activity in each brain region. In brief, based on the original reduction of Wong and Wang (64), this DMF model uses a reduced set of dynamical equations describing the activity of coupled excitatory (E) and inhibitory (I) pools of neurons to describe the activity of large ensembles of interconnected excitatory and inhibitory spiking neurons. The inhibitory currents, I(t), are mediated by GABA_A receptors, while excitatory synaptic currents, E(t), are mediated by NMDA receptors. In terms of connectivity, each brain region, n, consists of reciprocally connected excitatory and inhibitory pools of neurons, whereas coupling between two areas n and p occurs only at the excitatory-to-excitatory level where it is scaled by the underlying anatomical connectivity C_{np} (Materials and Methods: Structural Connectivity).

The following system of coupled differential equations expresses the DMF model at the whole-brain level:

\[
\begin{align*}
\frac{dE_n}{dt} &= W_{E} + G_{E} + J_{E} \left( \sum_{p} C_{np} \sigma_{E_n}(E_p) - J_{E_n} \right), \\
\frac{dI_n}{dt} &= H_{I} \left( \sigma_{I_n}(I_n) \right), \\
\end{align*}
\]

where E_n and I_n represent the total input current (in nano-amperes), the firing rate is denoted by the vector I(t) (in hertz), while the synaptic gating is denoted by the vector \( \sigma_{E_n}(E_p) \). The total input currents received by the E and I pools is converted by the neuronal response functions, \( H_{I}(\cdot) \), into firing rates, \( I_n(t) \), using the input-output function of Abbott and Chance (65), where the gain factors \( g_E = 310 \text{ nC}^{-1} \) and \( g_I = 615 \text{ nC}^{-1} \) determine the slope of H. Above the threshold currents of \( I_n^T = 0.403 \text{ nA}, \) and \( I_n^{I} = 0.288 \text{ nA} \), the firing rates increase linearly with the input currents. The shape of the curvature of H around \( I_n^T \) is given by the constants \( d_{E} = 0.16 \) and \( d_{I} = 0.087 \). GABA receptors with \( \tau_{GABA} = 0.01 \text{s} \) controls the average synaptic gating in inhibitory pools. NMDA receptors with a decay time constant \( \tau_{NMDA} = 0.1 \text{s} \) and \( \gamma = 0.641/1000 \text{ (the factor 1,000 is for expressing everything in milliseconds) control the synaptic gating variable of excitatory pools, } S_n^E. \) All excitatory synaptic couplings is weighted by \( J_{\text{E}_{A}} = 0.015 \text{nA} \), while the weight of recurrent excitation is \( w_n = 1.4 \). The overall effective external input is \( I_n = 0.382 \text{nA} \) with \( W_E = 1 \) and \( W_I = 0.7 \). Note that in Eqs. 6 and 7, the uncorrelated standard Gaussian noise, \( \nu_n \), has an amplitude of \( \sigma = 0.01 \text{nA} \).

Emulating the resting-state condition, we used parameters in the DMF model based on Wong and Wang (64) such that each isolated node exhibited the typical noisy spontaneous activity with low firing rate (\( r(t) = 3 \text{ Hz} \) observed in electrophysiology experiments (66-69). Furthermore, similar to Deco et al. (25), for each node n, we adjusted the inhibition weight, \( J_{\text{E}_{A}} \), such that the firing rate of the excitatory pools \( S_n^E \) remained clamped at 3 Hz—even when receiving excitatory input from connected areas. The algorithm for achieving feedback inhibition control is described by Deco et al. (25), where it was shown that using this leads to a better prediction of the resting-state FC and to a more realistic evoked activity.

The whole-brain network model used parcellated structural and functional MRI data from 90 cortical and subcortical brain regions (9), where each brain region \( n \) receives excitatory input from all structurally connected regions \( p \) into its excitatory pool, weighted by the underlying anatomical connectivity matrix, \( C_{np} \), obtained from dMRI (Materials and Methods: Structural Connectivity). Importantly, we used a global coupling factor \( G \) to equally scale all interarea E-to-E connections. Finding the optimal working point of the system simply requires finding the optimal global scaling factor, such that the simulated activity by the model is maximally fitting the empirical resting-state activity of participants in the placebo conditions.

The simulations were run with a time step of 1 ms for a range of \( G \) between 0 and 2.5 (with increments of 0.025). In order to emulate the empirical resting-state scans from nine participants, for each value of \( G \), we ran 200 simulations of 435 s each.

Please note that to transform the simulated mean field activity from our DMF model in to a BOLD signal, the generalized hemodynamic model of Stephan et al. (70) was used. This involves computing the BOLD signal in each brain area, n, arising from the firing rate of the excitatory pools \( S_n^E \), that an increase in the firing rate causes an increase in a vasodilatory signal, \( s_n \), subject to autoregulatory feedback. In proportion to this vasodilatory signal, the blood inflow \( f_n \) induces changes in deoxyhemoglobin content \( q_n \) and blood volume \( v_n \).

These biophysical variables are described by these equations:

\[
\begin{align*}
\frac{ds_n}{dt} &= 0.5 d_{E_n} + 3 - k_{s_n} - \rho (f_n - 1), \\
\frac{df_n}{dt} &= s_n, \\
\tau \frac{d\nu_n}{dt} &= f_n - \nu_n, \\
\frac{d\sigma_n}{dt} &= f_n (1 - \sigma_n) - \rho - q_n v_n \frac{\sigma_n}{v_n}, \\
\end{align*}
\]

where \( \tau = \) a constant, \( \rho = \) the resting oxygen extraction fraction, and the resistance of the veins is represented by \( \alpha \).

For each area, \( n \), the BOLD signal, \( B_n \), is a static nonlinear function of deoxyhemoglobin content, \( q_n \), and volume, \( v_n \), that comprises a volume-weighted sum of extravascular and intravascular signals:

\[
B_n = V_0 \left( k_1 (1 - q_n) + k_2 \frac{1 - q_n}{v_n} + k_3 (1 - v_n) \right).
\]

We used all biophysical parameters stated by Stephan et al. (70) and concentrated on the most functionally relevant frequency range for resting-state activity, i.e., both simulated and empirical and BOLD signals were bandpass filtered between 0.1 and 0.01 Hz (71-74).

Mutually coupled whole-brain neuronal and neurotransmitter system. Finally, in order to fully couple the whole-brain neuronal and neurotransmitter systems, we have to include an explicit description of the neurotransmitter dynamical system and the mutual coupling. We do this by modeling the dynamics of the neurotransmitter system through simulating the release-and-reuptake dynamics. These dynamics are then in turn coupled with the neuronal activity through the firing rate activity of the raphe brain region, source of the serotonin neurotransmitter.
Specifically, the concentration of serotonin is modeled through the Michaelis–Menten equations (26–28):

$$\frac{d[SN]}{dt} = -c[ER]_n[SN] - V_{max}[SN] \left( \frac{[SN]}{K_s + [SN]} \right)$$

where \([SN]\) is a vector, denoting the dynamics of the neurotransmitter concentration level in a brain region \(n\) and where the corresponding density of a serotonin receptor \(R_s\) is measured with PET. This is defining the interaction between the neuronal and neurotransmitter system through the first term on the right-hand side in Eq. 13 (i.e., brain raphé coupling). The \(c[ER]_n\) is outer product of the fiber density connectivity vector between the whole brain and the raphe nucleus, obtained through dMRI probabilistic tractography, and \(s = 5\) is a factor normalizing the activity such that the current generated by the neurotransmitter (Eqs. 14–16) is working in the nonlinear central sigmoidal part. For the second term on the right-hand side of Eq. 13, \(V_{max} = 1,300 \text{ nM}^{-1} \text{s}^{-1}\) is the maximum reuptake rate and \(K_s = 170\) is the substrate concentration where the uptake proceeds at one-half of the maximum rate (26–28).

The reverse coupling, i.e., the effect of the neurotransmitter system on the neuronal system, is described in Eqs. 14–16. Specifically, the neuronal activity is modeled as a dynamical system in Eqs. 14 and 15 (as a coupled variation of Eqs. 2 and 3) by generating an extra current on the excitatory pyramidal and GABAergic inhibitory neurons in the following way:

$$\dot{E}_i^H = \beta L_i + \frac{1}{2} \sum_{j=1}^{N} w_{ij} E_j + \sum_{j=1}^{N} J_{SN}^{[18]}(E_i, E_j), \quad \dot{E}_i^L = \beta L_i - \frac{1}{2} \sum_{j=1}^{N} w_{ij} E_j - \sum_{j=1}^{N} J_{SN}^{[18]}(E_i, E_j),$$

where \(\dot{E}_i^H\) and \(\dot{E}_i^L\) are the excitatory and inhibitory feedback coupling parameters from the neurotransmitter system to the neuronal activity. The density of a serotonin receptor \(R_s\) (as measured with PET) weighs the serotonin modulated current vector \(M_n\) (current for each brain region), which is given by the sigmoidal-like function commonly used in pharmacology and current equation (26):

$$\tau_M \frac{dM_n}{dt} = -M_n + \frac{j}{1 + e^{-\alpha(ERSn[i] + 1)}}$$

where \(\beta = 10, J = 0.1\), and \(\tau_M = 120\) ms (26). The last terms on the right-hand side of Eqs. 14 and 15 describe the coupling of the neuronal to the neurotransmitter system. In this way, both neuronal and neurotransmitter dynamical system are explicitly expressed and mutually coupled. Empirical fitting of mutually coupled whole-brain model to neuroimaging data. We used the following measurements to measure the empirical fit of the mutually coupled whole-brain system.

Comparing empirical and simulated PMS space measurements. We computed the symmetrized KLD between the simulated and empirical corresponding probabilities of the metastable substates, i.e., the probabilities of the extracted empirical centers after clusteringization:

$$KL(P_{emp}, P_{sim}) = 0.5 \sum_i P_{emp}(i) \log \left( \frac{P_{emp}(i)}{P_{sim}(i)} \right) + \sum_i P_{sim}(i) \log \left( \frac{P_{emp}(i)}{P_{sim}(i)} \right).$$

where \(P_{emp}(i)\) and \(P_{sim}(i)\) are the empirical and simulated probabilities on the same empirical extracted metastable substates \(i\).

Comparing empirical and simulated transition probabilities between metastable substates. We computed the entropy rate \(S\) of a Markov chain, with \(N\) states and transition matrix \(P\). The rate entropy \(S\) is given by the following:

$$S = S_1 + S_2 + \ldots + S_N,$$

where \(S_i = -p(i) \sum_{j=1}^{N} p(i,j) \log p(i,j),$$

The stationary probability of state \(i\) is given by \(p(i)\). For long realizations of the Markov chain, the probabilities of each state converge to the stationary distribution \(p\), which is the solution of the following equation:

$$p^2 = p.$$