### Posterior integration and thalamo-frontotemporal broadcasting are impaired in disorders of consciousness

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#### **Abstract:**

The study of the brain's dynamical activity is opening a valuable source of assistance for the clinical diagnosis of patients with disorders of consciousness (DOC). For example, dysfunctional spread of naturalistic and synthetic stimuli has proven useful to characterize hampered consciousness. However, understanding of the mechanisms behind loss of consciousness following brain injury is still missing. Here, we study the propagation of endogenous and *in-silico* exogenous perturbations in patients with DOC, based upon directed and causal interactions estimated from resting-state fMRI. We found that patients with DOC suffer decreased capacity for neural propagation and responsiveness to events. Particularly, that loss of consciousness is related to the malfunctioning of two neural circuits: the posterior cortical regions failing to convey information, in conjunction with reduced broadcasting of information from subcortical, temporal, parietal and frontal regions. These results seed light on the mechanisms behind DOC, thus opening new possibilities for clinical applications.

#### I. Introduction

Consciousness is a subjective experience. Internally perceived as the personal experience of "what is it like, to be you", the definition of consciousness and its origin are still a matter of scientific and philosophical debates without consensus<sup>1-4</sup>. Within the clinical context, however, practitioners treating patients with severe brain injuries and disorders of consciousness (DOC) face the daily reality to help their patients in the best possible manner, regardless of the exact definition of consciousness. Therefore, it is important to count with tangible and measurable correlates of consciousness in order to accurately assess the state of the patients at the behavioral level. The use of neuroimaging proxies can thus improve diagnosis and decision making, and contribute to clarifying the mechanisms behind pathological loss of consciousness<sup>5</sup>.

Behavioral assessment such as the response to sensory stimuli, pain or simple commands is the first line of action taken at bedside. From this perspective, it has proven useful to characterize consciousness based upon two components: wakefulness (the level of arousal) and awareness (the content of consciousness)<sup>6,7</sup>. Patients with severe brain injury can fall into a coma, which is characterized by the absence of both wakefulness and awareness. Patients surviving coma often recover signs of wakefulness, i.e. eye opening, but without manifestation of awareness of the self nor of the environment. Such state is known as unresponsive wakefulness syndrome (UWS) or vegetative state<sup>6</sup>. Some of these patients gradually regain awareness and progress into the so-called minimally conscious state (MCS), showing a wider range of non-reflexive behaviors such as visual pursuit, localization to pain or response to simple commands, although their ability to functionally communicate remains hampered<sup>8</sup>.

An upcoming approach to assess brain states in the clinical environment relies on the analysis of the brain's dynamical activity. It is well-known that neural activity is characterized by different frequency bands across sleep stages<sup>9</sup> or cognitive circumstances, and that local field potentials display intercalated epochs of bursting activity followed by silent periods during anesthesia<sup>10</sup>. Recent studies have shown that loss of consciousness leads to reduced spontaneous neural activity<sup>11</sup> and metabolism<sup>12,13</sup>, and that functional connectivity between brain or cortical regions is also significantly reduced<sup>7,13,14</sup>. Moreover, the fluctuating patterns of functional connectivity are altered during reduced consciousness, with shorter life-times and more random transitions between the patterns as observed in normal awake<sup>14–17</sup>.

Observing how external perturbations propagate through the brain constitutes an indirect window to probe the brain dynamics, and thus its state. For example, natural audio-visual stimuli presented to subjects undergoing general anesthesia or within deep sleep are still processed in the sensory cortices but fail to integrate at the higher level cortical regions <sup>18,19</sup>. Application of artificial perturbations such as transcranial magnetic stimulation triggers a response of the stimulated regions that is comparable in all cases, but a rapid decline in the propagation of the signals is found during deep sleep, anesthesia or patients with DOC<sup>20,21</sup>. These observations have been successfully employed to classify the level of consciousness both in patients and during anesthesia<sup>21</sup>. However, as the procedure focuses on the description of the whole-brain responses by a single number – the perturbational complexity index, it misses the directionality of the evoked causal interactions. These causal interactions have been demonstrated to be sensitive to different states of consciousness and moreover to hold explanatory power with respect to their neural mechanisms<sup>22,23</sup>.

In the present paper, we investigate the capacity of both endogenous and exogenous events to propagate along the brain in patients with disorders of consciousness as compared to normal wakefulness. By use of model-free and model-based analysis methods, all relevant information to characterize the potential of stimuli to propagate is extracted from the resting-state activity, as measured via functional MRI. Thus, bypassing the need to carry out clinical stimulation protocols. First, we studied how spontaneous endogenous events observed within the resting-state BOLD propagate and are subsequently integrated<sup>24</sup>. We found that the autocovariance relaxation times of the BOLD signals exhibit a spatial distribution in healthy controls which was disrupted in DOC patients, especially in the UWS group, followed by a significantly reduced capacity to integrate endogenous events. Then, we employed a model-based approach to estimate the effective connectivity between pairs of brain regions<sup>25–27</sup>. Since effective connectivity captures the directional causal relations, we could simulate the asymmetrical propagation of exogenous perturbations on the network in order to identify feedforward and backward effective pathways, and to recognize changes in the ability of brain areas to 'broadcast' or to 'receive' information. In particular, we found two well-differentiated subnetworks with altered propagation properties in the patients. The posterior regions of the cortex fail to convey information, while broadcasting of information is reduced in subcortical, temporal, parietal and frontal regions. These results evidence that patients with prolonged disorders of consciousness lack of the capacity for the integration of events that would lead to conscious perception.

#### II. Results

This study comprises resting-state fMRI (eyes-closed) of 33 healthy control (HC) subjects, 14 patients with unresponsive wakefulness syndrome (UWS) and 26 patients classified as in minimally conscious state (MCS). The diagnoses were made using repeated CRS-R assessments and confirmed with FDG-PET neuroimaging 12 to avoid including MCS\* patients 13.

#### Global integration of local endogenous events is hampered in lower conscious states

We started this study by investigating whether endogenous spontaneous events occurring locally propagate differently depending on the level of consciousness, across healthy controls, MCS patients or patients with UWS. For that, we employed the *intrinsic ignition* measure<sup>24</sup>. The level of global integration for a subject is calculated as the average integration triggered by all endogenous events identified in their resting-state BOLD session, see Methods. As shown in **Fig. 1a**, the mean intrinsic ignition was lowest in UWS patients implying that the endogenous BOLD events lead to a lower network response than in healthy controls and in MCS patients (HC =  $0.81\pm0.01$ , UWS =  $0.78\pm0.01$ , MCS =  $0.79\pm0.01$ , HC vs. UWS t(45)=6.6, p<0.0001, HC vs. MCS t(57)=4.6, p<0.0001, MCS vs. UWS t(38)=2.3, p=0.012). It shall also be noted that the number of observed intrinsic events was lowest in UWS patients, intermediate in MCS patients and highest in healthy controls (HC =  $14.1\pm3.6$ , UWS =  $7.6\pm2.9$ , MCS =  $11.0\pm3.6$ , HC vs. UWS t(45)=5.9, p<0.0001, HC vs. MCS t(57)=3.3, p=0.0017, MCS vs. UWS t(38)=2.9, p=0.0048).

#### Shorter relaxation-time of BOLD signals in low levels of consciousness

Measuring time-scales from signals – i.e, multivariate time-series – can reveal changes in the underlying mechanisms controlling the local dynamics and determining their operating regime. Specifically, the autocovariance profile of the BOLD signal for each brain area measures the duration for which the signal is altered before going back to pre-event baseline activity<sup>28</sup>. Here, we measure the autocovariance time constant  $\tau$ , also called the relaxation time or memory depth in the literature. Large  $\tau$  implies a longer lingering effect of a signal after an event or perturbation before it decays, thus suggesting that the brain region might remain available for processing longer.

At a whole-brain level, averaging over the  $\tau_i$  for all regions in one subject, we found that  $\tau$  was smaller in UWS patients (1.96±0.38) than in healthy controls (2.72±0.35; t(45)=6.5,

p<0.0001) and MCS patients (2.70±0.58; t(38)=4.2, p<0.001), see **Fig. 1b**. Looking at the region-wise spatial distributions, we found that in healthy controls  $\tau_i$  is heterogeneously distributed showing a gradient with shorter time constants in subcortical areas ( $\tau_i \sim 1.5$  sec) and longer ( $\tau_i \sim 3.5$  sec) in the frontal and in the parietal areas, see **Supplementary Fig. 1**. Importantly, the diversity of relaxation times is lost in the UWS patients with  $\tau_i$  being homogeneously distributed and dominated by small values. Compared to healthy controls, the decrease of  $\tau_i$  in UWS patients was most predominant in the bilateral thalamus, right caudate, left hippocampus, parahippocampus, bilateral posterior, middle and anterior cingulate, insula, inferior, middle, superior and dorsolateral frontal areas, **Fig. 1c and Supplementary Table 2**. In the case of MCS patients the heterogeneity of  $\tau_i$  distribution was practically recovered, **Supplementary Fig. 1**. Compared to the healthy controls, in MCS patients  $\tau_i$  was lower only in the bilateral thalamus and left medial prefrontal cortex, see **Fig. 1d and Supplementary Table 2**.

So far, the results obtained for the intrinsic ignition and the distribution of relaxation time constants from the resting-state BOLD signals indicate a breakdown in the signals' spatiotemporal structure that involves reduced propagation and integration capabilities of endogenous events in DOC patients, especially in the UWS group. For the remaining of the paper we shift to model-based analyses.

#### Whole-brain effective connectivity shows altered causal interactions in DOC patients

In order to identify alterations to the causal relations between the brain regions, we estimated whole-brain effective connectivity (EC) from the resting-state BOLD for each subject. The estimation of EC considers a model of Gaussian noise diffusion – the multivariate Ornstein-Uhlenbeck – on top of the anatomical connectivity as the generative dynamics<sup>25,27</sup> in order to capture the origin of the fluctuations in the BOLD; see **Fig. 2a** and Methods for further details. In short, EC estimation consists of identifying the most likely causal interactions that give rise to the observed BOLD signals, fitting both the interaction strengths between all pairs of ROIs and the levels of noise to stimulate each ROI, **Fig. 2a**.

At the whole-brain level, averaging across all EC links, we found that the EC of the UWS patients was higher than for the healthy controls or in MCS patients (HC = 0.015±0.002, UWS = 0.019±0.004, MCS = 0.014±0.003; HC vs. UWS t(45)=-4.7, p<0.0001, MCS vs. UWS t(38)=-4.0, p<0.001). A closer inspection of the pair-wise EC values revealed the presence of links that either increased or decreased in the UWS patients in respect to the healthy controls, **Fig. 2b**. The UWS patients showed increased EC for connections between subcortical and cortical regions (thalamus, caudate and putamen), but decreased EC in connections spanning posterior (i.e., parietal, occipital) to frontal (i.e., temporal and frontal) regions as well as between midline posterior regions (parietal, occipital) and middle frontal regions. The MCS patients showed especially lower EC in interactions from posterior to frontal and temporal regions and

midline regions encompassing the middle prefrontal and posterior cortex and the thalamus, see **Fig. 2c**, including regions important for long range connectivity and overlapping with key areas of the Default Mode Network.

#### Altered spatiotemporal propagation of exogenous perturbations

Having identified changes in specific pair-wise EC connections for both UWS and MCS patients, the question is now how do those alterations affect the propagation of information in the brain. To answer this question, we perform an *in-silico* perturbational study to assess how exogenous perturbations, applied to individual ROIs, spread along the network. Considering the same generative dynamical model as for the EC estimation, the effect of regional perturbations on the rest of the network can be analytically estimated<sup>26</sup>, see Methods. The spatiotemporal responses of nodal perturbations are encoded into the temporal response matrices R(t). The evolution of response matrices for the three study cases are shown in **Fig. 3a**. Specifically, a pair-wise element  $R_{ij}(t)$  represents the temporal response of area j to a unit perturbation applied on area i at time t=0. This conditional, pair-wise response encompasses all network effects from i to j acting at different time scales.

Figure 3a illustrates how the patterns of responses are progressively reshaped over time for the three study groups – healthy controls, UWS patients and MCS patients. The global brain responses (sum over all pair-wise responses) are shown in Fig. 3b. As seen, the global responses undergo a transient peak short after the initial perturbations and then decay as the effects of the stimuli dilute with time and the system relaxes back to its stationary state. This relaxation is also observed by the homogenization of the response matrices at the longer latencies in Fig. 3a. The global response curves for controls and MCS groups follow quite a similar behavior, both peaking at 18.2±2.9 and 15.6± 3.7 seconds respectively and taking peak values 0.30±0.03 and 0.28± 0.05. In the UWS patients, however, the global response peaks sooner (10.6±2.9 sec) (HC vs. UWS t(45)=8.2, p<0.0001, HC vs. MCS t(57)=3.0, p=0.0036, MCS vs. UWS t(38)=4.3, p < 0.001) and displays a higher peak (0.34 $\pm$ 0.05) (HC vs. UWS t(45)=-3.3, p= 0.0019, MCS vs. UWS t(38)=-3.1, p=0.0031) than for the controls and MCS groups, but then it decays notably faster. Quantitatively, we found that the area-under-the-curve in the time spanning 60-200 sec (modeled time) significantly decreases for the UWS group (0.08±0.06) and MCS patients (0.15± 0.04) compared to healthy controls (0.18±0.02) (HC vs. UWS t(45)=7.1, p<0.0001, HC vs. MCS t(57)=2.9, p=0.005, MCS vs. UWS t(38)=3.6, p < 0.001).

#### Broadcasting and integrative capabilities of brain regions across states of consciousness

Since EC identifies the directed causal interactions between the brain regions, this allows us to study the input and output relations for each area in respect to the exogenous perturbations. The row sum of the response matrices  $\mathbf{R}(t)$  represent the broadcasting capacity of a region (i.e., the response that a perturbation in one region elicits on all other areas) and the columns describe

the integrative capacity of the brain region (i.e., how much is a region affected by the perturbations applied to all other areas)<sup>26</sup>.

Figure 4 reveals that the three levels of consciousness are characterized by distinct spatial distributions of 'broadcaster' and 'receiver' (i.e., integrator) areas. Notably, in UWS patients no brain region stands out either as broadcaster or as integrator, except for the thalamus displaying relatively large receiving capacity. In the healthy controls, we found several regions with both significant broadcasting and receiving capacity: the bilateral occipital, calcarine, lingual, cuneus, precuneus, superior and inferior parietal, right superior temporal. Significant broadcasting-only capacity was found in the bilateral inferior, middle and superior temporal, right parahippocampal, putamen, bilateral insula, inferior parietal, supramarginal, precuneus, middle cingulum and right inferior frontal areas. On the other hand, the bilateral posterior cingulate cortex (PCC), precuneus, supramarginal gyrus, thalamus, middle cingulate, left anterior cingulate and right inferior frontal cortex displayed significant receiving-only capacity. The MCS patient group was characterized by globally reduced broadcasting and receiving properties compared to healthy controls, however they showed a relatively preserved receiving and broadcasting of information within bilateral occipital, cuneus, left calcarine, bilateral superior and inferior parietal, supramarginal, precuneus and the right PCC. Additionally, they presented preserved broadcasting properties in the bilateral middle temporal, postcentral, right calcarine and left inferior frontal cortices, and preserved receiving capacity in the bilateral thalamus, see Fig.4 and **Supplementary Table 3.** 

We ended our perturbative analysis by comparing the region-wise group differences of the patients in respect to the healthy controls, which are shown in Fig. 5. Following the severely hampered broadcasting and integrating capacity in UWS, reduced information broadcasting in UWS patients as compared to healthy controls was especially notable at the bilateral hippocampus, parahippocampus, thalamus, caudate, amygdala, putamen, insula, inferior/middle temporal, temporal pole, right superior temporal, fusiform, lingual, calcarine, occipital, anterior cingulate, right inferior and middle frontal cortices. The notorious lack of broadcasting capacity of the subcortical regions evidences a reduced activity of the whole network. A profound reduction of the capacity to receive information in the UWS patients compared to the healthy controls was found at the bilateral precuneus, PCC, lingual, calcarine, fusiform, middle occipital, middle / anterior cingulum, inferior / superior parietal, supramarginal, middle temporal, inferior frontal cortices and the middle prefrontal cortex. These regions encompass primary visual and auditory areas, but also higher integration areas in the PCC that have an important hub function within the whole-brain network, see Fig.5 and Supplementary Table 4.

The MCS patients showed a less pronounced picture of impaired information in- and out-flows. Compared to healthy controls they showed a significant reduction in the potential to broadcast information in the bilateral thalamus, parahippocampus, left hippocampus, bilateral insula, inferior / middle temporal, right superior temporal, bilateral fusiform and lingual cortices.

A reduced capacity to receive information at the bilateral precuneus, PCC, cuneus, right lingual, calcarine, bilateral middle cingulum and right middle temporal cortices. Finally, compared to MCS patients, UWS patients showed additional significant reduction in receiving and broadcasting of information at the left precuneus, occipital cortex, temporal and right superior parietal, thus indicating that the information flow in these areas might be the most important contributors to conscious information processing.

In summary, our model-based analysis to estimate effective connectivity and to simulate *in-silico* the response to exogenous perturbations allowed us to identify specific directed pathways that are disrupted in patients with DOC, and thus reveal relevant for the propagation and processing of consciousness.

#### **III. Discussion**

In the present paper, we have studied the neural propagation of endogenous and exogenous perturbations in the brain using model-free and model-based analysis methods, applied to the problem of elucidating the mechanisms behind loss of consciousness due to acquired brain injury. The methods here employed add significant value to the dynamical approaches for two main reasons. First, they rely on simple observables – the resting-state fMRI – and thus they do not require the execution of experimental exogenous stimulation protocols. And second, unlike previous approaches, they allowed us to investigate the directional causal interactions between brain regions, thus elucidating alterations in the broadcasting and the integrating capacities of individual areas or pathways, between normal awake and unconscious patients. Indeed, our main finding is that we could identify two distinct malfunctioning neural circuits in patients with DOC: the posterior cortical regions fail to convey information, in conjunction with reduced broadcasting of information from subcortical, temporal, parietal and frontal regions. These results show that patients with prolonged disorders of consciousness lack of the capacity for the integration of events that would lead to conscious perception.

In healthy controls we found that the relaxation time constants associated with the resting-state BOLD signals display a gradient distribution with shorter relaxation times in subcortical areas and longer time constants in the frontal and in the parietal areas, **Supplementary Fig. 1a.** Accordingly, analysis of exogenous *in-silico* perturbations revealed that the broadcasting of information flow is predominant in a broad range of cognitive modules, including the hippocampus, parahippocampal, temporal, posterior and inferior frontal regions. This subcortical-cortical loop has been proposed to mediate the sensory information to be globally 'accessible' to other cognitive functions through feedforward and feedback loops by the global neuronal workspace theory, and only when access to all cognitive modules occurs, sensory content is elevated to conscious perception<sup>29–31</sup>. Although the activity in the posterior regions is highly influenced by perturbation, suggesting that they have a large cause-effect

capacity to receive and integrate the information flow, one of the key principles of conscious perception in integration information theory<sup>1,2,32</sup>.

Regarding the patients with unresponsive wakefulness syndrome (UWS), the results observed were very much altered. First, the propagation of endogenous events occurring in the resting-state BOLD rapidly decay avoiding their subsequent integration, Fig. 1a. This is corroborated by the fact that the spatial distribution of relaxation times fades away in the UWS patients, with all areas taking short relaxation times (Figs. 1b and c) and evidencing that local activity doesn't properly propagate along the network. Especially frontal, parietal and higher-order cortices which ensure sensory information processing, need longer time to integrate diverse information<sup>33,34</sup>. Second, effective connectivity is reduced overall but interestingly some effective subcortical-cortical connections were found to significantly increase, Fig. 2b. The propagation of *in-silico* exogenous perturbations showed a rapid and large response followed by a fast decay, Fig. 3, since the information fails to propagate along the network in a sustained manner. Such early hyper-response has been previously seen in UWS patients<sup>35</sup> and in loss of consciousness due to generalized epilepsy possibly caused by excess electrical discharges in the brain<sup>36</sup>. The mechanism for this hyper-response is yet to be elucidated but it could be either due to the network being dominated by short local recurrent loops, or due to a lack of inhibition as in unconscious anesthetized ferrets<sup>37</sup>. Finally, the regional specificities for broadcasting and receiving that were observed in the healthy controls are vanished for the UWS patients. Only the thalamus stands-out, as an area with significant receiving capacity thus probably allowing its gateway function between the body and the brain.

The patients in minimally conscious state (MCS) studied here underwent through a coma and a UWS phases after brain injury, but later regained partial consciousness and cognitive functionalities. All the results found for the MCS patients show light alterations to those in the healthy participants, as expected from their partial functional recovery. The spatial gradient of relaxation time constants from the resting-state BOLD is recovered, **Supplementary Fig. 1**, except shorter time constants were still found in the thalamus and the left medial preforntal cortex. Effective connectivity was in general slightly below than those observed in control subjects but a significant reduction was found remaining in fronto-parietal connections and in between temporal regions, **Fig. 2c**. Regional broadcasting and receiving capacities to *in-silico* perturbations displayed a recovered scenario, see **Fig. 4** and see **Supplementary Table 3**. It is clinically relevant to understand why or how MCS patients could partially recover from the unresponsive wakefulness state. In the light of our results, it seems that an increase in receiving and broadcasting of information at the left precuneus, occipital cortex, temporal and right superior parietal is instrumental for the recovery of conscious information processing, **Fig. 5** 

In comparison with healthy controls, the UWS patients showed a reduction of receiving information in posterior regions, which implies sensory information integration is impaired already at the level of sensory regions to the high-level hub regions of the Default Mode

Network (i.e. PCC and Precuneus). Indeed, the lack of receiving of information in the sensory areas hampers a stimulus or event to reach awareness, as integration of external inputs is a prerequisite for consciousness [Herbet, 2014]. Our results provide a mechanistic explanation for how the ability to receive information in sensory and DMN hub regions alters cerebral information processing, which might be at the essence for the structural and functional anomalies in UWS patients. Although it is known that the PCC and Precuneus have decreased structural, functional and metabolism<sup>7,12,16,17,38</sup>, our results provide a mechanistic explanation that the ability to receive information in sensory and DMN hub regions is reduced in UWS patients. On the other hand, broadcasting was reduced in the subcortical regions (i.e., thalamus, caudate, putamen) and regions involved in higher cognitive function (i.e., temporo-parietal, anterior cingulate and frontal regions). This is aligned with the mesocircuit hypothesis<sup>39</sup> which states that the feedforward connections between these regions play a key role in reaching levels of (cortical) activity that support the stimuli to access consciousness processing. This has been recently confirmed in macaques using invasive-EEG, showing that integration at the thalamus, caudate, putamen and parietal cortex is a hallmark of conscious states<sup>40</sup>. Our findings unravel that human consciousness also relies on the broadcasting capacities from the thalamus, caudate and putamen leads to support the transmission of activity, functional integration and recurrent activity between subcortical and cortical neurons, all of which is lost in UWS patients. Interestingly, we also noted a decrease in receiving and broadcasting capacities in the bilateral temporal areas for DOC patients. Recent studies also noted altered structural<sup>41</sup> and functional loss in the temporal area in DOC patients<sup>7,13</sup>. To date, there is limited explanation for the involvement of the temporal cortex in consciousness. We speculate that a lesser involvement of the temporal areas in the information pathways could impede self-awareness and memory.

Our *in-silico* pertubational study revealed transient global changes to the dissemination of information that are similar to those we observed from the integration of endogenous events with the intrinsic ignition. In DOC patients, and especially in UWS patients, the brain's cause-effect capacity to respond is significantly lower than during normal wakefulness in healthy subjects. It seems that the observed spatiotemporal alterations of local event processing also hamper global integration and whole brain neural responses; as observed both after *in-silico* and endogenous perturbations. These results are in line with empirical studies using TMS, in which the recruitment of global neural activity after perturbation, both in space and time, has been found to be reduced during deep sleep, anesthesia and in DOC<sup>20,21</sup>. In conclusion, the cerebral capacity of propagation and integration of local, naturally occurring events into the entire network is affected by reduced states of consciousness and shares similarities with both information integration theory<sup>1,2</sup> and global neuronal workspace theory<sup>30,31</sup>. Although these theories have distinct concepts of consciousness, our results suggest that they might represent two sides of the same coin<sup>42-44</sup>.

#### IV. Methods

#### **Participants**

We included subjects with a pathological reduction or loss of consciousness after severe brain injury, so called disorders of consciousness (DOCs), as well as healthy control (HC) volunteers. Written informed consent was obtained from all HC participants and the legal representative of DOC patients for participation in the study. The local ethics committee from the University Hospital of Liège (Belgium) approved the study. Forty adult DOC patients, in which 26 in minimally conscious state (MCS) (7 females, age range 23-73 years; mean age  $\pm$  SD, 41  $\pm$  13 years) and 14 with the unresponsive wakefulness syndrome (UWS) (7 females, age range 20-74 years; mean age  $\pm$  SD, 49  $\pm$  16 years) and 33 age and gender matched HC (13 females, age range 19-72 years; mean age  $\pm$  SD, 40  $\pm$  14 years) without premorbid neurological problems were included. The diagnosis of the DOC patients was confirmed through two gold standard approaches. The first is the repeated behavioral assessment using the Coma Recovery Scale-Revised (CRS-R) by trained clinicians and the second used Fluoro-deoxyglucose Positron Emission Tomography (FDG-PET) neuroimaging as an objective test to complement behavioral assessment according to the procedure described by Stender et al. 12. Patients for whom these two diagnostic approaches disagreed were excluded from further analysis. Patients were behaviorally diagnosed through the best of at least 5 CRS-R assessments, evaluating auditory, visual, motor, oromotor function, communication and arousal<sup>45</sup>. Patient specific clinical information is presented in Supplementary Table 1.

#### **MRI Data Acquisition**

Structural (T1 and Diffusion Weighted Imaging, DWI) and functional MRI (fMRI) data was acquired on a Siemens 3T Trio scanner. 3D T1-weighted MP-RAGE images (120 transversal slices, repetition time = 2300 ms, voxel size = 1.0 x 1.0 x 1.2 mm3, flip angle = 9°, field of view = 256 x 256 mm2) were acquired prior the 10 minutes of BOLD fMRI resting state (i.e., task free) acquisition (EPI, gradient echo, volumes = 300, repetition time = 2000 ms, echo time = 30 ms, flip angle = 78°, voxel size = 3 x 3 x 3 mm3, field of view = 192×192 mm2, 32 transversal slices). Last, diffusion weighted MRI was acquired in 64 directions (b-value =1,000 s/mm2, voxel size = 1.8x1.8x3.3 mm3, field of view 230x230 mm2, repetition time 5,700 ms, echo time 87 ms, 45 transverse slices, 128x128 voxel matrix) preceded by a single unweighted image(b0). The DWI was acquired twice.

#### MRI data preprocessing

Preprocessing was performed using MELODIC (Multivariate Exploratory Linear Optimized Decomposition into Independent Components) version 3.14, which is part of FMRIB's Software

Library (FSL, <a href="http://fsl.fmrib.ox.ac.uk/fsl">http://fsl.fmrib.ox.ac.uk/fsl</a>). The preprocessing consisted of the following steps: the first five functional images were discarded to reduce scanner inhomogeneity, motion correction was performed using MCFLIRT, non-brain tissue was removed using BET, intensity was normalized, temporal band-pass filtering with sigma 100 sec was performed, spatial smoothing was applied using a 5mm FWHM Gaussian kernel, rigid-body registration and single-session ICA with automatic dimensionality. Then noise components and lesion-driven artifacts (e.g., head movement, metal, and physiological noise artifacts) were manually regressed out for each subject. Specifically, FSLeyes in Melodic mode was used to identify the single-subject Independent Components (ICs) into "good" for cerebral signal, "bad" for noise or injury-driven artifacts, and "unknown" for ambiguous components. Each component was evaluated based on the spatial map, the time series, and the temporal power spectrum<sup>46</sup>. FIX was applied with default parameters to remove bad and lesion-driven artifacts components<sup>46</sup>. Subsequently, the Shen et al (2015) functional resting state atlas (without cerebellum) was used for parcellation to obtain the BOLD time series of the 214 cortical and subcortical brain areas in each individual's native EPI space<sup>47</sup>. The cleaned functional data were co-registered to the T1-weighted structural image using FLIRT. Then, the T1-weighted image was co-registered to the standard MNI space by using FLIRT (12 DOF), and FNIRT<sup>48</sup>. This transformation matrix was inverted and applied to warp the resting-state atlas from MNI space to the single-subject functional data. Finally, the time series for each of the 214 brain areas were extracted using custom-made Matlab scripts using 'fslmaths' and 'fslmeants'.

#### **Structural Connectivity Matrix**

We computed an average whole-brain structural connectivity matrix from all healthy participants as described in our previous study<sup>17</sup>. Briefly, the b0 image in native diffusion space was co-registered to the T1 structural image using FLIRT<sup>48</sup>. Next, the T1 structural image was co-register to the MNI space by using FLIRT and FNIRT<sup>48</sup>. The resulting transformations were inverted and applied to warp the resting-state atlas from MNI space to the native diffusion space using a nearest-neighbor interpolation method. Then, analysis of diffusion images was performed using the FMRIB's Diffusion Toolbox (FDT) <a href="https://www.fmrib.ox.ac.uk/fsl">www.fmrib.ox.ac.uk/fsl</a>. Brain Extraction Tool was computed, and eddy current distortions and head motion were corrected using eddy correct tool<sup>49</sup>. Furthermore, the gradient matrix was reoriented to correct for the subject. Then, Crossing Fibres were modeled using the default BEDPOSTX parameters, and the probability of multi-fibre orientations was calculated to improve the sensitivity of non-dominant fibre populations<sup>49</sup>. Next, probabilistic tractography was calculated in native diffusion space using the default PROB- TRACKX settings<sup>49</sup> to estimate the connectivity probability of each brain area to each of the other 213 brain areas. Subsequently, to obtain the structural probability matrix, the value of each brain area was divided by its corresponding number of generated tracts. Given that

probabilistic tractography does not capture fiber directionality, the SCnp matrix was symmetrized by computing their transpose matrix SCpn and averaging both matrices.

Finally a group structural connectivity (SC) mask was obtained by averaging the all HC subjects' SC matrix and applying a threshold of 80% to maintain the top 20% of strongest connections to binarize the SC. This SC mask was used to constrain the functional connectivity matrix for the whole brain EC computation.

#### **Data Analysis**

#### **Intrinsic Ignition**

Intrinsic ignition describes the influence of local endogenous events – spontaneously occurring – over the whole-brain network and their subsequent integration<sup>24</sup>. See Deco et al. (2017) for details<sup>24</sup>. Local events are defined as significantly large fluctuations taking place in the resting-state BOLD signals. First, the BOLD signals were transformed into z-scores,  $z_i(t)$ , and binarized by imposing a threshold  $\theta$  such that the binary signal takes value 1 if  $z_i(t) > \theta$  and 0 otherwise [Tagliazucchi, 2012]. Here we considered  $\theta = 2$  standard deviation. For every endogenous event identified, we calculated the subsequent integration of the event by the network. Therefore, the phase-locking matrix of the network was calculated for a time-window of 4TR post-event. Phase-locking matrices account for the instantaneous level of pairwise synchronization, see below. Integration is then calculated as the area-under-the curve delimited by the size of the largest component in the binarised phase-locking matrix, for all thresholds from 1 to 0. The mean intrinsic ignition is finally calculated as the average integration triggered by all events occurring in the resting-state BOLD for a subject. Higher values of intrinsic ignition correspond to rich and flexible brain dynamics whereas lower values correspond to poor and rigid, structurally driven brain dynamics.

#### **Phase-locking matrices**

The instantaneous level of pairwise synchronization was calculated by the phase-locking value between two brain regions. First, the BOLD signals were filtered within a narrowband of 0.04-0.07 Hz. Then the instantaneous phases  $\varphi_k(t)$  were computed using the Hilbert transform for each BOLD signal individually. This yields the associated analytical signal which represents a narrowband signal s(t) in the time domain as a rotating vector with an instantaneous phase  $\varphi(t)$  and an instantaneous amplitude, A(t). That is,  $s(t) = A(t)\cos(\varphi(t))$ . Given the instantaneous phases  $\varphi_i(t)$  and  $\varphi_k(t)$  calculated two brain regions from their corresponding

BOLD signals, the pairwise synchronization  $P_{jk}(t)$  was defined as the cosine similarity of the two phases:

$$P_{ik}(t) = cos(|\varphi_i(t) - \varphi_k(t)|).$$

Thus,  $P_{jk}(t) = 1$  when the two regions are in phase,  $P_{jk}(t) = 0$  when they are orthogonal and  $P_{jk}(t) = -1$  when they are in anti-phase.

#### Relaxation time constants $(\tau)$

In order to obtain information about the operating regime of brain regions, we measured the relaxation time constant  $\tau$  from the BOLD signals. Specifically, we measured the time constant of the autocovariance for each brain region individually, using time shifts from 0 to 1 TRs. Given that  $Q_{ij}^{0}$  and  $Q_{ij}^{1}$  are the zero-lag and 1TR-lag covariance matrices from the empirical BOLD, the time constants  $\tau_{i}$  are calculated as:

$$\tau_i = -\frac{1}{a(v_i|u)},$$

where  $a(v_i|u)$  corresponds to the slope of the linear regression of  $v_i = [log(Q_{ij}^0), log(Q_{ij}^1))]$  by u=[0,1]. Apart from the information extracted out of the regional time constants, the calculated  $\tau_i$  were also employed to inform the estimation of effective connectivity.

#### **Estimation of effective connectivity**

We estimated whole-brain effective connectivity from the resting-state BOLD signals considering the multivariate Ornstein-Uhlenbeck (MOU) process as the generative dynamical model of the BOLD<sup>25,27</sup>. See Gilson et al (2016) for details<sup>25</sup>. The MOU is a model of Gaussian noise diffusion on a network that has been popular to study the relation between the anatomical connectivity and the whole-brain network dynamics<sup>25,50</sup>. Given a structural connectivity matrix **A**, the MOU is defined as:

$$dx_{i} = \left(-\frac{x_{i}}{\tau_{i}} + A_{ij} x_{j}\right) dt + dB_{i},$$

where  $x_i$  corresponds to the activity (BOLD signal) of a brain region i,  $\tau_i$  is the time constant characterizing the exponential decay and dB is a colored noise given by a covariance matrix  $\Sigma$ .

The zero-lag  $Q^0$  and 1TR-lag  $Q^1$  covariance matrices of this model can be analytically calculated. The model is thus fitted to empirical data by a Lyapunov optimization procedure such that the distance between the empirical and the estimated  $Q^0$  and  $Q^1$  covariances is minimized<sup>25</sup>. The optimization process was initialized considering A as the binarized structural connectivity matrix in order to restrict the optimization to links identified via diffusion imaging. The estimations were performed using the pyMOU python package.

#### In-silico exogenous perturbational analysis

Considering the MOU as the generative dynamical model for the diffusion of noise in a network, the network responses to local perturbations can be analytically estimated; see Gilson et al.  $(2019)^{26}$ . In particular, we characterise the Green function of the MOU for a given connectivity matrix A. The Green function describes the temporal network responses at times t > 0, due to a unit perturbation applied at a given brain region i at time t = 0. For the MOU process, the spatiotemporal responses are given by:

$$R(t) = ||J^0||(e^{Jt} - e^{Jt}),$$

where J is the Jaccobian of the MOU process,  $J_{ij} = -\frac{\delta_i}{\tau} + A_{ij}$  and  $J_{ij}^0 = -\delta_{ij}/\tau$ , is the Jaccobian associated to the leakage term alone, characterising the decay rate of the system.  $||J^0||$  is a normalization term to make analysis across networks comparable. The response matrices R(t) encode the spatio-temporal responses to nodal perturbations. In other words, a pair-wise element  $R_{ij}(t)$  represents the temporal response of area j to a unit perturbation applied on area i at time t=0. This conditional, pair-wise response encompases all network effects from i to j acting at different time scales. Note that in Ref. Gilson et al.,  $(2019)^{26}$  the network responses R(t) were referred to as 'dynamic communicability C(t)'. Here we adopted a nomenclature that is clearer and conceptually more precise in order to facilitate the interpretation of results.

As in Ref. Gilson et al.  $(2019)^{26}$ , in the present study, the connectivity matrices **A** are the effective connectivity matrices previously estimated for each subject. Hence, the propagation of responses to exogenous perturbations are constrained by the strength of the directed, causal interactions between every pair of brain regions.

The global network response r(t) is the sum of all pairwise responses at each time point:

$$r(t) = \sum_{i,j=1}^{N} R_{ij}(t),$$

accounting for the total excitability of a network to exogenous perturbation.

Since effective connectivity estimates the directed, causal pairwise interactions between brain regions, and the response matrices R(t) are constrained upon effective connectivity, R(t) account for the asymmetric interactions between brain regions. The broadcasting capacity of a region i is calculated as the sum of all responses exerted by region i on the rest of brain areas, and the receiving or integration capacity is given by the sum of responses elicited on region i, by the perturbations at all areas. That is, the broadcasting and receiving capacities of a node are calculated as the row and column sum of the response matrices R(t) at each time point t:

Broadcasting capacity: 
$$r_i^+(t) = \sum_{j=1}^N R_{ij}(t)$$
, and

Receiving capacity: 
$$r_i(t) = \sum_{j=1}^{N} R_{ji}(t)$$
.

Note that in Refs. Gilson et al. (2019)<sup>26</sup> the broadcasting and receiving capacities are referred to as out-communicability and in-communicability respectively.

#### Statistical analysis

For the model free measure, two-sample t-tests were used to assess group differences in global Intrinsic Ignition and relaxation time constant  $\tau$  at the whole brain level (Bonferroni correction for 3 groups). Then we investigated local between group differences in regional relaxation time constant  $\tau$  using two-sample t-tests with Bonferroni correction (p<0.05) accounting for the number of regions (i.e., N = 214).

For the model free measure, first, we assess between group differences in whole-brain total (i.e., receiving and broadcasting) communicability. An ANOVA with Tukey post hoc comparison, Bonferroni corrected for 200 timepoints of integration, was employed. Second, to investigate local broadcasting and receiving properties, we considered the area under the receiving and broadcasting curves separately for every brain region. We identified regions with relatively high communicability within groups (i.e., the different states within the DOC) with a one-sample t-test with FDR <0.05 correction for the number of regions (i.e., N = 214). Last, between group differences in regional receiving and broadcasting information were assessed with two-sample t-tests. Between group statistics were corrected using Bonferroni correction (p<0.05) accounting for the number of regions (i.e., N = 214).

#### **Code and Data Availability:**

After the acceptance of the manuscript, the code used for this study will be available at: <a href="https://github.com/RajanikantPanda/Ignition">https://github.com/RajanikantPanda/Ignition</a> and Information flow for DOC

The multimodal neuroimaging data used in this experiment are available upon request with appropriate procedures.

#### **Authors' contributions**

RP, ALG, JA, GZL, MG, GD and SL designed research. JA and GZL supervised the research. JA, AT, CM, OG and the Coma Science Group Collaborators acquired the data. RP, ALG and AE preprocessed the data. RP, ALG, JA and GF analyzed the data. MG and GZL designed the computational model and optimized the code as per this research study. RP, JA, ALG and GZL wrote the manuscript. All authors interpreted the results and contributed to the editing of the manuscript.

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### **Figures and Tables**

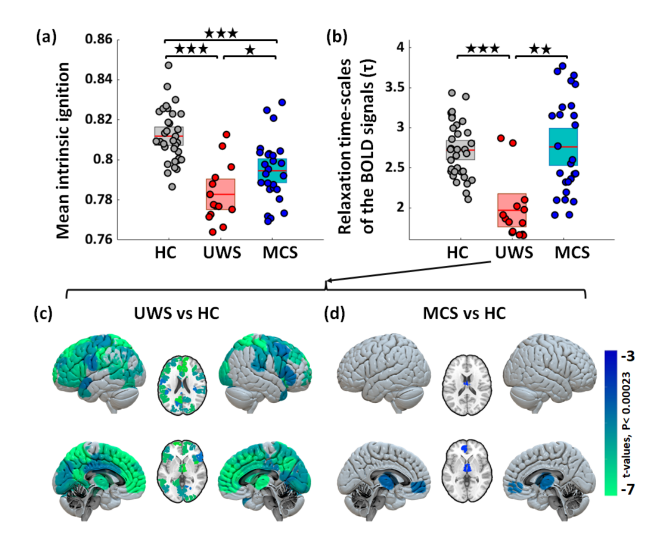


Figure 1. Changes in endogenous properties from resting-state BOLD signals. Healthy controls (HC), unresponsive wakefulness syndrome (UWS) and minimally conscious state (MCS) (a) Comparison of mean intrinsic ignition for the three groups, illustrating the reduced capacity to integrate endogenous spontaneous events in patients with DOC. (b) Relaxation time-scales of the BOLD signals ( $\tau$ ) at the whole-brain level shows significant reductions in UWS and MCS patients compared to HC. Stars reflect the Bonferroni corrected (for three groups) significance levels (\*=p-value<0.016; \*\*= p-value<0.001; \*\*\*= p-value<0.0001). (c,d) Maps of significant differences in regional distributions of  $\tau$  between patients and controls. The color bar represents the t-values of significant between-group differences (Bonferroni corrected for 214 tests, p-value=0.05).

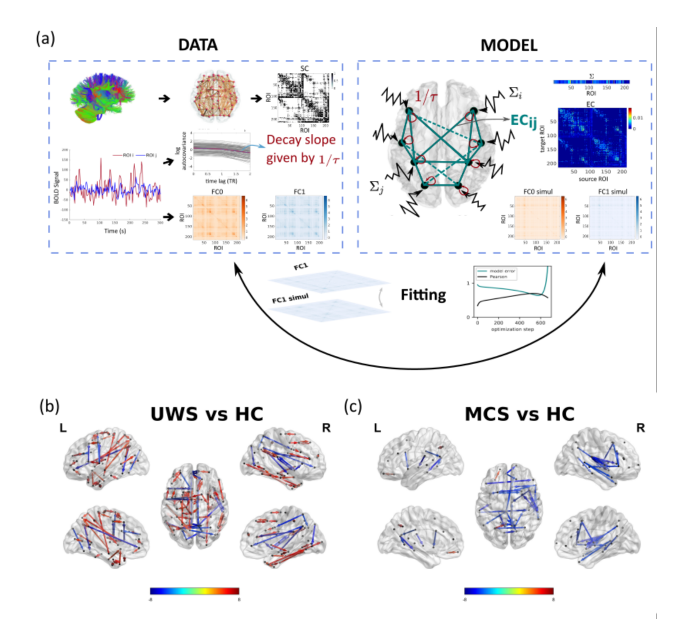


Figure 2. Comparison of effective connectivity (EC) between healthy controls and patients.

(a) Schematic representation of the fitting procedure leading to estimation of EC. Considering a model of noise diffusion – the multivariate Ornstein-Uhlenbeck process – the whole-brain network model is constrained using structural connectivity obtained from diffusion imaging and then fitted to reproduce the empirical resting-state data. In particular, to fit the zero-lag and 1TR-lag covariance matrices (FC0 and FC1), and the regional noise level  $\Sigma_i$ . (b, c) Maps of significantly different EC connections between patients and controls. UWS patients show connections with both decreased and increased EC (decreased in fronto-temporal, frontal-parietal and midline regions; increased in subcortical and wide cortical areas). MCS patients show decreased EC in fronto-temporal and interhemispheric midline connections. Blue and red arrows

indicate lower and higher EC respectively in patients as compared to HC subjects. The directional connections in the glass brain represents connections with significant between-group differences (Bonferroni corrected for 11395 tests, p-value < 0.05) are represented.

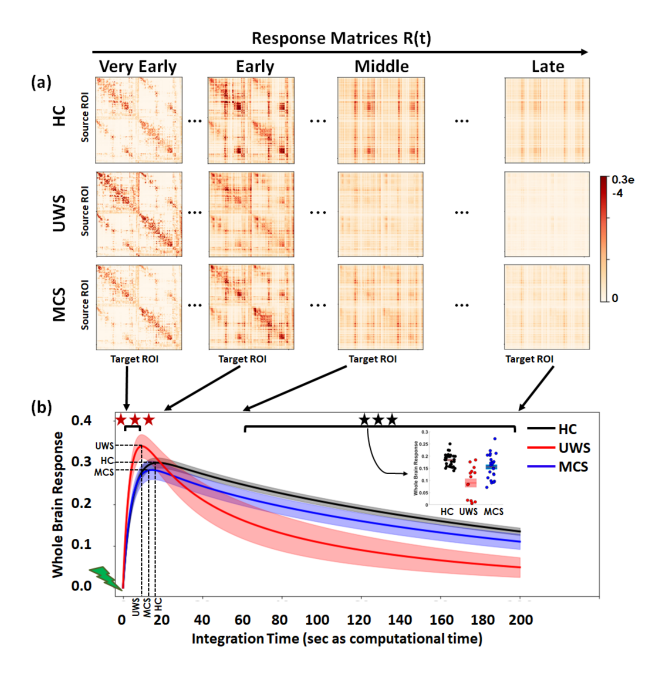
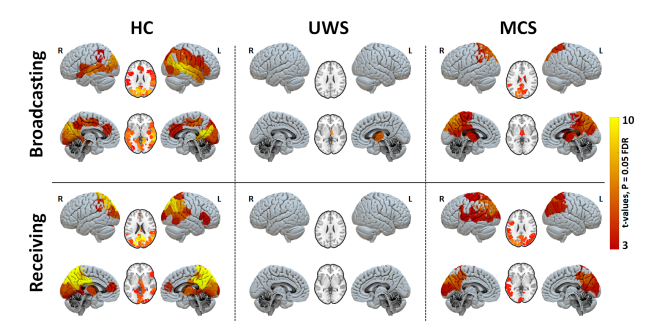


Figure 3. *In-silico* propagation of exogenous perturbations. (a) Temporal evolution of the response matrices R(t) for healthy controls (HC) and patients (unresponsive wakefulness syndrome, UWS; minimally conscious state, MCS) at different times (early = 2 sec, middle = 20 sec, late 60 sec and very late = 200 sec). Matrix elements  $R_{ij}(t)$  represent the conditional response at region j due to a unit perturbation applied at region i at time t = 0. Note that here time

corresponds to the arbitrary simulation time after the *in-silico* perturbation is applied and thus it does not correspond to actual time, although the time-constants governing the evolution were estimated from the BOLD signals. The colorbar represents the relative strength of the response between brain regions (unitless). (b) Whole brain response curves for the three study cases reflecting the sum of all pair-wise responses at each time point post-stimulus. Shaded areas represent the 95% confidence intervals across subjects. Black stars indicate a difference in global responses between all three groups (Bonferroni corrected for 100 tests/time points, p-value=0.05). Red stars indicate the early epoch during which UWS patients display a larger response than HC and MCS (Bonferroni corrected for 100 tests/time points, p-value=0.05). Inset: Area-under-the-curve for the three global response curves in the time range t = 60 - 200 sec, quantifying the differences across the three groups.



**Figure 4. Region-wise broadcasting and receiving capacities due to exogenous perturbations.** Maps of significantly large broadcasting and receiving capacities for the three study groups (healthy controls, HC; unresponsive wakefulness syndrome, UWS; and minimally conscious state, MCS). The color code represents the t-values. Only regions with significantly high values are presented in each case (FDR corrected p-values <0.05 for 214 tests (ROIs)).

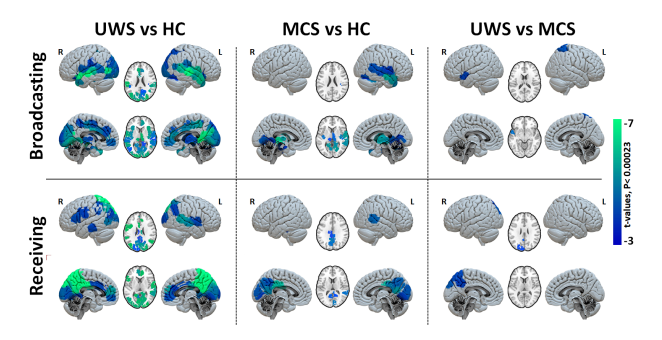
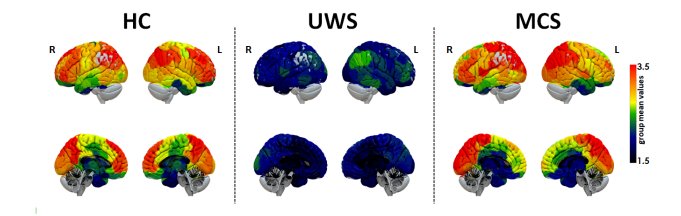
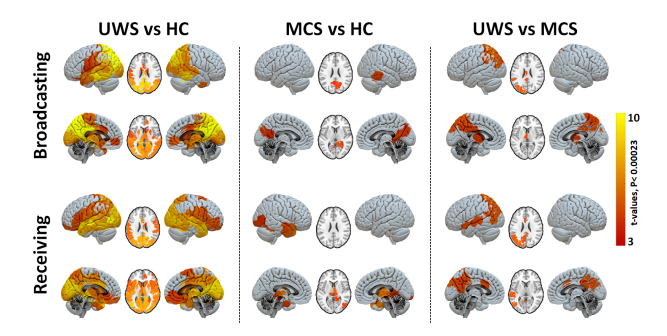


Figure 5. Group comparison of regional broadcasting and receiving capacities after exogenous perturbations. Maps of regional contrasts in broadcasting and receiving capacity between study groups. Color bar represents the t-values for regions with significant between-group differences (Bonferroni corrected p-values <0.05 for 214 tests (ROIs)).

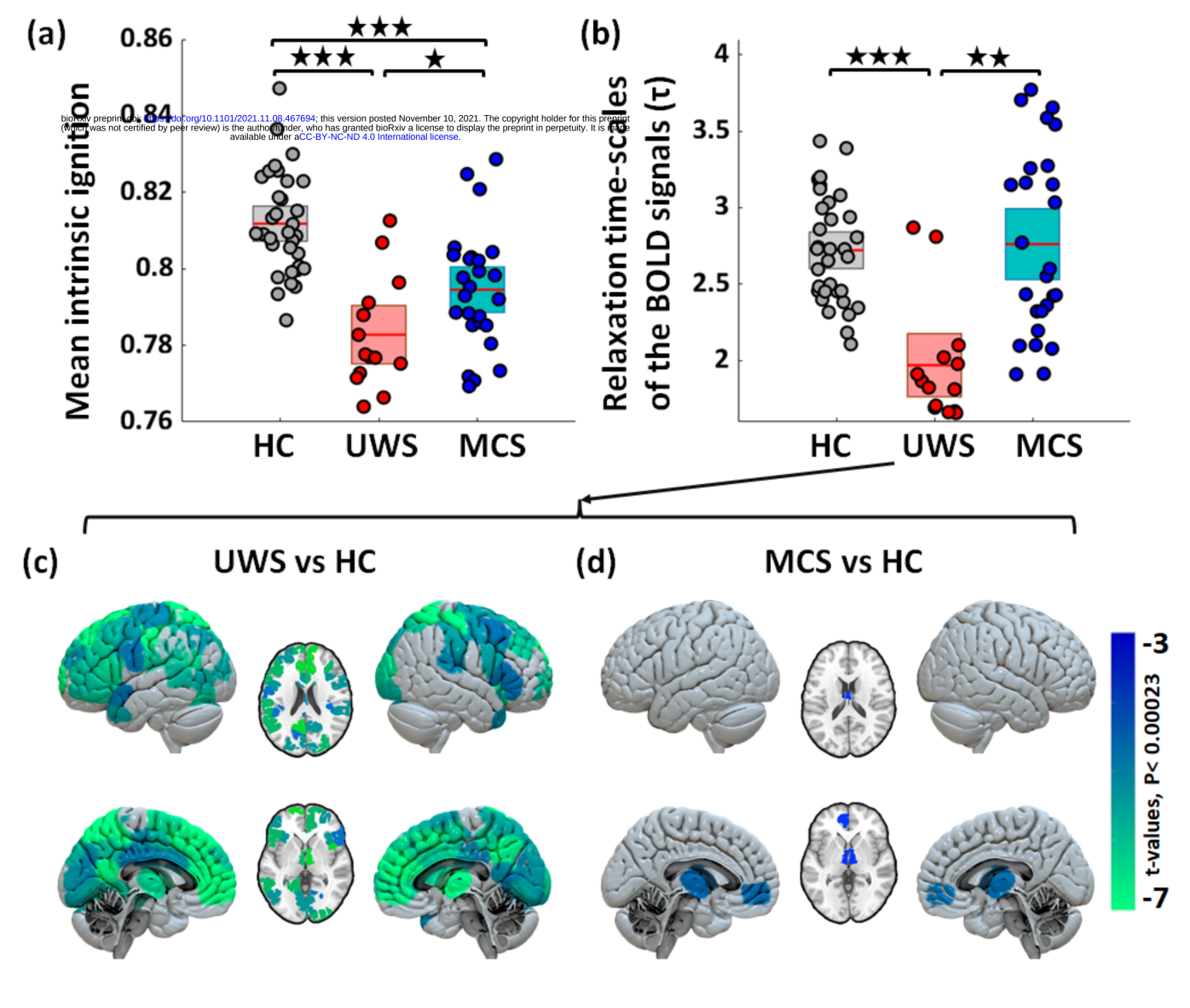
#### **Supplementary Material**

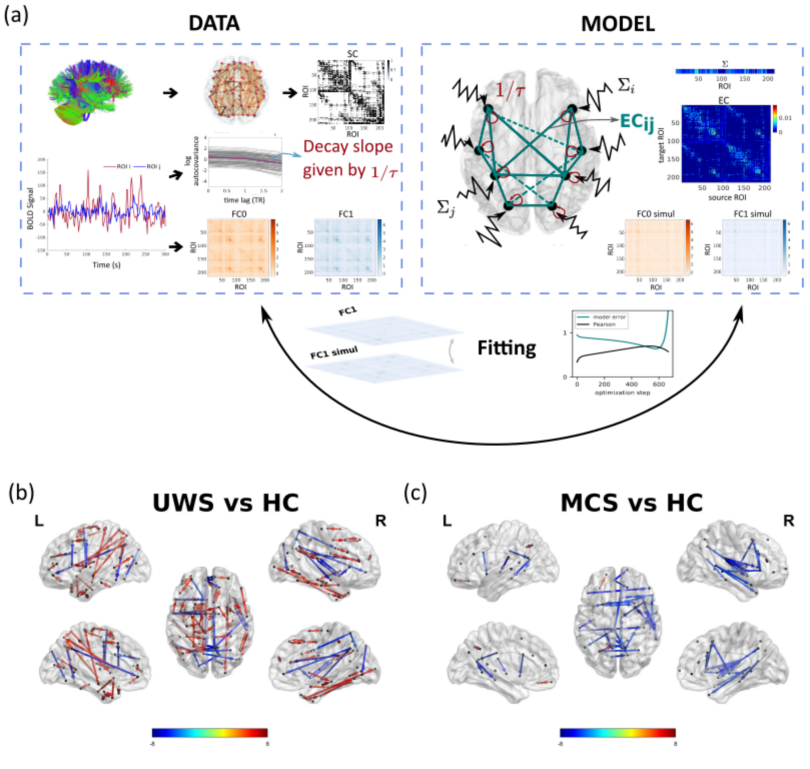


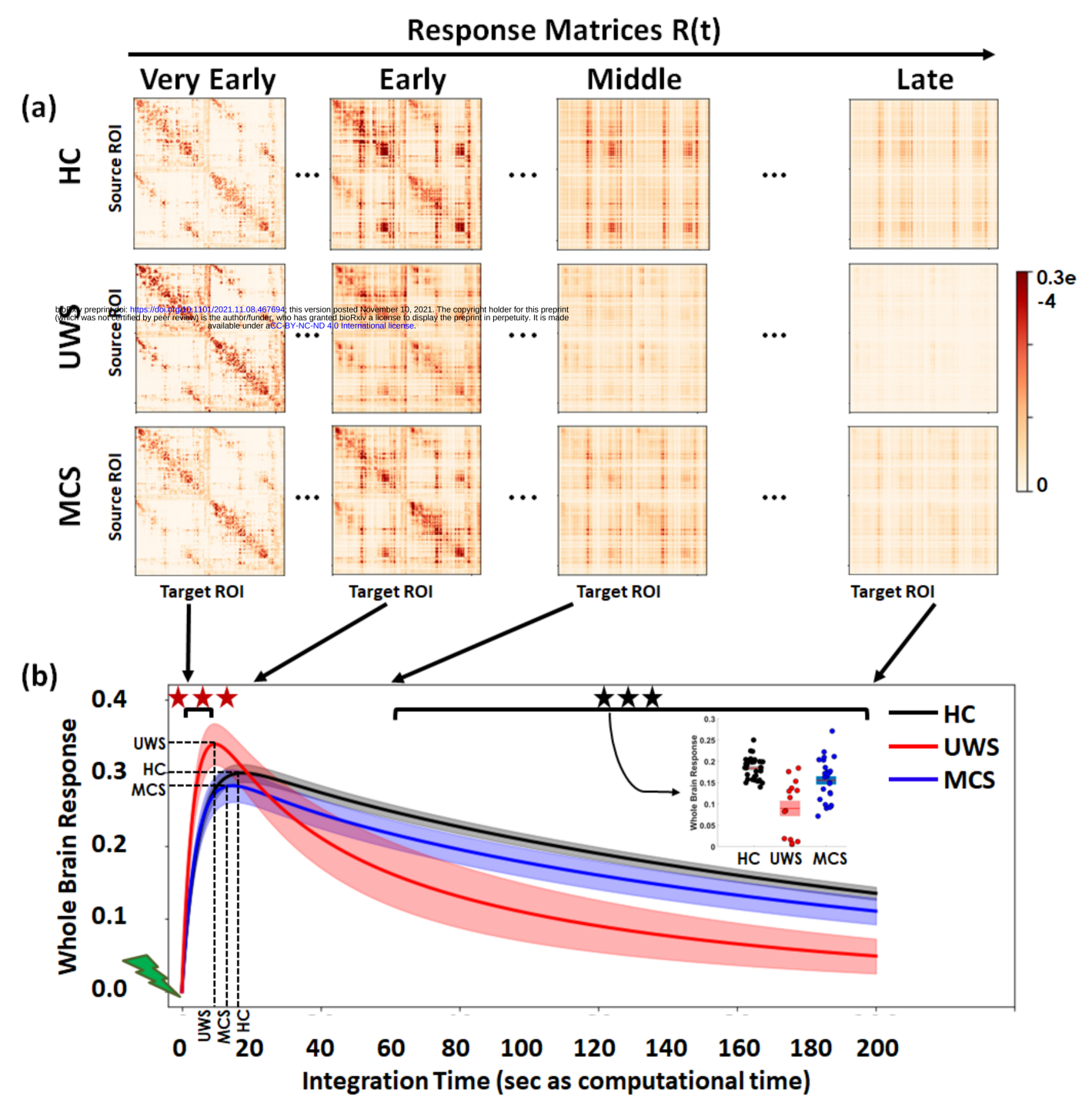
Supplementary Figure 1. Spatial maps showing the regional distribution of relaxation time constants ( $\tau$ ) as calculated for empirical BOLD signals for each area. Healthy controls display a spatial heterogeneous distribution of  $\tau$ , while UWS patients are characterized by short time constants overall. The spatial distribution of regional  $\tau$  is very much recovered in MCS patients.



Supplementary Figure 2. Brain regions' hyper-response (higher growth with sudden decay) differ in (a) UWS patients compared to HC (b) MCS patients compared to HC (c) UWS compared to MCS patients for receiving and broadcasting capacities. Color bar represents the t-values for regions with significant between-group differences (Bonferroni corrected for 214 tests).







## **UWS vs HC**

# MCS vs HC

## **UWS vs MCS**

