Signature of consciousness in brain-wide synchronization patterns of monkey and human fMRI signals

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A B S T R A C T

During the sleep-wake cycle, the brain undergoes profound dynamical changes, which manifest subjectively as transitions between conscious experience and unconsciousness. Yet, neurophysiological signatures that can objectively distinguish different consciousness states based are scarce. Here, we show that differences in the level of brain-wide signals can reliably distinguish different stages of sleep and anesthesia from the awake state in human and monkey fMRI resting state data. Moreover, a whole-brain computational model can faithfully reproduce changes in global synchronization and other metrics such as functional connectivity, structure-function relationship, integration and segregation across vigilance states. We demonstrate that the awake brain is close to a Hopf bifurcation, which naturally coincides with the emergence of globally correlated fMRI signals. Furthermore, simulating lesions of individual brain areas highlights the importance of connectivity hubs in the posterior brain and subcortical nuclei for maintaining the model in the awake state, as predicted by graph-theoretical analyses of structural data.

1. Introduction

Daily life is divided into periods of consciousness with subjective and reportable experience, which are repeatedly replaced by epochs with absence of consciousness (Koch, 2018). These different states of consciousness and their transition are generated and tightly regulated by sophisticated physiological mechanisms residing in the cortex and subcortical structures (e.g. the thalamus, brainstem nuclei and basal forebrain), which orchestrate the sleep-wake cycle (Scannell et al., 2017). In addition, pharmacological interventions during anesthesia (Alkire et al., 2008) can interfere with the physiology of consciousness and poise the brain to unconscious dynamics. At the same time, the precise organization of the anatomical connectivity plays a pivotal role in the generation of conscious experience, as evidenced by loss of consciousness after severe brain damage in patients with coma or unresponsive wakefulness syndrome (Gosselies et al., 2014). Understanding how such interplay between the anatomical structure and the physiology of neuronal networks generates different levels of consciousness remains a major challenge in neuroscience research.

To open a window on brain functioning during different levels of consciousness, researchers have traditionally gained insights from neuronal oscillations that can be captured by recording macroscopic electromagnetic activity macroscopically (EEG, MEG) (Stevner et al., 2019), mesoscopically (Nir et al., 2011) or by single cell recordings (Steriade et al., 1993). These studies delineate sleep into REM and
non-REM sleep, which is separated into three different stages (N1, N2, N3), each with a distinct electrophysiological profile (Hobson and Pace-Schott, 2002).

More recently, electrophysiological findings have been complemented by fMRI studies, which focus on the study of functional relationships between brain areas, as measured e.g. by BOLD signal correlations (Masbour and Hudetz, 2018). Such functional connectivity (FC) can be contrasted with the underlying structural connectome (SC) to examine how the variety of pathways, constrained by the anatomical connectivity, is expressed in the FC during different consciousness levels. Overall, the strength of the functional connectivity is largely reduced during unconscious states, even though specific resting state networks remain intact (Bovereux et al., 2010; Spoormaker et al., 2010). Moreover, during loss of consciousness structural constraints are more rigidly expressed in the FC as opposed to the awake state (Barttfeld et al., 2015; Tagliazucchi et al., 2016b).

In addition, fMRI signals tend to display an organized, wave-like flow across the entire brain with high integration in the awake state, while propagation remains confined to resting state networks with high modularity scores during sleep (Boly et al., 2012; Mitra et al., 2015b; Tagliazucchi et al., 2013). Furthermore, it was shown that these correlation metrics are dynamic and show fluctuations across time (Betzel et al., 2016; Fukushima et al., 2018a, 2017; Shine et al., 2016), which is informative about the level of consciousness (Barttfeld et al., 2015; Demertzi et al., 2019; Haimovici et al., 2017; Uhrig et al., 2018).

Here, we study how large-scale synchronization of the resting state fMRI signal changes across conscious and unconscious states (sleep stages N1, N2, N3 and anesthesia), and whether it can serve as an objective marker for the state of consciousness. Coherence across large parts of the brain has been observed previously in fMRI recordings of humans (Deco et al., 2017b) and monkeys (Liu et al., 2018a; Turchi et al., 2018), and is accompanied by activity changes in neuromodulatory nuclei. We investigate how various degrees of shared large-scale fluctuations relate to metrics such as FC, SC-FC relationship, integration and modularity. The empirical findings are then contrasted with simulations from a whole-brain model and we further examine whether it can reproduce the pattern of global fluctuations seen across different consciousness conditions. Finally, the impact of brain lesions on global dynamics is studied via in-silico modeling.

2. Results

In this study, we analyzed brain activity of different vigilance states in two different fMRI datasets, which contain both conscious and unconscious experimental conditions. One dataset was recorded in humans (N = 18) and comprises awake, resting state activity together with different stages of sleep (N1-N3) with a parcellation of 90 brain areas (Supplementary Table 1). The other dataset stems from monkey recordings (N = 3) during wakeful resting and two different levels of propofol anesthesia (moderate and deep) with a spatial resolution of 432 regions (Supplementary Table 2). We searched for a physiological metric that can differentiate conscious from unconscious states and can be explained mechanistically by a whole-brain computational model. Our hypothesis revolved around differences in brain-wide synchronization patterns, indicating fluctuations of fMRI activity shared across many brain areas (Scholvinck et al., 2010). To visualize synchronization, we first transformed the fMRI signals into a point process by representing all positive fMRI peaks as discrete events (Tagliazucchi et al., 2012) (Fig. 1A). This approach takes the entire signal across all brain areas into account, irrespective of inter-area variations of oscillation power in different frequency bands. The raster plots revealed pronounced differences between the awake and unconscious states (Fig. 1B). Awake brain dynamics was characterized by frequent large-scale population events that spanned large parts of the brain in both human and monkey resting state data (e.g. Fig. 1B top left, red rectangle). Interspersed were periods with local synchronization and absence of brain wide activity. In contrast, during unconscious states large-scale synchronization was entirely absent with dominant local synchronization between a few areas (e.g. Fig. 1B bottom left, blue rectangle) in both human and monkey datasets.

To quantify these varying patterns of global and local synchronization, we calculated the Fano factor (FF), a measure widely applied in the analysis of population spiking activity (Brunel and Hakim, 1999; Hahn et al., 2017; Kumar et al., 2008), which captures higher order correlation and global synchrony between discrete events (Fig. 1C, see Methods for a detailed definition). The FF equals ~1, when there is no correlation across brain areas, resembling a random Poisson process (Fig. 1C, right). Synchronization across brain regions increases the FF to values >1, with brain wide synchronization events being associated with FF >=1 (Fig. 1C, left).

We subsequently computed FFs for sliding windows of different sizes and their distribution for each vigilance state. The distributions decayed exponentially for small window sizes (Fig. 1D), while they generally followed a gamma distribution for larger window sizes in human and monkey datasets (Fig. 1E, K5-test; human: window size = 5TR, D = 0.02±0.006, p < 0.001; window size = 25TR, D = 0.03±0.01, p < 0.001; monkey: window size = 5TR, D = 0.06±0.006, p < 0.001; window size = 25TR, D = 0.23±0.07, p < 0.001). The shape parameter (beta) of these gamma distributions, increasing for higher right skewness, was subsequently used to quantify the level of synchronization.

Notably, the distributions clearly differed between levels of consciousness in the human and monkey data with the exception of awake and N1 states in humans, which showed a similar synchronization profile (window size = 5TR, one-way rm-ANOVA: F3, 51 = 1.9 × 10^5, p < 0.001, ϵ = 1 in the human and F2,28 = 1.48 × 10^5, p < 0.0001, ϵ = 0.6 in the monkey; posthoc test: p < 0.001 for all pairs except for awake vs. N1 with p = 0.99). In the awake state, the distributions were more shallow and right skewed (Fig. 1D and E) with higher beta values due to a higher prevalence of large-scale synchronization between brain areas. Unconscious states were associated with steeper and more narrow distributions due to missing large-scale synchronization (Fig. 1D and E) and reduced beta values. Importantly, the beta parameter also distinguished between different levels of unconsciousness as it progressively decreased when subjects drifted into deeper levels of sleep or anesthesia. The same results held, when negative fMRI peaks were used for the calculation of FF distributions (Supplementary Fig. 2). To test the robustness of this novel marker, we calculated beta for different window sizes and indeed found that the separation of consciousness states remained unchanged up to a window size = 50 TR (Fig. 1F). Additionally, these findings suggest that the varying levels of large-scale synchronization, as indexed by the FF distribution, could serve as a robust marker for different vigilance states and the depth of unconsciousness.

2.1. Computational model

What might be the origin of these declining levels of large-scale synchronization as brain dynamics traverses from awake to gradually deeper levels of unconsciousness? To tackle this question, we devised a whole-brain computational model that aimed at reproducing these changes in global dynamics. This model simulated local fMRI signals for each brain area of two different parcellations (human: N = 90 areas (Tzourio-Mazoyer et al., 2002), monkey (Paxinos et al., 2009): N = 432 areas), using the normalized Hopf bifurcation model (see Methods). The brain areas were connected on the basis of diffusion-tensor imaging data from humans and monkeys (Adlrur et al., 2012; Zakszewski et al., 2014). The core parameter of the model is the bifurcation parameter (alpha), which captures the excitability of each node, i.e. its responsiveness to incoming input from other brain areas (Supplementary Fig. 3A). We uniformly varied alpha across all brain areas mimicking changes in global excitability mediated by diffusely projecting neuromodulator systems (e.g. acetylcholine, noradrenaline), whose activity is known to change across the sleep-wake cycle (Scammell et al., 2017). Depending on the values of alpha, the model displayed three qualitatively different
activity states. When alpha was kept small, the dynamics of the model was governed by synchronization between a few brain areas, but remained free of large-scale synchronization (Fig. 2A, left). This dynamical regime was strikingly similar to the empirical pattern of unconscious states in the monkey and human data (Fig. 1A, bottom). At intermediate values, the dynamics was characterized by periods of asynchronous activity, as seen with smaller alpha, but sporadically interrupted by events of global synchrony (Fig. 2A, center). This activity mode was reminiscent of the dynamics seen in the awake humans and monkeys (Fig. 1B, top). Further increase of alpha was accompanied by a dynamical state with highly synchronized and oscillatory activity across the entire model brain (Fig. 2A, right). More generally, the behavior of the model characterizes a bifurcation from exclusively local synchrony (alpha < bifurcation) to global synchronization (alpha > bifurcation) with elements of both types of synchronization being present close to the bifurcation point (alpha ~ bifurcation).

Next, we computed Fano factor distributions for the simulated fMRI signals. These distributions were also governed by gamma distributions, which approached an exponential function with small time windows, similar to the empirical counterpart, for alpha < or ~ bifurcation (Fig. 2B). The highly synchronized regime, however, deviated from this behavior and showed a clear peak in the FF-distribution caused by
the global oscillation. After normalization, such that the bifurcation appeared at \( \alpha = 0 \) (see Methods, Supplementary Fig. 3B), we fitted gamma distributions for alpha values < 0 and showed that the obtained beta values increased monotonically as alpha approached the bifurcation point (Supplementary Fig. 3C). Thus, different levels of synchronization in the model mapped directly onto the FF distribution. This allowed us to find the alpha values of the model, where the beta values of empirical and simulated FF distribution were identical, i.e. their difference was close to zero. We found that the awake state was close to the bifurcation point with both locally and globally dominated synchronization in both human and monkey data. As subjects lost consciousness, the alpha values dropped progressively with deeper levels of sleep or anesthesia, indicating a gradual shift in the dynamics towards smaller synchronization events (Fig. 2C-F). These results suggest that changes in global excitability are the main determinant of different levels of large-scale synchronization in the simulated and empirical data.

### 2.2. Model predictions

The present findings demonstrate that the model can mirror the time-varying properties of global versus local synchronization across different levels of consciousness. We therefore tested whether the simulations can also correctly predict the change of other metrics across vigilance states. First, we calculated the grand-average functional connectivity (FC) of fMRI signals for each state of the empirical and model data. In accordance with previous findings (Barttfeld et al., 2015), average FC between brain areas was highest in the awake state and significantly decreased with onset and progression into deeper levels of unconsciousness in both human and monkey data (Fig. 3A, left, one-way rm-ANOVA: \( \text{F3,51} = 0.99 \times 10^3, p < 0.0001, \epsilon = 0.42 \) in the human and \( \text{F2,28} = 1.81 \times 10^3, p < 0.0001, \epsilon = 0.78 \) in the monkey; posthoc test: \( p < 0.001 \) for all pairs). Importantly, the model also showed higher average FC during the awake state and lowest FC during the deepest levels.
of unconsciousness (Fig. 3A, right). Next, we examined the relationship between structural and functional connectivity, which has been shown to undergo changes across consciousness states (Barttfeld et al., 2015; Tagliazucchi et al., 2016b). Similarly, we detected an increase of the SC-FC relation with loss of consciousness as compared to the awake state. This indicates stronger expression of direct anatomical connections in the FC during unconscious states, whereas in the awake state further network effects emerge beyond pair-wise interactions (Fig. 3B, left, one-way rm-ANOVA: F3,51 = 1.13 * 10^3, p < 0.0001, ε = 0.45 in the human and F2,18 = 2.56 * 10^3, p < 0.0001, ε = 0.92 in the monkey; posthoc test: p < 0.001 for all pairs). Again, the model correctly reproduced the behavior of the SC-FC relationship for different levels of consciousness (Fig. 3B, right). Other descriptors that are used to characterize consciousness include the level of integration and segregation, which are in general inversely related (Deco et al., 2015). Integrated activity is characterized by synchronization across multiple brain areas, while modularity refers to the prevalence of locally confined processing. In the human and monkey data, integration significantly increased,
as brain activity approached wakefulness (Fig. 3C, left, one-way rm-ANOVA: F3,51 = 0.911 10^{-3}, p < 0.0001, \epsilon = 0.43 in the human and F2,28 = 4.36 10^{-3}, p < 0.0001, \epsilon = 0.84 in the monkey; post hoc test: p < 0.001 for all pairs) due to rising levels of brain-wide synchronization. In contrast, local activation dominated during unconscious states with more segregation, quantified by the modularity value that we obtained for the optimal partition when breaking the FC matrix into clusters (Fig. 3D, left, one-way rm-ANOVA: F3,51 = 1.04 10^{3}, p < 0.0001, \epsilon = 0.65 in the human and F2,28 = 4.92 10^{-3}, p < 0.0001, \epsilon = 0.59 in the monkey; post hoc test: p < 0.001 for all pairs). The model correctly predicted the change of integration and modularity across vigilance states (Fig. 3C and D, right).

2.3. Analysis of different synchronization states

Previous studies have highlighted that different consciousness levels can be characterized by the relative prevalence of different functional connectivity states (Barttfeld et al., 2015; Demertz et al., 2019) with varying levels of fMRI signal correlations and SC-FC relationship. Here, we hypothesize that these states reflect different levels of large-scale synchronization with a frequency of occurrence that is determined by the level of consciousness. To test this hypothesis, we split the data into short bins (size = 5 TR) and concatenated bins with similar Fano factor values, representing similar global synchronization levels, into a new dataset. The FF range was chosen to be 1 + n < FF < 5 + n, with n being incremented by one for each new dataset. Thus, we obtained datasets with increasing large-scale synchronization, each of which was considered a different state. This procedure was repeated for both empirical and model data, and for each vigilance state.

We first examined how the average amplitude of the fMRI signal, measured as the mean z-score of the positive peak amplitude, changed across different synchronization states for each vigilance state. The analysis of the human data showed that the BOLD amplitude steadily rises with increase in global synchronization, indicating that large-scale synchronization is associated with stronger local signals (Fig. 4A, left). Importantly, these results were accurately predicted by the model. A similar behavior was found in the empirical and modeled monkey data for the awake state, while the fMRI signal amplitude slightly dropped with stronger synchronization in the two anesthetized states of the empirical data (Fig. 4A, right).

Next, we studied how the mean FC evolves across the different synchronization states. As the global synchronization increased in the empirical human data, the mean FC also grew stronger for all vigilance conditions (Fig. 4B, left). However, the mean FC for most large-scale synchronization states was higher in the awake state as compared to the unconscious conditions. Likewise, highly synchronized states also showed the highest FC and were only found in the awake condition. The model was able to replicate the empirical findings. In the empirical and modeled monkey data, similar results were obtained with more pronounced FC differences between awake and unconscious conditions (Fig. 4B, right).

We then studied to what extent the SC is expressed in the FC as a function of different global synchronization states. We found that with increasing levels of large-scale synchrony, the similarity between empirical FC and SC in humans decreases and reaches minimum values during highly synchronized states prevalent only during the awake condition (Fig. 4C, left). A similar decrease was seen in the model, even though very small SC-FC correlation values were also found in unconscious states. The same results held in the monkey data, with the exception that high SC-FC similarity was found predominately in weakly synchronized states during anesthesia (Fig. 4C, right).

The integration measure also increased with rising levels of global synchronization across all consciousness conditions in humans and monkeys, a result that was accurately reproduced by the model (Fig. 4D). In contrast, modularity decreased with large-scale synchrony in human empirical and simulated data (Fig. 4E, left). Interestingly, modularity in the monkey initially decreased with larger FF values and subsequently increased again in the anesthetized states, showing a U-shaped behavior in both empirical and modeled data. In contrast, the awake state was characterized by a monotonic decrease of modularity with higher FF values as seen in humans.

Overall, this analysis suggests that different global synchronization states with varying levels of FC, expression of SC, integration and modularity exist across different vigilance conditions. The level of consciousness can then be distinguished based on the composition of synchronization states, which, as the model shows, is a function of the global excitability level across the entire brain.

2.4. Single trial analysis

The previous results were obtained by computing synchronization metrics from data merged across all trials and subjects of a consciousness state. To test whether the composition of synchronization states can also delineate consciousness states at the single-trial level, we computed FF distributions and the other FC-based metrics for each trial separately. In humans, we recorded one trial per consciousness state with a duration that ranged between ~90 and 1000 TR (see Supplementary Fig. 4 for details). We found significant beta value differences between the awake state and unconscious states (N2 and/or N3) (one-way rm-ANOVA: F3,51 = 5.7, p < 0.002, \epsilon = 0.93), which was repeated for the other measures except modularity, while the separation between unconscion states was not consistent across sleep stages (Fig 5A, Supplementary Fig. 5). Moreover, when comparing consciousness states within single subjects, there was a monotonic decrease of beta values from the awake state to N3 sleep only in a few cases, while most subjects did not show a consistent change of beta across consciousness states (Supplementary Fig. 6). Notably, the beta values and other measures were highly overlapping across states, prohibiting an unambiguous identification of consciousness states. As we have shown above, each consciousness state can be defined by a distribution of synchronization events, in which reduced global synchrony occurs across all states, while larger events are mainly prevalent during the awake stage and N1 sleep. Thus, disambiguation of states requires sufficient sampling to provide accurate statistics, which may not be the case in our single trial analysis due to short trial duration.

To test whether these findings can be replicated by the human model, we simulated several short trials (250 TR) and averaged across trials (Fig. 5B). Similar to the empirical counterpart, the beta values monotonically decreased with the depth of sleep and with statistical differences across states as seen in the empirical data, while they remained highly overlapping. In contrast, when we performed simulations with long trials (10,000 TR), the overlap disappeared and each trial could be unequivocally assigned to a particular consciousness state (Fig. 5C).

In monkeys, where the trial duration was fixed at 500 TR, the awake state was significantly different from the two unconscious states for all measures (Fig. 5D, Supplementary Fig. 5, one-way rm-ANOVA: F2,60 = 8.6, p < 0.0001, \epsilon = 0.45 in the human and F2,18 = 2.56 10^{-3}, p < 0.0001, \epsilon = 0.6 in the monkey). Yet, the values were overlapping between the states, which we reproduced in simulations with short trials (500 TR, Fig. 5E). Similar to humans, the overlap vanished with longer trials (Fig. 5F).

Overall, these results suggest that short trials are less suited for an unambiguous separation of different consciousness states due to insufficient sampling of large-scale synchronization events across time.

2.5. Model lesions and global synchronization

The correspondence between measures of the empirical and model data suggests that the model may be a viable tool to investigate the impact of specific brain areas on consciousness. We thus employed a lesion approach in which the excitability (i.e. the bifurcation parameter
alpha) of individual nodes in the model was strongly reduced, mimicking damage after stroke or mechanical trauma (Adhikari et al., 2017). To quantify the consequences of node damage on model dynamics and its associated level of consciousness in the empirical data, we introduced a lesion index ($\lambda$), which indicates the loss of global synchronization in the simulations with respect to the synchronization level seen in the awake state of the empirical recording (see Methods for details). A value of $\lambda = 1$ reflects model synchronization equivalent to the awake data, while we consider $\lambda < 0.85$ as equal to loss of consciousness.

Fig. 4. Various metrics change with the level of global synchronization (FF states) across different consciousness conditions. (A) Average peak amplitude (z-score) within a time window of 5 TR as a function of FF state in empirical and modeled human (left) and monkey data (right). (B) Mean direct FC as a function of global synchronization states for human and monkey data. (C) SC-FC similarity for different global synchronization states in humans and monkeys. (D-E) Same as in (C) for integration and modularity, respectively.
When each area was lesioned separately, the impact on large-scale synchronization was modest for most regions and yielded a lesion index that remained above the threshold for dynamics associated with consciousness in both human and monkey data (Fig. 6A). However, the lesion of a few areas strongly reduced brain-wide synchronization with $\lambda$ descending to values $\approx 0.4$ in some cases. In humans, these areas were all found in the posterior brain, with the precuneus and posterior cingulate cortex having the most pronounced effect (Fig. 6B, left panel). This is in striking congruence with previous experimental findings pointing at a posterior hotspot for consciousness (Koch et al., 2016). Notably, in simulations of monkey data the strongest influence on global synchronization was also found in the posterior cingulate cortex (area 23) and the precuneus (area PGM) together with subcortical nuclei, in particular higher order thalamic nuclei (Fig. 6B, right panel), which according to empirical studies play an important role in modulating consciousness (Schiff, 2008).

To gain a better understanding of how these nodes contribute to global dynamics and putatively to the maintenance of consciousness, we conducted a graph theoretical analysis of the human and monkey structural connectome. We first tested the structural matrices for the presence of a rich club, i.e. network hubs that are highly interconnected with other brain areas and also strongly linked between themselves. Based on the k-density metric (see Methods, Supplementary Fig. 7), we found several rich club nodes in the human, including the precuneus and posterior cingulate cortex, and monkey connectome, which were mainly located in subcortical nuclei and area PMG (precuneus) (Fig. 7A and B, asterisks). Even though a majority of these areas had a strong impact on the level of large-scale synchronization, a large number of brain regions with a low lesion index did not belong to the rich club network (Figs. 6B and 7A,B). We thus searched for a metric that would better reflect the role of each node in maintaining brain-wide synchronization and the awake state. The node communicability, a measure that captures the propagation of activity across the network after the stimulation of a given node, proved to be a good predictor of synchronization levels (Fig. 7A and B). In general, we found an inverse relationship such that areas with a lower lesion index typically had higher communicability values in human and monkey datasets. Surprisingly, however, in humans we found two separate linear relationships between communicability and lesion index, one with a steep slope where the impact of a lesion was high and another with a shallower slope. We hypothesized that this finding is rooted in the difference of node communicability between different hemispheres, since taking out a node in one hemisphere influences large-scale synchronization both ipsilaterally and contralaterally only if strong interhemispheric connections exist. Thus, a node may have strong communicability in the presence of only ipsilateral connections, but a lower impact on large-scale synchronization, as its activity does not spread to the other hemisphere. We thus calculated communicability separately for each hemisphere and quantified the asymmetry between both communicability values. A value of $\sim 1$ indicates high asymmetry with a node exclusively impacting activity in one hemisphere, while asymmetry $\sim 0$ indicates a strong propagation within both hemispheres. We indeed found that high communicability
areas can show different degrees of asymmetry across the two hemispheres (Supplementary Fig. 8). Despite their similar communicability, areas with reduced asymmetry such as the precuneus and posterior cingulate cortex influence brain-wide synchronization strongly, while other nodes with very high asymmetry such as the putamen influence synchronization primarily in one hemisphere with an attenuated impact on brain-wide synchrony (Fig. 7C).

3. Discussion

Here, we have shown that the pattern of large-scale synchronization across the brain is a new signature of consciousness and can readily distinguish different consciousness conditions in human and monkey fMRI data. It separates awake from unconscious conditions and also delineates various degrees of non-REM sleep and propofol anesthesia. Changes in brain-wide synchrony are accompanied by alterations of other measures of consciousness such as FC, the SC-FC relationship, integration and modularity. Strikingly, the statistics of these metrics were accurately reproduced by a whole-brain model and explained by different global excitability levels that naturally generate different configurations of large-scale synchrony. Additionally, graph theoretical analyses together with model lesion simulations show that the anatomical properties of specific brain areas in the posterior brain play a crucial role in maintaining the level of large-scale synchronization typical of the awake state.

The existence of brain-wide synchronization in fMRI data naturally raises the question whether it reflects a physiological signal (Scholvinck et al., 2010) or should be considered as an artifact caused by e.g. head movements that are usually regressed out (Liu et al., 2017), as was also done here. However, recent studies have demonstrated that large-scale synchrony of fMRI signals not only respect precisely orchestrated electrophysiological changes (Liu et al., 2018a, 2015), but is also tightly linked to acetylcholine release of the basal forebrain, since its lesion abolishes brain-wide synchronization (Turchi et al., 2018). Changes in global synchrony were also found in schizophrenia (Hahamy et al., 2014; Yang et al., 2014). Moreover, the fMRI signal shows wave-like propagation (Majeed et al., 2011; Mitra et al., 2014, 2015a) across the entire brain during the awake state with large-scale synchronization (Mitra et al., 2015b), while waves remain confined to specific resting state networks during slow-wave sleep (Mitra et al., 2015b). Lastly, both empirical and model distributions decay exponentially and lack a peak at their tail, where highly widespread synchrony due to artifacts would be expected.

Previous studies have highlighted that fMRI signal synchronization between brain areas fluctuates over time, creating different functional connectivity states (Barttfeld et al., 2015; Keilholz et al., 2017). These FC states are tied to different levels of modularity (Betzel et al., 2016; Fukushima et al., 2018a), integration (Shine et al., 2016), varying SC-FC relationship (Barttfeld et al., 2015; Fukushima et al., 2018b), behavior (Shine et al., 2016) and the level of consciousness (Barttfeld et al., 2015; Demertzi et al., 2019). Here, we show that large-scale synchronization evolves at a fine-grained temporal scale and changes strength across a wide range in the awake state, but a largely reduced range during unconscious conditions in both empirical and modeled human and monkey data. We show that global synchrony is related to other measures such as mean FC strength, SC-FC relationship, modularity and integration. These metrics move along a continuum with a range that changes with the consciousness level, rather than reflecting a few discrete states that are correlated with behavior (Shine et al., 2016; Wang et al., 2016). During the awake state, consciousness is maintained over a large range of synchronization states including those that are primarily seen during loss of consciousness with a lack of global synchrony. Such absence of global synchrony, which we found to dominate during the N3 state, is also consistent with reports that during sleep both the BOLD signal and slow waves/spindles in electrophysiological recordings propagate only locally (Mitra et al., 2015b; Nir et al., 2011; Vyzazovskiy et al., 2011). The temporary absence of large-scale synchrony and integration may indicate a low arousal state with drowsiness, which is not yet low enough to suppress consciousness, even though a clear correlation with microsleep was not found (Demertzi et al., 2019). On the other hand, highly integrated states have been associated with accurate task performance (Shine et al., 2016) and are predicted by both the integrated information and global neuronal workspace theories of consciousness (Dehaene and Changeux, 2011; Tononi et al., 2016), as conscious processing would require large-scale integration and spread of neuronal activity (‘ignition’) (Deco et al., 2017b; van Vugt et al., 2018). In contrast, other studies suggest that such strong global synchronization is accompanied by a drop of arousal as defined by a reduction of alpha activity and enhanced delta/theta oscillations in simultaneously recorded electrophysiological signals (Chang et al., 2016; Wong et al., 2013), which is reversed by alertness-promoting caffeine (Wong et al., 2013), and putatively caused by reduced ACH concentrations after a large-scale synchronization event (Liu et al., 2018b). It is possible that such brain-wide synchronization with high Fano factors reduces functional differentiation of neuronal activity, another hypothesized prerequisite for consciousness (Nacci et al., 2018; Tononi et al., 2016). Even though systems close to a bifurcation, as seen in our study for the awake state, theoretically maximize their number of possible states (Shew et al., 2011; Shew and Plenz, 2013), the occasional undifferentiated and more homogeneous activation of large parts of the brain would thus diminish alertness. A key component of enhanced differentiation during the awake state might be enhanced inhibition (Rudolph et al., 2007), which allows competition between different neuronal representations and fine-grained activity patterns. Indeed, active suppression of activity has only been found during the awake state, as evidenced by a specific FC configuration, containing strong correlation and anticorrelations between brain areas (Barttfeld et al., 2015; Demertzi et al., 2019; Fox et al., 2005; Wong et al., 2013). A possible scenario is that such inhibition is reduced during strong global synchrony, which triggers a large-scale propagation of excitation with lower levels of differentiation. An important limitation of our marker is...
that the global synchronization index is not sensitive to anticorrelations. Likewise, the presented model does not incorporate mutual inhibition between brain areas and can thus not reproduce the anticorrelated FC states seen during the awake state, a problem which will be addressed in future studies.

Another limitation of our marker of consciousness is that it is based on a distribution, which requires sufficient sampling across time to give a robust separation of consciousness states. If sampling is too short the estimation of distribution parameters is inaccurate and consciousness states can have similar marker values, as shown by our single-trial analysis. Yet, our model results suggest that with sufficient recording time states can be distinguished with high fidelity. In addition, the existence of distributions of synchronization strength, which differ in their tail of large-scale synchronization across consciousness states, but otherwise overlap in terms of small-scale synchronization, imply that in both conscious and unconscious states there are epochs of neuronal activity in parcellated fMRI data, which are statistically not distinguishable. Thus, more research is needed to overcome this limitation of distribution-based markers and correctly detect consciousness at any given moment in time.

What is the origin of the fluctuations in large-scale synchrony, integration and separation seen in empirical data (Deco et al., 2015; Fukushima and Sporns, 2018)? There are two scenarios: First, the cortex inherits excitability fluctuations originating in neuromodulatory nuclei or second, fluctuations are created intrinsically in the cortex.

Our whole-brain model creates fluctuations intrinsically and can faithfully reproduce the statistics of global synchronization and connectivity measures at a fine temporal scale, given a fixed level of global excitability, and across a variety of consciousness conditions. Such fluctuations within a continuum of largely synchronized and desynchronized states is an expected property of a dynamical system close to a Hopf bifurcation point (Jobst et al., 2017), which resembles the awake state in humans and monkeys. This idea is similar to the concept of criticality and neuronal avalanches (Beggs and Plenz, 2003; Hahn et al., 2017; Hahn et al., 2010; Liang et al., 2020; Shew and Plenz, 2013), where a system poised to the critical point generates fluctuations at various spatial scales with occasional avalanches that spread across the entire system, akin to the fMRI waves seen in awake humans (Mitra et al., 2015a). Moving away from the bifurcation or to the subcritical regime by reducing global excitability levels reduces the size of fluctuations and avalanches are restricted to local propagation within strongly connected resting state networks (Tagliazucchi et al., 2016a). Our model indeed suggests that the transition from awake to deeper stages of sleep or anaesthesia is linked to a gradual decrease in global excitability and large-scale functional connectivity.

However, there is also experimental evidence that the variability of functional connectivity is at least partly driven by changes in neuromodulation. One study showed that integrated FC states appear during cognitive tasks and are accompanied by a dilation of pupil diameter indicative of enhanced activity of subcortical neuromodulator nuclei, in
particular the noradrenergic locus coeruleus (Shine et al., 2016). Another study linked the occurrence of global synchronization during resting state to a decrease in cholinergic basal forebrain activity suggesting that a decreased ACH tone is causally related to widespread propagation of neuronal activity (Liu et al., 2018a). Furthermore, inactivation of the nucleus basalis disrupts the presence of large-scale synchronization, while keeping the structure of local resting state network synchronization intact (Turci et al., 2018). This finding is reminiscent of our model result where reduced global excitability diminished global synchrony. Thus, it remains to be studied how fluctuations in neuromodulation are causally related to individual connectivity and synchronization states. It is unclear to what extent neuromodulatory changes are stochastic and independent of cortical activity or are to some extent under the control of cortical dynamics.

Our whole-brain modeling also allowed us to mimic lesions of individual areas due to e.g. stroke, and study their effect on global synchrony as seen during the awake state. We found a few key areas that are crucial for maintaining the model in a condition akin to the awake state and their damage resulted in a transition into model dynamics in which brain-wide synchronization events were absent similar to unconscious states. Surprisingly, these areas were located in the back of the brain, centered around the ‘posterior hotspot’ of consciousness described in the literature (Boly et al., 2017; Koch et al., 2016; Tagliazucchi, 2017; Vogt and Laureys, 2005). The most influential areas in the model were the precuneus and posterior cingulate cortex in both humans and monkeys, which have been previously found to be key areas in maintaining consciousness (Alkire et al., 2008; Cavanna, 2007). In the monkey, thalamic nuclei had a large impact, as described previously in empirical studies (Schiff, 2008). In the human parcellation the thalamus was only represented by one node per hemisphere with less impact on consciousness, explained by its weaker node communicability of the anatomical connections as compared to highly influential areas (Fig. 7A). Using graph theoretical analysis, we identified a few areas with large impact on large-scale synchrony that are part of the structural rich-club, a highly interconnected network of hubs that define the default node network in the posterior brain (van den Heuvel and Sporns, 2011; Zamorano-López et al., 2011; Zamorano-López et al., 2010). More generally, the effect of individual brain areas on global synchronization was attributed to the degree of node communicability, i.e. how strongly activity initiated in a given area propagates across the brain. We also found that strong interhemispheric connections play a crucial role in mediating synchrony across the entire brain. This effect was most pronounced in humans and much less in monkeys. We can only speculate about the causes for this difference between species. Aside from existing anatomical differences, it is also conceivable that the difference in parcellation or tractography, which is known to confound interhemispheric connections specifically, may play a role. Overall, these results demonstrate that excitability and information transfer through single nodes is not only determined by neuromodulation, but also by the activity flowing in and out of crucial nodes in the network. These nodes then play a paramount role in keeping overall global excitability at a level necessary for conscious processing.

In conclusion, we described a novel marker for different consciousness states that is based on large-scale synchronization of the fMRI signal across the brain and can be solidly explained by changes of global excitability in a whole brain model. This model also suggests that the maintenance of consciousness is dependent on graph theoretical properties of individual brain areas, making conscious experience vulnerable to lesion of specific nodes.

4. Methods

4.1. Human study

4.1.1. Participants

For this study, approved by the local ethics committee, we collected data from 63 young healthy subjects, who provided written informed consent and were reimbursed for their participation. Eight subjects remained awake during scanning and were not included in the study, leaving five subjects who reached at least N1 sleep (36 females, mean ± SD age of 23.4 ± 3.3 years). Simultaneous EEG-fMRI was recorded in the evening after a regular sleeping schedule. For this study, we only included 18 participants, who went through all three sleep stages (N1, N2 and N3).

4.1.2. Acquisition of fMRI and EEG and artifact processing

We simultaneously recorded EEG via a cap (modified BrainCapMR, Easycap) and fMRI (1505 vol of T2*-weighted echo planar images, TR/TE = 2080 ms/30 ms, matrix 64 × 64, voxel size 3 × 3 × 2 mm³, distance factor 50%; FOV 192 mm²) with a 3T Siemens Trio scanner. Polysomnography was performed, recording chin and tibial EMG, EOG, EEG bi-polarly (sampling rate 5 kHz, low pass filter 1 kHz) and EEG with 30 channels using FCz as the reference (sampling rate 5 kHz, low pass filter 250 Hz); We recorded pulse oxymetry and respiration with sensors from the Trio (sampling rate 50 Hz) and MR scanner compatible devices (BrainAmpMR+, BrainAmpExG; Brain Products), which aided sleep scoring during fMRI recordings (Jahneke et al., 2012). MRI and pulse artifacts were removed with the average artifact subtraction (AAS) method (Allen et al., 1998) as implemented in Vision Analyzer2 (Brain Products) and thereafter by objective (CBC parameters, Vision Analyzer) ICA-based rejection of residual artifact components, after which the EEG had a sampling rate of 250 Hz (Jahneke et al., 2012). Likewise, motion artifacts in the EEG were removed based on ICA using the Vision An- alyzer2 software. An expert scored sleep stages manually according to the AASM criteria (Iber et al., 2007).

4.1.3. fMRI preprocessing

Using Echo planar imaging (EPI) data were realigned, normalized (MNINet space), spatially smoothed (Gaussian kernel, 6-mm* full width at half maximum) with the Statistical Parametric Mapping software (SPM8, www.fil.ion.ucl.ac.uk/spm) and resampled to a 4 × 4 × 4 mm³ resolution for artifact removal. Then we regressed out cardiac, respira- tory (both estimated using the RETROICOR method (Glover et al., 2000)) and motion-induced noise. For data analysis, the data were first bandpass filtered in the range of the 0.01–0.1 Hz (Cordes et al., 2001) us- ing a second order Butterworth filter or between 0.07 and 0.09 Hz to compare empirical with model data (see below). All voxels were then projected onto the automated anatomic labeling (AAL) atlas (Tzourio- Mazoyer et al., 2002) (90 areas, 76 cortical regions, 14 subcortical re- gions, excluding cerebellar regions, Supplementary Table 1) by averag- ing their time courses within each AAL cluster. Alternatively, we also computed the first principal component of the voxel time-series and compared it with the averaged time course. The results show that the two time series are highly similar (Supplementary Fig. 1A) with a correlation coefficient of 0.96 ± 0.04 (Supplementary Fig. 1B).

4.1.4. Anatomical dataset acquisition in humans with DTI

The human SC-matrix was obtained at Aarhus University, Denmark using DTI (3T Siemens Skyra scanner) in 16 healthy right-handed par- ticipants (11 men and five women, mean age: 24.75 ± 2.54). Subjects with a history of psychiatric or neurologic disorders were not included in this study. The T1 acquisition of the structural MRI was performed with a voxel size of 1 mm³, a reconstructed matrix size of 256 × 256, echo time (TE) of 3.8 ms and repetition time (TR) of 2300 ms. The DTI data were acquired using TR = 9000 ms, TE = 84 ms, flip angle = 90°, reconstructed matrix size of 106 × 106, voxel size of 1.98 × 1.98 mm with slice thickness of 2 mm and a bandwidth of 1745 Hz/Px. Moreover, the data collection was carried out with 62 optimal nonlinear diffusion gradient directions at b = 1500 s/mm² and roughly one nondiffusion weighted image (DWI; b = 0) per 10 diffusion-weighted images was ob- tained. In addition, DTI images were taken with opposing phase encoding directions (anterior to posterior or posterior to anterior). Similar to the fMRI data, we parcellated the structural scans into 90 areas using the
AAL template. We used the linear registration tool from the FSL toolbox (www.fmrib.ox.ac.uk/fsl, FMRIB (Jenkinson et al., 2002)) to coregister the EPI image to the T1-weighted structural image. The T1-weighted image was coregistered to the T1 template of ICBM152 in MNI space (Collins et al., 1994). The resulting transformations were concatenated and inverted and further applied to warp the AAL template (Tzourio-Mazoyer et al., 2002) from MNI space to the EPI native space, where interpolation using nearest-neighbor method ensured that the discrete labeling values were preserved. SC maps for each participant using the DTI data were acquired and processed based on the two datasets with different phase encoding. To construct the SC matrix, we estimated connections between nodes defined by the AAL atlas using probabilistic tractography and averaged across subjects.

4.2. Monkey study

4.2.1. Animals

Three rhesus macaques (Macaca mulatta, 1 male, 2 females, weight 5–8 kg, age 6–12 years) were tested for each arousal state (awake, moderate propofol sedation, deep propofol anesthesia). All procedures were conducted in accordance with the European convention for animal care (86–406) and the National Institutes of Health’s Guide for the Care and Use of Laboratory Animals. Macaque studies were approved by the institutional Ethical Committee (protocols #10-003).

4.2.2. Propofol anesthesia protocol

Detailed methodology was described in our previous studies (Barttfeld et al., 2015; Uhrig et al., 2018). The propofol anesthesia levels (moderate and deep anesthesia) were defined using a clinical score, the monkey sedation scale (spontaneous movements, response to juice presentation, shaking/prodding, toe pinch and corneal reflex), determined at the beginning and the end of each scanning session and continuous electroencephalography (EEG) monitoring (Barttfeld et al., 2015; Uhrig et al., 2018). Under deep propofol anesthesia (general anesthesia), monkeys did not respond to any stimuli. EEG Propofol anesthesia levels were defined as follows (Barttfeld et al., 2015; Uhrig et al., 2018): level 1, awake state, posterior alpha waves and anterior beta waves; level 2, light propofol sedation, increasing of the amplitude of alpha waves and anterior diffusion of alpha waves; level 3, moderate propofol sedation, diffuse and wide alpha waves, and anterior theta waves; level 4, deep propofol anesthesia (general anesthesia), diffuse delta waves, waves of low amplitude and anterior alpha waves (10 Hz); level 5, very deep sedation (deeper then level of general anesthesia), burst suppression.

The awake monkeys were trained to be injected with an intravenous propofol bolus (5–7.5 mg/kg i.v.; Fresenius Kabi, France) to induce anesthesia, followed by target-controlled infusion (Alaris PK Syringe pump, Carefusion, CA, USA) of propofol (moderate propofol sedation: 3.7–4.0 microg/ml; deep propofol anesthesia 5.6–7.2 microg/ml) based on the ‘Paedusor’ pharmacokinetic model (Absalom and Kenny, 2005). During moderate propofol sedation sessions, a muscle-blocking agent was co-administered (cisatracurium, 0.15 mg/kg bolus i.v. followed by continuous i.v. infusion at a rate of 0.18 mg/kg/h, GlaxoSmithKline, France) to prevent artifacts related to potential movements during the fMRI acquisition. Monkeys were intubated and ventilated. Heart rate, non-invasive blood pressure, oxygen saturation, respiratory rate, end-tidal CO2, cutaneous temperature was monitored (Maglile, Schiller, France) and recorded online (Schiller, France) (Barttfeld et al., 2015; Uhrig et al., 2018). Temperature, hemodynamic and ventilation parameters were kept constant during each experiment (Barttfeld et al., 2015; Uhrig et al., 2018).

4.2.3. Awake state protocol for the monkeys

For the awake state acquisitions, monkeys were implanted with an MR-compatible headpost and trained to sit in the spinhix position in a primate chair (Uhrig et al., 2014; Wang et al., 2015). Monkeys sat in the dark inside the MRI without any task (Barttfeld et al., 2015; Uhrig et al., 2018).

4.2.4. Monkey fMRI acquisition

Monkeys were scanned with a customized single transmit-receiver surface coil on a 3-T horizontal scanner (Siemens Tim Trio, Erlangen, Germany). Before each scanning session, monocrystalline iron oxide nanoparticles (MIOM, Feraheme; AMAG Pharmaceuticals; 10 mg/kg, i.v.) were injected into the monkey’s saphenous vein (Barttfeld et al., 2015; Uhrig et al., 2018). Each functional scan contained 500 gradient-echoplanar whole-brain images (TR = 2400 ms; TE = 20 ms; and 1.5-mm3 voxel size). 27 awake runs, 25 moderate propofol sedation runs and 31 deep propofol anesthesia runs were acquired.

4.2.5. Monkey fMRI preprocessing

Monkey fMRI images were reoriented, realigned and rigidly coregistered to the anatomical template of the monkey Montreal Neurological Institute space (MNI) (Frey et al., 2011; Uhrig et al., 2014). We also regressed out movement parameters resulting from rigid body correction for head motion. We removed the global signal from the images to minimize the potential respiratory and cardiac effects associated with the propofol administration. Voxel time series were parcelled according to the Paxinos atlas (Paxinos et al., 2009) (see Supplementary Table 2), encompassing 432 cortical and subcortical areas. The signals were then band-pass filtered between 0.01 and 0.1 Hz or in the range of 0.07–0.09 Hz to compare empirical with model data (see below).

4.2.6. Anatomical dataset acquisition in monkeys with diffusion tensor imaging (DTI)

DTI data collected in 271 young rhesus macaques in the age-range of 0.7570 to 4.2027 years (Adluru et al., 2012) were used for the creation of a macaque SC matrix. Data collection in macaques was performed using protocols approved by the University of Wisconsin Institutional Animal Care and Use Committee (IACUC). Before undergoing MRI acquisition, the monkeys were anesthetized using ketamine (15 mg/kg). MRI scanning was performed using a GE SIGNA 3T scanner. MR scanning was performed using a two-dimensional, echo-planar, diffusion-weighted, spin-echo sequence with the following parameters: repetition time = 10 s; echo time = 77.2 ms; field of view=14 cm; matrix=128 × 128; 2.5 mm thick contiguous slices; echo-planar echo spacing=800 µs. Diffusion imaging (b = 1000 s/mm2) was performed in 12 non-collinear directions with one non-diffusion weighted image and the acquisition was repeated six times and averaged. The volumes were eddy-current corrected using FSL and then brain extraction was performed in a semi-automated manner (Adluru et al., 2012). Following spatial normalization of the single-macaque DTI images to monkey MNI space (Frey et al., 2011) using DTI-TK, an advanced DTI spatial normalization tool (http://www.nitrc.org/projects/dtitk), a DTI template was computed as their Log-Euclidean mean. A brain parcellation with 432 distinct areas in MNI space, defined according to the Paxinos atlas (Paxinos et al., 2009) was used for probabilistic tractography (Behrens et al., 2003). Accordingly, an SC matrix, representing the strength of connectivity between pairs of network nodes (i.e., atlas regions), was estimated.

4.3. Analysis of global synchronization

For the analysis of global synchronization we first extracted discrete events from all empirical and modeled data by assigning a time stamp to each positive peak of the z-scored data, similar to previous studies (Deco et al., 2017b; Tagliazucchi et al., 2012). As a control we also repeated subsequent analysis on events that were based on negative peaks, which yielded the same results (Supplementary Fig. 1A-B). Next, we chunked the data into sliding windows with different sizes ranging between 5 and 50 TR. The window was advanced in steps of 1TR. To quantify global synchronization, we took advantage of the Fano factor,
which was used in classical point process analysis to assess the presence of globally synchronized spiking activity in E-I networks (Brunel and Hakim, 1999; Hahn et al., 2017; Kumar et al., 2008). To this end, we calculated the sum of all events found within 1 TR of a window, which resulted in a sequence of counts with length equal to the window size. The Fano factor (FF) of the count sequence was then derived based on the following relationship:

$$ FF = \frac{\text{var(counts)}}{\text{mean(counts)}} $$

(1)

The value of the Fano factor gives insight into the correlation structure of the events within each time window. FF is 1 indicates that all the events appear independently from each other and seem to have been drawn from a random Poisson process (Fig. 1F). In the presence of large-scale synchronization of events across many brain areas, the FF increases strongly (FF > 1), as the synchrony now confines most events to a short period of time (1–2 TRs), which is followed by a period with only few events due to synchronization of the negative peaks. FFs were computed for each time window and summarized in a FF distribution. To quantify the overall level of global synchronization, we fitted a Gamma distribution which is defined as follows:

$$ N(F, \beta, \theta) = \frac{e^{-\theta/F}}{\Gamma(\beta)} \theta^{\beta}; \beta > 0, \theta > 0 $$

(2)

where $\beta$ denotes the shape parameter, $\theta$ the scale parameter and $\Gamma$ represents the gamma function. Subsequently, we used the shape parameter $\beta$ to quantify the level of consciousness by assigning a single number to each condition that reflects the overall distribution of large-scale synchrony.

4.4. Statistical analysis

To test for significant differences between beta values and other metrics across different consciousness conditions at the group level, we applied a jackknife approach and created N datasets (N is number of subjects in humans and trials in monkeys) for each vigilance condition, each time concatenating N-1 subjects/trials and leave one out. The new datasets were then assessed using a one-way repeated measure (rm) ANOVA and posthoc tests with Bonferroni correction. The significance level was set to $p < 0.05$. In case the sphericity ($\varepsilon$) assumption of the rm-ANOVA was violated, as assessed by a Mauchly test, the $p$-values and degrees of freedom were adjusted by the Huynh-Feldt estimate of $\varepsilon$. To compute the goodness of fit for the gamma distributions fitted to the Fano factor distributions, we used a one-sample Kolmogorov-Smirnov test.

4.5. Functional connectivity

To quantify statistical dependencies between empirical and modeled fMRI time series, we calculated the Pearson correlation coefficient across all brain area pairs, which were subsequently averaged for each consciousness condition or Fano factor state.

4.6. SC-FC relationship

The structure-function relationship for each consciousness condition was assessed by computing the Pearson correlation coefficient between the structural and functional connectivity matrices (human and monkey) for different consciousness levels or Fano factor states.

4.7. Integration

We used an integration measure based on the largest connected component of the FC matrix, as defined previously (Deco et al., 2015). As a first step, the FC matrix was binarized by setting connections above a threshold $\theta$ to 1 and discarding connections below the threshold, which has a range between 0 and 1. We then searched for the largest connected subgraph, i.e. the nodes that are connected without further connections to the remaining nodes of the entire matrix. The size of the largest component (i.e. the number of nodes pertaining to the subgraph) was calculated for each $\theta$ and integrated across all tested thresholds. To obtain integration values between 0 and 1, the result was normalized by the number of areas N and integration steps. This integration measure was calculated for all consciousness levels and Fano factor states.

4.8. Modularity

The activity during different consciousness levels (or Fano factor states) is organized into functional communities. Therefore, to quantify the level of segregation in the functional connectivity (FC) matrices we performed a community detection procedure on them. Partitioning the networks into clusters was performed using the Louvain method, which optimizes the weighted Newman modularity (Newman, 2006). This measure evaluates the “goodness” of the partition, giving insights to what extent the FC matrices can be broken down into nonoverlapping subnetworks. Optimization of modularity implies that the connectivity strength within a group of brain areas is maximized, while the strength between different groups is minimized. Thus, modularity provides a quantitative indication of the level of functional segregation in the FC. This metric was applied to different consciousness conditions and Fano factor states.

4.9. Computational whole-brain model

To study the mechanisms behind global synchronization, we employed a whole-brain model consisting of either 90 nodes, mimicking the parcellation of areas in human data, or 432 nodes, corresponding to the empirical monkey parcellation. The modeled brain areas were connected based on the patterns of long-range connections that were extracted by DTI from humans and monkeys. We used the measured fiber density as a proxy for the connectivity strength in the model. To simulate the dynamics of the local fMRI signal, we used the normal-form of the Hopf bifurcation (Deco et al., 2017a, 2017b; Hahn et al., 2019), which corresponds to a transition from a focus equilibrium with damped, noisy oscillations to a limit cycle behavior with self-sustained oscillations, similar to the behavior of networks of excitatory and inhibitory neurons (Brunel, 2000). The dynamics is described by the following set of coupled differential equations:

$$ \frac{dx_j}{dt} = \frac{dRe(z_j)}{dt} = \left[ x_j - x_j^2 - y_j^2 \right] x_j - \omega y_j + G \sum_i C_{ji} (x_i - x_j) + \beta \eta_j(t) $$

(3)

$$ \frac{dy_j}{dt} = \frac{dIm(z_j)}{dt} = \left[ x_j - x_j^2 - y_j^2 \right] y_j + \omega x_j + G \sum_i C_{ji} (y_i - y_j) + \beta \eta_j(t) $$

(4)

where $\beta \eta_j$ represents additive Gaussian noise with standard deviation $\beta = 0.01, \omega_j$ is the intrinsic node frequency, which was translated to the BOLD frequency $f_j$ via the following expression: $f_j = \frac{\omega_j}{\pi}$. The BOLD frequency for each model node was set to 0.085 Hz and filtered between 0.075 and 0.095 Hz in the human data simulations, or to 0.04 Hz in the monkey model with filtering between 0.03 and 0.05 Hz. These filters were necessary to remove high frequency noise imposed by the Gaussian noise and allow comparison with empirical Fano factor distributions. These model frequencies were chosen, since the empirical data filtered around the same frequency bands can reproduce the same Fano factor distributions that are seen with a more broadband filtering (0.01 – 0.1 Hz). The variable $x_j$ was used to model the local fMRI signal. The parameter G scales each connection in the anatomical connectivity matrix by a constant value, which was set to 0.1 in this study. For each consciousness level we performed a simulation of 100 min. For the single-trial analysis we simulated 18 trials of 250 TR for each consciousness state of the human model. For the monkey model we simulated 21, 26 and 31 trials of 250 TR for the awake, light and deep anesthesia case.
respectively. These simulations were compared with ten long trials of a duration of 10,000 TR.

In an unconnected node, the dynamical behavior is determined by the bifurcation parameter $a_j$ which exhibits a supercritical Hopf bifurcation at $a_j = 0$. This means that the activity transitions from a stable focus equilibrium, governed by resonance and noisy oscillations at $a_j < 0$, to a stable limit cycle at $a_j > 0$ with self-sustained oscillations. However, as soon as a node is embedded in a large scale network, the impinging connections change the bifurcation point to negative values. In order to find $a_j$ at the bifurcation for the large-scale anatomical network of the human and monkey DTI data, we took advantage of the fact that the Fano factor distributions (see above) for small window sizes (5 TR), followed an exponential distribution when $a_j$ was set below the bifurcation (Fig. 2B). As soon as $a_j >$ bifurcation, the network produced global oscillations, which was visible as a bump in the FF distribution and thus indicated a deviation from the exponential FF decay. We thus defined the bifurcation point as the value of $a_j$, where the exponential fit, as assessed by a Kolmogorov-Smirnov distance, increased to values above 0.1 (Fig. 2C). The $a_j$ value at the bifurcation point was $\sim 0.66$ for the human and $\sim 0.33$ for the monkey. SC matrices where then normalized to zero (Fig. 2D-G). To fit the model to the data, we performed a grid search to find $a_j$, where the distance $|\hat{\beta}_\text{emp} - \hat{\beta}_\text{mod}|$ of the empirical and modeled FF distributions was minimized.

4.10. Model lesion analysis

To study the role of individual brain areas for model dynamics and global synchronization, we conducted lesion simulations by setting the bifurcation parameter of single nodes $j$ to a value of $a_j = -2.65$ in humans and $a_j = -1.31$ for monkeys, while keeping the other nodes at values that correspond to the awake state ($\sim 0.663$ for humans, $\sim 0.329$ for monkeys). The impact of a single node lesion on global synchronization was then quantified by the lesion index $\lambda$, which we defined as follows:

$$\lambda = \frac{\hat{\beta}_{\text{awake}}}{\hat{\beta}_{\text{lesion}}}$$

where $\hat{\beta}_{\text{awake}}$ represents the beta value of the FF distribution found in the awake state (human or monkey) and $\hat{\beta}_{\text{lesion}}$ indicates the beta value after the node lesion with a simulation time of 10 min. The lesion index $\lambda$ ranges between 0 and 1, and reflects the fraction of global synchrony after the lesion as compared to the awake state in either monkey or human data models. Thus, $\lambda \sim 1$ indicates that the global synchronization is barely affected by the lesion, while $\lambda < 1$ is the result of a strong influence of a given node on global dynamics.

4.11. Graph theoretical analysis

The human and monkey SC matrices were studied using several graph analysis tools. A rich club was identified in both SC networks. Rich-club identification implies the calculation of the $k$-density, a recursive calculation that computes the density of the remaining network after all nodes with degree $k' \leq k$ have been removed, until no nodes are left (Zamora-López et al., 2010). The case $k = 0$ corresponds to the density of the original network. If the $k$-density grows with $k$, it means that the nodes with largest degree are densely interconnected with each other. If the $k$-density decays, it means that the nodes with largest degree are indeed not connected with each other. The two SC networks display a growing $k$-density function and clearly contain a rich-club, taken as the remaining subset of hubs with node degree $k' > 47$ in humans and $k' > 108$ in monkeys.

Communicability (M) is a graph metric that quantifies the “influence” one node exerts over another given the path structure of the network ( Estrada and Hatano, 2008). It accounts for the fact that (i) information flows along all possible paths, not only across the shortest ones, and (ii) that shorter paths are more relevant, while they influence decays with the length of the path (Bettinardi et al., 2017). Given the adjacency matrix $A$ of the matrix, communicability is defined as the matrix exponential, $e^A$, which can be decomposed as the power series:

$$C = e^A = 1 + A + \frac{A^2}{2!} + \frac{A^3}{3!} + \frac{A^4}{4!} + \ldots,$$

where the powers $(A^n)_{ij}$ represent the number of paths of length $n$ that run from node $i$ to node $j$. The communicability of every node, taken as the column sum of the matrix $C$, was then correlated with the lesion index.

To quantify the asymmetry of communicability between the two hemispheres, we first computed communicability of a given node for the right ($r$) and left hemisphere ($l$) separately and then calculated the asymmetry ($S$) according to the following expression:

$$S_{rl} = \frac{|M_l - M_r|}{M_l + M_r}$$

Asymmetry values range between 0 and 1, such that $S_{rl} \sim 1$ indicates full asymmetry and $S_{rl} \sim 0$ complete symmetry.

Declaration of Competing Interest

The authors declare no competing financial interests.

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Author contributions

G.H. and G.D. designed the computational study. G.H. and G.Z. analyzed the data. G.H. performed the numerical simulations. H.L. designed the EEG-fMRI experiment and with E.T. collected the EEG and BOLD-fMRI human data. M.L.K. provided the human DTI data. L.U. and B.J. collected and preprocessed the monkey fMRI data. D.M. provided the monkey DTI data. G.H. wrote the first draft of the paper, which was finalized by all the remaining authors.

Supplementary materials

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