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Connectivity changes underlying spectral EEG changes during propofol-induced loss of consciousness

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Abstract

The mechanisms underlying anesthesia-induced loss of consciousness remain a matter of debate. Recent electrophysiological reports suggest that while initial propofol infusion provokes an increase in fast rhythms (from beta to gamma range), slow activity (delta to alpha) rises selectively during loss of consciousness. Dynamic causal modeling was used to investigate the neural mechanisms mediating these changes in spectral power in humans. We analyzed sourcereconstructed data from frontal and parietal cortices during normal wakefulness, propofol-induced mild sedation and loss of consciousness. Bayesian model selection revealed that the best model for explaining spectral changes across the three states involved changes in cortico-thalamic interactions. Compared to wakefulness, mild sedation was accounted for by an increase in thalamic excitability, which did not further increase during loss of consciousness. In contrast, loss of consciousness *per se* was accompanied by a decrease in backward cortico-cortical connectivity from frontal to parietal cortices, while thalamo-cortical connectivity remained unchanged. These results emphasize the importance of recurrent cortico-cortical communication in the maintenance of consciousness and suggest a direct effect of propofol on cortical dynamics.

Introduction

The mechanisms underlying anesthesia-induced loss of consciousness (LOC) are unknown. Previous neuroimaging studies performed during anesthesia have shown a breakdown of connectivity in fronto-parietal networks (Lee et al., 2009; Boveroux et al., 2010). Whether this decreased connectivity is due to a direct cortical action of anesthetics, or to a disruption of cortico-thalamo-cortical connectivity, remains unclear (Boveroux et al., 2010). Some authors suggest that anesthetics may act by depressing thalamic function, impairing its role either as an arousal promoter, or as a cortical read-out (Alkire et al., 2008). Many studies have reported a reduction of thalamic metabolism and blood flow during anesthesia (Fiset et

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al., 1999; Alkire and Miller, 2005; Alkire et al., 2008). However, during the first few minutes of propofol-induced LOC, thalamic activity seems to be preserved, while cortical activity is depressed (Velly et al., 2007). It has also been suggested that anesthetic effects on the thalamus may be largely indirect (Alkire et al., 2000; Schiff and Plum, 2000; Franks, 2008). Indeed, the metabolic and electrophysiological effects of anesthetics on the thalamus in animals are abolished by removal of the cortex (Nakakimura et al., 1988; Angel, 1993; Vahle-Hinz et al., 2007). In short, the thalamus versus cortex debate remains active in the anesthesia literature (Alkire et al., 2008).

A recent study investigated EEG correlates of propofol-induced LOC in man (Murphy et al., 2011b). Mild sedation (with a slower response to command but preserved consciousness) was characterized by a rise in high frequency power in scalp EEG. Conversely, LOC was paired with the appearance of high-amplitude delta waves, on a persistent fast rhythmic background (Murphy et al., 2011b). Furthermore, the same default network areas - i.e. precuneus / posterior cingulate cortex (PCC) and medial prefrontal / anterior cingulate cortex (ACC) - appeared to support both travelling slow waves and high frequency EEG power increases. This observation is puzzling, because it is thought that the generation of fast and slow frequencies are mediated by opposing cellular mechanisms (Steriade, 2006): fast rhythms are produced when neurons are in a depolarized 'awake' state, while slow rhythms are generated when neurons are hyperpolarized. The simultaneous presence of both activity modes in the same EEG (as well as their distinct profile of emergence across levels of sedation) therefore suggests distinct mechanisms in their generation.

The present study investigated the neural mechanisms underlying EEG spectral power changes observed during propofol-induced unconsciousness. This enquiry has been enabled by recent advances in the analysis of EEG data, specifically, dynamic causal modeling (DCM) (Friston et al., 2003). DCM uses a biologically informed generative model that embodies known constraints on the generation of EEG signals (Kiebel et al., 2009). Bayesian model inversion and comparison then provide a method for testing candidate models of causal interactions among brain sources in a distributed network. In particular, DCM for steady-state responses (SSR) allows one to explain neural mechanisms underlying the differences in spontaneous EEG power spectra (Moran et al., 2009) and was used here to quantify changes in neural coupling during propofol-induced unconsciousness.

Material and methods

Data acquisition

The data used in this study have been reported previously in (Murphy et al., 2011a) in an analysis of functional connectivity using coherence measures. In brief, the study was approved by the Ethics Committee of the Faculty of Medicine of the University of Liege and written informed consent was obtained from all participants. None of the subjects had a history of head trauma or surgery, mental illness, drug addiction, asthma, motion sickness, or previous problems during anesthesia. Fifteen minute spontaneous 256-electrode highdensity EEG recordings were acquired in eight subjects (mean age 22 ± 2 y, 4 males) in four different states: normal wakefulness, sedation (slower response to command, Ramsay scale score 3), LOC with clinical unconsciousness (no response to command, Ramsay scale score 5), and recovery of consciousness (Ramsay et al., 1974). Subjects were considered fully awake or to have recovered consciousness if the response to verbal command ('squeeze my hand') was clear and strong (Ramsay 2), in sedation if the response to verbal command was clear but slow (Ramsay 3) and in LOC, if there was no response (Ramsay 5-6). For each consciousness level assessment, Ramsay scale verbal commands were repeated twice. The most comfortable supine position was used to avoid painful stimulation related to position. All recordings were performed with eyes closed.

Propofol was infused through an intravenous catheter placed into a vein of the right hand or forearm. Throughout the study, the subjects breathed spontaneously and additional oxygen (5 liter min⁻¹) was given through a loosely fitting plastic face-mask. Computer-controlled intravenous infusion of propofol (Alaris® TIVA) was used to obtain constant effect-site concentrations. The propofol plasma and effect-site concentrations were estimated using a three-compartment pharmacokinetic model (Marsh et al., 1991). Average arterial blood concentrations of propofol were $1.91 \pm .52 \text{ mcg/ml}$ for sedation and $3.87 \pm 1.39 \text{ mcg/ml}$ for LOC.

Data pre-processing

Thirty seconds artifact-free EEG epochs from each condition were selected for analysis. The raw signals were filtered from 0.5-40 Hz. A standard set of electrodes was co-registered to the magnetic resonance image of an individual whose head closely approximates the Montreal Neurological Institute (MNI) space. The electromagnetic forward model comprised four nested spheres. The inverse matrix was calculated using a constrained, depth and orientation weighted, truncated singular value decomposition regularized (10^{-3}) L2 minimum norm (Grave de Peralta Menendez et al., 2001). The source space was a set of 2447 cortical voxels (7 mm³) that were identified based on the MNI probabilistic brain atlas. The source space was restricted to the cortex because cortical pyramidal neurons are the most likely generators of the EEG (Kirschstein and Kohling, 2009). For the power analysis, we source localized about five minutes of artifact free EEG from each subject in each condition. Source modeling was performed in GeoSource (Electrical Geodesics Incorporated), while all other calculations were performed in MATLAB (The Math Works). From this source reconstruction we selected two regions of interest based on a previous study showing increased gamma power during LOC compared to wakefulness (Murphy et al., 2011b). Figure 1 shows group mean source-level power spectra and cross spectral densities (in semi-log and log-log scales) from the frontal and parietal ROIs during wakefulness, sedation and LOC. SnPM (statistical nonparametric mapping) was used to compare frequency spectra between behavioral states (Nichols and Holmes, 2002). Power spectrum changes statistics were performed on a 1Hz-by 1Hz frequency bin basis and were run in MATLAB (The Mathworks) using SnPM - a permutation-based image analysis scheme that accounts for multiple comparisons (Nichols and Holmes, 2002).

Extraction of frontal and parietal source activity

The rationale of the present work was to investigate changes in a fronto-parietal network that caused changes in the power spectrum observed during two levels of propofol sedation. In DCM, this is modeled explicitly in terms of neuronal subpopulations that influence each other through intrinsic (within area) and extrinsic (between-area) connections. The parameters of these models control the expression of synaptic responses resulting from these influences. Model inversion allows us to estimate these parameters and evaluate the evidence for one model or network architecture, relative to others. In brief, we extracted the source-reconstructed signals from two sources or regions of interest, previously determined to be the major contributors to increases in gamma power (25-40 Hz) during propofolinduced unconsciousness (see above) Increases in gamma power during loss of consciousness are particularly interesting, given that gamma frequency activity is thought to be important for consciousness (Llinas et al., 1998). Source-reconstructed signals were modeled with DCM to characterize the mechanisms underlying changes in power spectra across the three states. Bayesian model selection compared the evidence of a model with exclusively cortical sources to that of two other thalamo-cortical models. These thalamocortical models comprised one or two thalamic sources connected reciprocally with the cortical sources. Having established the best model, we then quantified the changes in neural excitability and connectivity with sedation and LOC.

For DCM, we summarized the activity of each region (frontal and parietal) using its principle eigenvariate. The first eigenvariate of frontal and parietal activity was calculated from 9 dipolar sources (vertices) per region, over a 30 second period in each condition. These were the time series used for subsequent dynamic causal modeling: DCM uses an inbuilt AR process to compute a regularized frequency domain representation (found in the spectral toolbox of SPM8 http://www.fil.ion.ucl.ac.uk) from the time series. We evaluated three DCMs for each subject that explained differences among the conditions (wakefulness, sedation and LOC) in terms of changes in intrinsic and extrinsic coupling.

Network models or hypotheses

DCM is usually used in a hypothesis-driven fashion: it does not explore all possible models but tests specific mechanistic hypotheses (Boly et al., 2011). Here, we examined three models, corresponding to hypotheses about the neural mechanisms underlying the spectral changes observed during propofol-induced loss of consciousness (Figure 2.A). Our first model comprised two cortical areas, connected by forward and backward connections. A second model incorporated a (hidden) thalamic source, reciprocally connected to both cortical areas (i.e. no data were reconstructed in this region). The third model used two thalamic modules, with a similar architecture, but separately connected to each cortical source.

Dynamic causal modeling

The idea behind DCM is that M/EEG data are the response of a dynamic input–output system to endogenous fluctuations or exogenous (experimental) inputs. One models the measured M/EEG as the response of a network of sources, where each source corresponds to a neural mass model of several subpopulations. In DCM for steady-state responses neuronal dynamics are assumed to be driven by endogenous (neuronal) fluctuations with an unknown but parameterized spectral density. This sort of DCM is useful, for example, in sleep research, where there are no external or exogenous stimuli (Moran et al., 2009). We employ these modeling assumptions here, assuming the endogenous neuronal fluctuations were a mixture of pink (one over f) and white noise.

Assuming the system operates in a steady-state around its fixed point, we can linearize the nonlinear differential equations (Figure 2.B) describing its response in the frequency domain. These differential equations depend on the neural mass model of sources in the network and the effective connectivity or coupling among them. This linearization allows one to compute the transfer function mapping from the endogenous (neuronal) fluctuations to the sensors, where the transfer functions are the Fourier transforms of the source's kernels (see Figure 2.B). Applying this transfer function to the spectral density of these inputs provides the predicted spectral activity in each region. This allows us to establish a mapping from the system parameters to the predicted frequency spectrum (Moran et al., 2007). Furthermore, we can model differences between two or more spectra, acquired under different conditions, as a consequence of changes in parameters. These parameters correspond here, to intrinsic or extrinsic connections, where intrinsic connections are controlled by the maximum amplitude of post-synaptic potentials (Kiebel et al., 2007). The basic idea is to manipulate the (real) system (e.g. by changing levels of propofol sedation) and model the experimental effects in terms of changes in coupling. Estimated parameter changes across the three levels of consciousness served as summary statistics in classical tests (analysis of variance and post-hoc *t*-tests) of condition-specific effects over subjects.

In detail, we used a neural mass model for each source, which describes the dynamics of three neuronal subpopulations. Each population has its own (intrinsic) dynamics governed by the neural mass equations: see (Moran et al., 2009), but also receives endogenous input

from surrounding cortex or from other sources. The sources and their interactions are fully specified by first-order differential equations that are formally related to neural mass models used in computational models of M/EEG (Breakspear et al., 2006; Rodrigues et al., 2010). These differential equations describe the evolution of hidden neuronal states (membrane potentials and currents) in the subpopulations comprising each source. For the cortex, these subpopulations are layer specific, with spiny stellate cells in the granular layer that are reciprocally connected to pyramidal cells in infra and supra-granular layers. Inhibitory interneurons in the infra and supra granular layers are, in turn, reciprocally connected to the pyramidal cells. Connections between regions originate and terminate at specific cortical layers. In this way, DCM for steady state responses (SSR) allows one to specify hierarchical neuronal networks with anatomically grounded connectivity rules (Felleman and Van Essen, 1991). Prefrontal and parietal cortices are thought to be hierarchically arranged with laminar specific bidirectional coupling (Jacobson and Trojanowski, 1977; Schwartz and Goldman-Rakic, 1984). Generally, forward connections originate in superficial pyramidal cells (in supragranular layers II and III) and terminate in the granular layer IV (Pandya, 1995; Rouiller and Durif, 2004). In contrast, reciprocal backward connections arise from the deep pyramidal cells (in (infragranular layers V and VI) and terminate outside of layer IV (Winguth and Winer, 1986; Rouiller and Durif, 2004; Moran et al., 2009). While our model lumps together supra and infra-granular pyramidal cells, it accommodates these asymmetries by retaining their major functional characteristics; namely, forward connections arise in pyramidal cells and target spiny stellate cells in layer VI, whereas backward connections target pyramidal and inhibitory cells in supra and infra granular layers. The thalamic architecture was based on the model used by Robinson et al. (Robinson et al., 2001) and Breakspear et al. (Breakspear et al., 2006) in previous characterizations of EEG spectra and seizure activity: thalamic sources comprised excitatory (relay cells) and inhibitory (reticular cells) subpopulations that were reciprocally connected. The thalamus sent forward connections to both cortical areas and received backward connections. We assume that the depolarization of pyramidal cell populations generates observed M/EEG data. In our case (because we used source reconstructed activity), there was a direct mapping between the activity of pyramidal cells and the virtual sensor in the two cortical sources.

Bayesian Model Inversion and Estimation

Model inversion means fitting the model to the empirical data; here the cross spectral densities from the cortical sources. In DCM a Variational Bayesian scheme called Variational Laplace (Friston et al., 2007) is used to approximate the conditional or posterior density $q(\theta)$, Over parameters by maximizing the negative free energy

 $F = \ln p(y|m) - KL[q(\theta), p(\theta|y, m)]$ (1)

Here KL is the divergence between the true $p(\theta | y, m)$ and approximate $q(\theta)$ posteriors. It can be seen from Equation 1 that the negative free energy is also a lower bound on the log model evidence, ln p(y/m) (the probability of the data y given a model m, see (Friston and Stephan, 2007) for details. This quantity is used for model selection when testing a series of candidate architectures; in this case a model with cortical sources was compared with two thalamo-cortical models. A random effects Bayesian model selection (BMS) analysis was performed (Stephan et al., 2009) to select the best model. The maximum a posteriori (or conditional mean) of the parameters encoding changes in connectivity then serve to quantify differences among experimental conditions. We modeled the three condition-specific effects, corresponding to wakefulness, sedation and LOC, in each individual DCM. Effects were modeled in terms of changes (relative to the first condition) in the intrinsic connectivity (excitability) in all sources and the extrinsic connectivity between sources. To assess the consistency and significance of these effects over subjects, subject-specific

changes were entered into an ANOVA with anesthetic depth as a within subject factor. Twotailed probabilities at p < 0.05 (Bonferroni corrected for multiple comparisons, i.e. for 9 tests) were used to assess the significance of changes in parameters. Post-hoc *t*-tests were then used to investigate the directionality of effects over conditions. An additional analysis was also performed on Model 2 parameters to test for an interaction between backward versus forward cortico-cortical connectivity changes during propofol sedation. To verify our inference, we performed two additional ANOVA analyses on Model 1 parameters, searching for 1) an effect of level of consciousness on backward connection strength and 2) an interaction between backward versus forward cortico-cortical connectivity changes during propofol sedation.

Results

The two regions of interest we investigated were located in precuneus/posterior cingulate and in medial prefrontal cortex. While initial propofol infusion was accompanied by an increase in fast frequencies in the beta-gamma range, both regions showed a further significant increase in slow frequency power (delta to alpha range) with the emergence of clinical unconsciousness (Figure 1). To characterize the neuronal mechanisms underlying these changes in power spectrum, we extracted the time-series of these two regions during wakefulness, mild sedation and LOC and modeled them with DCM, under steady-state assumptions. We inverted three models for each subject (Figure 2.A). Model 1 comprised the two cortical areas that were connected reciprocally by backward and forward extrinsic connections. In Model 2, a thalamic source was added (Robinson et al., 2001; Breakspear et al., 2006), connected reciprocally to both cortical areas by forward (thalamo-cortical) and backward (cortico-thalamic) connections (Figure 2.B). Model 3 comprised the two cortical areas, each connected reciprocally with its own thalamic region. Bayesian model comparison revealed that the best model for explaining the data across subjects and conditions was Model 2 (a cortico-thalamic network with a common thalamic source) with a 92% posterior probability. Figure 3 shows individual fits between predicted and observed source-level power spectra and cross spectral densities as computed for Model 2 by DCM.

In order to quantify changes in connectivity and excitability across conditions, we then examined the parameters of the best model (Model 2). Classical inference on the parameter estimates for each subject revealed a significant effect of condition on specific model parameters: one effect showed significant changes in thalamic excitability (Bonferroni corrected p = 0.01) and the other a change in backward cortico-cortical connectivity from frontal to parietal cortices (Bonferroni corrected p = 0.036, Figure 4, left panel). Post-hoc analyses revealed that thalamic excitability significantly increased during mild sedation as compared to wakefulness; and that this hyper-excitability persisted during loss of consciousness (Figure 4, right panel). Hence thalamic excitability mirrored the changes in fast rhythms that accompany propofol infusion but did not change with LOC. On the other hand, backward cortico-cortical connectivity was preserved during mild sedation, but showed a significant reduction with loss of consciousness - thus following the expression of slow power changes in EEG (Figure 4, right panel). We observed a significant interaction between backward and forward connectivity changes with propofol-induced sedation stages (p = 0.006). Thalamo-cortical connectivity and cortical excitability did not show significant changes across sedation stages. A complementary analysis of Model 1 (cortico-cortical model) parameters confirmed a reduction in backward connections during propofol-induced loss of consciousness (effect of the level of consciousness on backward connectivity: p =0.022). We also reproduced the significant interaction between backward and forward connectivity changes during propofol sedation (p = 0.026) under this model.

Discussion

To the extent that the models of thalamo-cortical interactions examined here are representative, DCM indicates that propofol-induced LOC is associated with a selective breakdown of cortico-cortical backward connectivity, while thalamo-cortical connectivity, thalamic excitability and cortical excitability are preserved. These results extend previous studies reporting decreased cerebral connectivity during anesthesia-induced LOC using resting state fMRI (Martuzzi et al.; Boveroux et al., 2010) or TMS-EEG (Ferrarelli et al., 2010). They suggest that propofol-induced LOC is due to impairment of direct cortical interactions, rather than to a thalamic inhibiting effect (Alkire et al., 2008; Franks, 2008). There are several issues that previous studies of propofol and *functional* connectivity could not resolve. Previous resting state fMRI studies could not differentiate between impaired thalamo-cortical or cortico-cortical connectivity, or between an impairment of backward versus forward connections (Boveroux et al., 2010). Previous EEG functional connectivity studies (Cimenser et al., 2011; Murphy et al., 2011a; Supp et al., 2011) provided contradictory results, suggesting increased coherence during propofol-induced LOC. The present study resolves these issues, through the analysis of *directed effective* connectivity using DCM. Our results support the view that propofol-induced LOC may be linked to an impairment of direct communication between cortical areas. This selective cortical effect is in line with previous studies reporting aroused thalamic EEG activity, but marked cortical slowing during propofol-induced LOC (Velly et al., 2007) or spike-wave seizures (Englot et al., 2009). Since bilateral thalamic lesions can cause vegetative state (Adams et al., 2000), thalamic activity might be necessary, but not sufficient, for consciousness. The common denominator in the neural correlates of consciousness might rather be changes in distributed cortico-cortical processing.

According to the DCM conducted here, during propofol-induced LOC, backward corticocortical connectivity was impaired, while forward connectivity was preserved. Our findings do not point to a global decrease in connectivity. The significant interaction between backward and forward cortico-cortical connectivity changes during propofol-induced LOC rather suggests a selective impairment of backward connections, in the context of preserved forward connectivity. Our results are in line with previous scalp EEG studies reporting loss of front-to-back connectivity during propofol-induced sedation (Lee et al., 2009; Ku et al., 2011). The present finding supports a role for recurrent processing in higher-order cortices in conscious perception (Lamme, 2006; Boly et al., 2011). It is probably the case that both forward and backward connections are important for consciousness (Boly et al., 2011). Furthermore, the present results do not preclude more widespread physiological changes under other anesthetics. Rather, they suggest that backward connectivity in higher-order associative cortex may be a critical bottleneck for anomalies of conscious awareness.

Increases in the power of high frequencies, present from the start of propofol infusion, appeared to be linked to thalamic hyper-excitability, broadcasting abnormal rhythms to the cortex. According to DCM, thalamic excitability increased during both sedation and LOC as compared to wakefulness. BMS suggested that a thalamic source interconnected with both cortical regions was the best model to explain our data. If thalamic involvement was restricted to specific nuclei (such as the pulvinar or geniculate, mainly connected to posterior cortex, or the anterior thalamus, mainly connected to anterior cortex), Model 3 would have won in BMS. DCM allows a principled comparison of different models (Penny et al., 2004), where the best model is not necessarily the most complex: in our case, the second model had greater evidence despite the fact it was simpler (had fewer free parameters) than the third model. Some studies show interconnections between PCC, ACC and associative thalamic nuclei in humans; e.g. the centromedial thalamus (Eckert et al., 2011). In monkeys, calbindin-positive matrix cells, concentrated within intralaminar

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thalamic nuclei, project diffusely to superficial layers of cortex (Jones, 2002). Cells in intralaminar nuclei can fire at high frequencies (Alkire et al., 2008). Thalamic hyperexcitability is in line with reports showing that propofol increases the discharge probability of thalamic relay cells (Angel and LeBeau, 1992; Bonhomme et al., 2001) and with a recent modeling study suggesting that increased thalamo-cortical coherence could explain propofol-induced changes in power spectra, especially in the alpha range (Ching et al., 2010). The present study highlights how DCM can separate mixed power spectrum patterns, and provide a biophysically grounded explanation for them. Here, the two different phenomena expressed in the changes of power spectra (in the low and high frequencies) were related to two distinct neuronal mechanisms.

Several technical issues of the present work deserve discussion. DCM is a hypothesis-driven method: it does not explore all possible models but tests specific mechanistic hypotheses (Boly et al., 2011). Results of DCM studies are only valid within the space of the models tested. Our three models correspond to three hypotheses about generic changes in brain connectivity during propofol-induced LOC. The presence of high delta power in the EEG may reflect the expression of neuronal up and down states at smaller temporal and spatial scales; DCM for SSR however focuses on population dynamics under steady-state assumptions; i.e., condition-specific changes in power spectra (Moran et al., 2009). Statedependent coupling in DCM is modeled using nonlinearities, which can be thought of as mediating activity-dependent changes in synaptic efficacy. In DCM for SSR these nonlinearities are fairly simple and reduce to a sigmoid activation function for each neuronal subpopulation (Moran et al., 2009). In the present case, this model captures most changes in power present in source-level EEG (see Figure 3). The implicit mesoscopic characterization of synaptic responses (Moran et al., 2007; Moran et al., 2009) provided a reasonable account of electrophysiological data in a number of validation studies in humans or animals (Moran et al., 2008; Moran et al., 2011b; Moran et al., 2011a).

We used source-reconstructed EEG responses, obtained using L2-norm algorithm (Murphy et al., 2009), to access frontal and parietal signals under the different anesthetic conditions. The present study thus does not attempt to fully account for scalp EEG changes during propofol sedation, but rather accounts for selective changes in a fronto-parietal network (encompassing ACC and PCC regions) that may represent propofol-induced changes in thalamo-cortical interactions. However, we cannot exclude the possibility that other networks may have shown different changes during propofol sedation. Our rationale was to choose two regions with demonstrable differences in power between wakefulness and propofol-induced LOC - here the ACC and PCC - to model with DCM. Furthermore, numerous studies have implicated the default network (Martuzzi et al.; Boly et al., 2009; Boveroux et al., 2010; Vanhaudenhuyse et al., 2010; Samann et al., 2011) and PCC (Laureys et al., 2004; Vogt and Laureys, 2005; Boveroux et al., 2008; Vanhaudenhuyse et al., 2010) in consciousness. Finally, the choice of ACC and PCC allowed us to compare our DCM results to the coherence analyses reported in (Murphy et al., 2011a). As source-reconstructed EEG signal-to-noise ratio decreases with sources depth (Goldenholz et al., 2009), we chose to model a thalamic third driver as a hidden source in our analysis, rather than to extract the thalamic signal itself - an approach previously used in a number of EEG computational modeling studies (Breakspear et al., 2006; Ching et al., 2010; Victor et al., 2011a). The architecture of this hidden source was chosen to model reticular and relay nuclei in the thalamus (as in (Robinson et al., 2001; Breakspear et al., 2006)). Without actual data from thalamus, we can only infer the presence of a thalamic relay. However, it is possible that this hidden source could reflect other brainstem contributions. Having said this, cortical-basal ganglia loops outputs involve the thalamus (Herrero et al., 2002; Redgrave et al., 2010) - in this context, a modulatory effect mediated by basal ganglia would be accounted for by a change in cortico-thalamic interactions our model. Our hypothesis that the thalamus was

involved in propofol-induced changes is motivated by several studies suggesting a role for the thalamus in anesthesia-induced LOC (Alkire and Miller, 2005; Alkire et al., 2008). It also appeals to previous studies using a thalamo-cortical model to explain changes in EEG in pathological (Victor et al., 2011b) or pharmacological (Ching et al., 2010) LOC. Further work in animals (as in (Imas et al., 2005b, a)) may provide direct thalamic measurements to corroborate the present findings.

Beyond the implications for the mechanisms underlying propofol-induced sedation, our results may provide further evidence for a key role of backward connections in the maintenance of consciousness. Backward connections seem to be impaired in the vegetative state (Boly et al., 2011) and under isoflurane sedation (Imas et al., 2006). They are also thought to play a role in normal conscious access (Fahrenfort et al., 2007). The present study suggests that loss of consciousness may be induced by a selective impairment of cortico-cortical backward connectivity, even in the presence of preserved subcortical inputs to the cortex.

In conclusion, our results show that a breakdown of recurrent cortical processing can lead to propofol-induced LOC, even in the presence of preserved thalamus arousal and excitability. The present findings support the hypothesis of a direct cortical effect of propofol anesthetic. In the future, cortico-cortical connectivity measures could be tested as a clinical measure for the depth of propofol sedation.

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Cross-spectral density



Figure 1.

Power spectrum changes across vigilance states in two sources showing increased gamma power in LOC, compared to wakefulness (p < 0.05, corrected using SnPM). The frontal source or region of interest (ROI) is a portion of the anterior cingulate and the posterior ROI is a portion of the posterior cingulate. Semi-logarithmic plots of power spectra from each region and cross-spectral density between the two regions are displayed in waking (W, black), mild sedation (MS, blue) and LOC (red); the inserts are the corresponding log-log plots. The blue bars in the middle panel show frequency band specific changes in power during the transition from waking to sedation and then to LOC (when considered significant

at p<0.05, corrected using SnPM). W: wakefulness, MS: mild sedation, LOC: loss of consciousness.



Figure 2.

A. Model specification: the three DCMs used for Bayesian model comparison. Sources in each model receive endogenous neuronal fluctuations or innovations (modeled as a mixture of pink and white noise) that elicit neural responses characterized by a power spectrum that depends upon the model parameters. Random effects Bayesian model selection showed that the model with a common thalamic source (Model 2) had the greatest evidence (with a 92% posterior probability). Model 2 was thus selected for subsequent quantitative analysis of effective connectivity changes across the three experimental conditions. B. Equations of motion describing neuronal interactions in thalamic, posterior cingulate and anterior cingulate regions. These dynamics are based on a neural-mass model that has been used

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extensively in the causal modeling of electromagnetic data (Jansen and Rit, 1995; David and Friston, 2003; David et al., 2005; Moran et al., 2008). It represents a minimal description of synaptic processing in multiple populations, in terms of synaptic convolution of pre-synaptic inputs and a non-linear mapping between the resulting depolarization and firing rates. The kernels, from which we derive the transfer functions, obtain analytically from the Jacobian $\Im = f(x, u, \theta)$ of hidden neuronal states, x(t) and a mapping (forward model) $s(x, \theta) : x \rightarrow s$ that couples hidden states to the observed signals in PCC and ACC. Neuronal fluctuations or input are denoted by u(t). For region *i*, and input or innovation *k*, the kernel is

$$\kappa_{i}^{k}(\tau,\theta) = \frac{\partial s_{i}(t)}{\partial u_{k}(t-\tau)} \\ = \frac{\partial s_{i}(t)}{\partial g(t)} \frac{\partial g(t)}{\partial x(t)} \frac{\partial x(t)}{\partial x(t-\tau)} \frac{\partial x(t-\tau)}{\partial \dot{x}(t-\tau)} \frac{\partial \dot{x}(t-\tau)}{\partial u_{k}(t-\tau)} \\ = \frac{\partial g_{i}}{\partial x} \exp\left(\Im\tau\right) \Im^{-1} \frac{\partial f}{\partial u_{k}}$$

The Jacobian is augmented by a matrix of inter-regional delays D, $\Im \leftarrow (I + D\Im)^{-1}\Im$. The kernels are hence functions of the model's equations of motion and output mapping. The output here is determined largely by pyramidal cell activity (~80%) in each cortical area. See Tables 1 & 2 of (Moran et al., 2009) for the prior values of model parameters.

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Figure 3.

This figure shows that the single subject model fits (solid lines) for source-level power spectra and cross spectral densities as computed by DCM could satisfyingly account for the original power spectrum data (in dashed lines). For each subject (S1-8), wakefulness, mild sedation and LOC data are displayed respectively in black, blue and red. Model fits are provided both as semi-log and log-log plots. CSD: cross-spectral density.



Figure 4.

Connectivity parameter analysis of the winning model (i.e. Model 2, the model with a common thalamic source). Left panel. Quantitative parameter analysis revealed two effects of changes in conscious level on network parameters (p<0.05, represented in dark red): changes in thalamic (T) excitability and changes in backward fronto-parietal (F-P) connectivity. Right panel. Average change in thalamic excitability and fronto-parietal backward connectivity parameters over subjects (with standard errors shown as red bars) for mild sedation (MS) and LOC as compared to wakefulness. Post-hoc analyses revealed that changes in thalamic excitability were already present in mild sedation stage as compared to wakefulness (p<0.05), and persisted during LOC as compared to wakefulness (p<0.05), but without a significant further increase as compared to mild sedation. In contrast, backward fronto-parietal connectivity was not significantly modified during mild sedation, but showed a selective decrease (p<0.05) during LOC as compared to wakefulness. The (adimensional) changes are relative to the value in the first (waking) condition. T: thalamus, F: frontal, P: parietal, MS: mild sedation, LOC: loss of consciousness.