Effects of transient unilateral functional brain disruption on global neural network status in rats: a methods paper

Willem M. Otte1,2*,†, Kajo van der Marel1†, Kees P. Braun2 and Rick M. Dijkhuizen1,3

1 Biomedical MR Imaging and Spectroscopy Group, Image Sciences Institute, University Medical Center Utrecht, Utrecht, Netherlands
2 Department of Pediatric Neurology, Brain Center Rudolf Magnus, University Medical Center Utrecht, Utrecht, Netherlands
3 Molecular Imaging Program at Stanford, Stanford University, Stanford, CA, USA

INTRODUCTION

Focal brain damage, such as after ischemic stroke or in epilepsy, not only affects local neuronal networks, but can also have significant effects on the function of remote brain regions. For instance, constrained behavioral output (Carter et al., 2010) and cognitive comorbidities (Tatemichi et al., 1994; Motamedi and Meador, 2003) after stroke and epilepsy have partly been attributed to functional disturbances of distant brain regions that were not primary affected. In addition, spontaneous partial recovery of sensorimotor and behavior output in the weeks and months following stroke may be explained by functional plasticity of remote and unaffected brain regions (Langhorne et al., 2009; Stinear, 2010). As brain functions are believed to emerge from interacting regions within large-scale neural networks, the brain’s intrinsic capacity to compensate for focal injury is linked to reorganization of surviving functional networks at a global level (Grefkes and Fink, 2011).

A sophisticated set of measures, using the concept of graph theory, has been used to describe alterations in functional network organization following focal brain damage (Rubinov and Sporns, 2010), with promising opportunities for applications in clinical settings (Fornito et al., 2013). Obviously, changes in functional networks are strongly dependent on the structural status of neuronal connections that link brain regions. Recent studies have reported a high similarity between functional network connectivity and underlying structural neuronal connections (Achard et al., 2008; Honey et al., 2009). However, it is largely unknown how functional networks reorganize in response to transient focal disturbances without structural damage. Characterizing the impact of transient localized functional brain disruptions on large-scale neural networks can provide important insights into a network’s resilience, which might lead to improved understanding of underlying causes of comorbidities in diseases with localized brain damage, such as focal epilepsy and ischemic stroke.

Global neural networks are known to have a hierarchical structure with the existence of a number of well-connected and highly central hub regions (Hagmann et al., 2008; Bullmore and Sporns, 2012) that form a so-called “rich-club,” which reflects a network’s

Keywords: rat brain, resting-state fMRI, transient brain impairment, neural network, rich-club effect
capacity to segregate and integrate multisensory information in a united manner (Van Den Heuvel and Sporns, 2011; Harriger et al., 2012). These well-connected regions play a key role in global signal processing between different parts of the network, and functional disturbance of hub regions can have critical impact on overall network functioning. In this study we aimed to map the impact of transient functional brain disruption on global neural networks in the absence of structural damage in a newly developed reproducible rat model. To that aim, we characterized the brain’s functional network organization with resting-state fMRI (rs-fMRI) after unilateral pentobarbital-induced brain “silencing.” We specifically focused on effects on well-connected hub regions using a relatively new framework that provides quantitative measures of the organization of highly connected rich-clubs (Van Den Heuvel and Sporns, 2011; Harriger et al., 2012; Collin et al., 2013). So far it has remained unknown whether rodent brain networks contain rich-club connections and if so, how these connections should be interpreted. Our hypothesis was that (i) rat brains have functional rich-club connections, and (ii) that transient hemispheric functional disruption affects the rich-club organization in both the disrupted and unaffected network regions.

METHODS

RAT MODEL

We developed a rat model of transient functional hemispheric disruption using unilateral focal anesthesia, similar to the intracarotid sodium amobarbital procedure for the “Wada test” in humans (Wada and Rasmussen, 2007). This procedure was chosen to temporally disrupt intra- and interhemispheric functional connectivity (FC) of the ipsilateral hemisphere. The protocol was approved by the Ethical Committee on Animal Experiments of the University Medical Center Utrecht and Utrecht University. Experiments were carried out in accordance with the guidelines of the European Communities Council Directive.

Three healthy adult Sprague-Dawley rats (Charles River Laboratories International Inc., MA, USA) were mechanically ventilated with 2% isoflurane in air: O2(7:1; rate 52–59 min⁻¹). To catheterize the right carotid artery, the right common, and external carotid arteries were exposed by a ventral incision in the neck. A small catheter was advanced into the external carotid artery in the direction of the common carotid artery until the blunt catheter tip reached the bifurcation of the internal and external carotid arteries. Blood flow from the common into the internal carotid artery was ensured to be maintained (that is, the catheter tip remained in the ligated external common artery). The catheter was kept in place with a 5-0 suture and instant glue. The incision was closed and animals received 5 mL glucose-saline solution to compensate for loss of water and minerals.

Directly after surgery, rats were placed in an MR-compatible stereotactic holder and immobilized with earplugs and a tooth-holder. Blood oxygen saturation and heart rate were continuously monitored with a pulse-oximeter with the probe positioned on a hindpaw. In addition, expired CO₂ was continuously monitored with a capnograph, and body temperature was maintained at 37.0 ± 0.5°C using a feedback-controlled heating pad. End-tidal CO₂ levels were kept within the normal range, equivalent to arterial pCO₂ levels between 35 and 45 mmHg (calculated from previous calibration measurements), by adjusting ventilation volume and rate when necessary.

MRI

MRI experiments were done at 4.7 T with a 125 mm internal diameter gradient-coil insert. A Helmholtz volume coil and an inductively coupled surface coil were used for signal excitation and detection, respectively.

Repetitive blood oxygenation level-dependent rs-fMRI scans were acquired using a gradient echo single shot echo-planar imaging (EPI) sequence [repetition time (TR)/echo time (TE) = 500/19 ms; 7 1.5-mm coronal slices; field-of-view = 32 × 32 mm; in-plane voxel resolution, 0.5 × 0.5 mm; pulse angle = 35°; temporal scan resolution = 500 ms; total scan time = 15 min]. Exactly 10 min prior to rs-fMRI acquisition, end-tidal isoflurane concentration was reduced to 1.0%, which was maintained during rs-fMRI.

After 7.5 min rs-fMRI acquisition, 175 µL of 2 mg/kg pentobarbital was slowly injected intracarotidly. This concentration based on an electroencephalography (EEG) titration/dose-response experiment, which showed that suppression of EEG activity occurred at this threshold level at 1% isoflurane anesthesia (data not shown). EEG activity returned to baseline within 15 min.

In one animal, perfusion MRI was conducted (TR/TE = 100/25 ms) following rs-fMRI, using gradient echo EPI (TR/TE = 100/25 ms; 11.5-mm coronal slice; field-of-view = 32 × 32 mm; in-plane voxel resolution, 0.5 × 0.5 mm; pulse angle = 90°) in combination with an bolus injection of 50 µL of 1.0 mmol/mL gadobutrol through the intracarotid catheter. Serial images were inspected to verify unilateral bolus arrival.

EEG

To validate functional MRI measurements, EEG was recorded immediately following rs-fMRI acquisitions, at 1.0% isoflurane anesthesia outside the MR scanner. To that aim a small medial incision was made in the skin covering the skull and the pericranium. A hole in the skull above the left and right primary somatosensory cortices was made with a 300 µm micro drill. Next, epidural EEG was recorded from the bilateral somatosensory cortices with a reference electrode above the cerebellum. EEG activity was recorded for 15 min in two animals. After 7.5 min of EEG measurements, 175 µL of 2 mg/kg pentobarbital was slowly injected intracarotidly. The time between this injection and the first injection in the MR scanner was >1 h. EEG measurements were conducted using a homebuilt multichannel amplifier, and band-pass filtered between 0.1 Hz and 250 Hz, using a National Instruments™ NI USB-6211 DAQ with a sampling rate of 1000 Hz per channel.

IMAGE PROCESSING AND ANALYSIS

Rs-fMRI images were motion-corrected, spatially smoothed, band-pass filtered between 0.01 and 0.1 Hz, and nonlinearly matched with the Paxinos and Watson rat brain atlas (Paxinos and Watson, 2006) for FC analysis. The first (prebarbital) and last (postbarbital) 500 images were used to extract time series for 32
regions-of-interest (ROIs) in each hemisphere, and to calculate pair-wise FCs between all ROIs from the correlation coefficient \( r \). ROIs were carefully selected to basically include all bilateral neocortical and subcortical gray matter atlas regions (see Figure 2). The cerebellum was excluded from analysis. All ROIs were large enough to contain multiple voxels at the applied fMRI resolution, and were functionally connected to at least one other region.

### ESTIMATION OF WEIGHTED RICH-CLUB EFFECT

Pre- and postbarbital FCs were averaged across animals to construct two \( 64 \times 64 \) node networks (each node is a cortical or subcortical ROI in one of the hemispheres). From these networks ipsilateral and contralateral \( 32 \times 32 \) node sub-networks were extracted. Negative FCs were excluded to allow calculations according to the weighted rich-club methodology (Opsahl et al., 2008).

The weighted rich-club effect characterizes the tendency of network hubs to engage in stronger or weaker FCs among them than expected by chance. The calculation involves multiple steps. First, all network nodes are ranked in terms of prominence. We used node strength \( s \) as a measure of prominence. Strength is defined as the sum of FCs attached to the connections originating from a network node (Barrat et al., 2004). For increasing values of \( s \), we selected the group (the club) of all nodes whose strengths were larger than \( s \). In this step we obtain a series of increasingly selective clubs. For each of these clubs we counted the number \( E_{s}, \) of connections connecting the club members, and measured the sum \( W_{s} \) of the FCs attached to these connections.

In the next step we determined the ratio \( \phi_w(s) \) between \( W_{s} \), and the sum of the FCs attached to the \( E_{s} \), strongest connections within the network. Mathematically:

\[
\phi_w(s) = \frac{W_{s}}{\sum_{l=1}^{E_{s}} w_{l}^{rank}}
\]

where \( w_{l}^{rank} \geq w_{l+1}^{rank} \) with \( l = 1, 2, \ldots, E \) the ranked FCs on the connections of the network. \( E \) is the total number of connections. With this formulation we determined the fractions of FCs shared by rich nodes compared with the total amount they could share if they were connected through the strongest connections of the network.

A high value of \( \phi_w(s) \) is not sufficient to account for the weighted rich-club effect as higher-strength nodes in a network have a higher probability of sharing connections with each other just by chance. In other words, networks where connections are randomly established could have a non-zero value of \( \phi_w(s) \). To control for this effect we normalized \( \phi_w(s) \) relative to the average weighted rich-club effect, \( \phi_{\text{random}}(s) \), found in 1000 random networks generated using a reshuffle procedure. This procedure reshuffles the FCs globally in the networks, while keeping the topology intact.

The weighted rich-club effect \( \rho_w(s) \) at strength \( s \) is the ratio:

\[
\rho_w(s) = \frac{\phi_w(s)}{\phi_{\text{random}}(s)}
\]

When \( \rho_w(s) \) is > 1.0 the functional network has a positive weighted rich-club effect, with nodes in the rich-club concentrating most of their connections toward other club members relative to random networks.

### EEG ANALYSIS

EEG signals were analyzed in consecutive epochs of 30 s. For each epoch the correlation coefficients between the temporal fluctuations of the EEG signals from left and right somatosensory cortices were calculated for the delta (1–5 Hz), theta (5–8 Hz), alpha (8–15 Hz), beta (15–30 Hz), gamma (30–250 Hz), and broad (1–255 Hz) frequency bands.

### RESULTS

Perfusion MRI data demonstrated that the catheterized carotid artery predominantly supplied the ipsilateral hemisphere allowing for selective hemispheric brain silencing with pentobarbital (Figure 1). Perfusion was strongest in the cortical areas, whereas levels in subcortical regions were lower. We also detected some contrast agent arrival in the anterior contralateral hemisphere, which can be explained by delivery via the circle of Willis.

### NETWORK PROMINENCE

The prominence of all network nodes, measured as the strength, is plotted in Figure 2 for the prebarbital (baseline) and postbarbital whole-brain functional networks, as calculated from the rs-fMRI experiments. The ten most prominent prebarbital ROIs included mostly somatosensory regions, namely, the ipsi- and contralateral dysgranular zone of the primary somatosensory cortex; the contralateral forelimb region, upper lip region and barrel field of the primary somatosensory cortex; and the ipsilateral jaw region of the primary somatosensory cortex. The agranular and granular insular cortex and the facial nuclei were also prominent network nodes before pentobarbital injection.

Following unilateral pentobarbital injection, a subgroup of these ROIs lost prominence (Figure 2). This was most pronounced for the contralateral barrel field of the primary somatosensory cortex and the agranular insular cortex. In contrast, the prominence of the contralateral primary motor cortex increased toward the highest value in the postbarbital network. The ROIs presenting with the largest changes in prominence were not confined to a single anatomical region.

### FUNCTIONAL CONNECTIVITY

The average baseline functional network, i.e., before pentobarbital injection, displayed strong connectivity within and between hemispheres. After pentobarbital injection the ipsilateral hemisphere revealed increased intrahemispheric connections with concomitant decrease of interhemispheric FC (Figure 3A). Increased intrahemispheric FC was found in both hemispheres. Several ipsilateral regions retained strong network strengths with other regions, including the facial nucleus, premotor cortex, cingulate cortex area 1, and the granular insular cortex.

### WEIGHTED RICH-CLUB EFFECT

The bilateral functional network was characterized by a strong weighted rich-club effect, with \( \rho^W(s) > 1.0 \) over a range of prominence thresholds (Figure 3B). This rich-club effect was not affected by ipsilateral FC disruption. In contrast to the whole brain network, the \( \rho^W(s) \) values were significantly decreased in the ipsilateral sub-network following unilateral pentobarbital
FIGURE 1 | Perfusion MRI frames, demonstrating clear unilateral delivery of the contrast agent in the hemisphere ipsilateral to the side of cannulation. The frames were acquired at 100 ms intervals from the start of bolus administration until 1.5 s later (the temporal frame sequence runs from top left to bottom right). The frames are in coronal plane at 1 mm from bregma. Presence of contrast agent is displayed using a color scale: dark for low contrast concentrations and bright for high contrast concentration (in arbitrary units). The green rectangles mark the ipsi- and contralateral hemispheres.

injection (Figure 3C). This pattern of decreased $\rho^w(s)$ values was to a lesser extent also seen in the contralateral sub-network.

DATA VARIATION
The variation between individual functional networks is shown in Figure 4. The standard deviation of the FC values slightly increased post pentobarbital injection as shown from the density plots. The standard deviation range was 0–0.3 prior to injection and 0–0.4 post injection.

ELECTROENCEPHALOGRAPHY
The EEG measurements, acquired in the same animals directly after imaging, supported our MRI-based observation of decreased interhemispheric resting-state FC. Interhemispheric EEG synchronization decreased after unilateral pentobarbital injection in the alpha, theta, and broadband frequencies (Figure 5). The synchronization in the other frequency bands increased temporarily (gamma) or remained unchanged (beta).

DISCUSSION
Our pilot study shows that transient functional global network disruptions in response to unilateral and reversible functional brain disturbance can be effectively studied using a straightforward rodent model in combination with rs-fMRI. Our rat data support the concept that densely connected rich-club regions play a central role in global brain communication, as has been previously shown in humans. In addition, we found that global network hub configurations can be significantly affected by temporary functional disruption of a hemisphere.

A rodent model that allows selective and reversible functional hemispheric silencing has not been described before. Our results show that this model can be useful to assess effects of (transient) brain impairment on functional networks. The concept of this model was based on recent FC and network data presented by Douw et al. (2009, 2010). They showed that FC within and between hemispheres changed in persons with epilepsy during an intra-arterial amobarbital injection—the WADA test—as measured with EEG (Douw et al., 2009). They concluded that the effects of unilateral anesthesia on FC in the contralateral hemisphere should not be ignored. Direct comparison of low-frequency rs-fMRI connectivities with the different EEG frequency bands is difficult, and more data are required to elucidate differences and similarities between rs-fMRI and EEG patterns. Nevertheless, the observed decrease in interhemispheric EEG FC in the delta and theta bands by Douw and colleagues corresponds with our interhemispheric EEG results (Figure 5). Over the ipsilateral hemisphere the delta and theta band synchronization increased after amobarbital injection in humans, which also agrees with our findings of increased ipsilateral intrahemispheric FC in rat brain after pentobarbital injection. Apparently, unilateral functional brain disruption by amobarbital injections is confined to subverted functional outcome (that is, speech and language comprehension in humans) rather than absolute inhibition of low-frequency FC. This idea is further supported by recent FC measurements at an anesthetic depth characterized by unresponsiveness, where reductions in connectivity were shown to be partial and not complete (Hudetz, 2012).
Douv et al. also assessed EEG data acquired during WADA-test using whole-brain functional network analysis (Douv et al., 2010). Their results revealed an increased random network organization after unilateral functional disruption with amobarbital in people with epilepsy (Douv et al., 2010). However, the network topologies within the injected and non-injected hemispheres were not studied separately in this group of patients. In our study we found no difference in rich-club characteristics between pre- and postbarbital whole-brain networks. Differences in weighted rich-club effects were only found in sub-network analyses of individual hemispheres (Figure 3). The asymmetric presentation of prebarbital hub regions (that is, mostly contralateral sensorimotor areas) is in line with studies in healthy rats that reported right-sided population bias and functional lateralization (Glick and Ross, 1981; Alonso et al., 1991).

A positive weighted rich-club effect characterizes the tendency of network hubs to engage in stronger functional connectivities among them than expected by chance. We found significantly
FIGURE 3 | Pre- and postbarbital (A) whole-brain functional connectivity (FC) networks, displayed as matrices. Each row and column stands for a region-of-interest in the hemisphere ipsi- or contralateral to the side of cannulation. Matrix elements are colored according to FCs values, measured as the correlation coefficient \( r \) between regions-of-interest. Red indicates no FC; yellow indicates high FC. Interhemispheric FCs are delineated with blue squares; intrahemispheric FCs with green squares. Weighted rich-club effects (y-axis) are plotted against increasing levels of strength (x-axis) for the entire brain network (B), and the contra- and ipsilateral (C) hemispheric networks. The maximum strength threshold depends on the size of the network. The strength thresholds were increased until networks started to disintegrate. This explains the different scaling of the x-axis in the hemispheric networks as compared to the entire brain network. Significant differences in weighted rich-club effects between prebarbital (blue line) and postbarbital (red line) networks are indicated as * [permutation analysis; \( p < 0.01 \) considered significant].

decreased \( \rho (s) \) values in the ipsi- and contralateral sub-networks after unilateral pentobarbital injection. Similar positive rich-club organizations have been described at a neuronal level in the nematode worm Caenorhabditis elegans (Towlson et al., 2013) and at a larger scale in human neocortical networks (Van Den Heuvel and Sporns, 2011). Our findings in rodent brain networks, consolidate the hypothesis that connections according a rich-club layout may be a general and scale-invariant principle of brain network organization (Towlson et al., 2013). Nonetheless, further validation of this hypothesis is essential, given the small sample size used in this study.

The rich-club connections in the structural human brain network are considered to be the backbone for global brain communication (Van Den Heuvel et al., 2012). The majority of these connections are long-distance neural pathways, such that the rich-club phenomenon is costly in terms of consumed energy and space. However, the trade-off for higher cost is higher performance in terms of network integration; up to 70% of all network’s shortest paths travel through rich-club connections.

Whether structural network principles also hold for functional rich-club networks remains to be determined, but we expect the functional rich-club organization to follow the underlying structural organization to a large extent. On the other hand, functional rich-club connections may be much more flexible in acutely responding to environmental changes, such as induced by anesthesia. This is, for instance, underscored by the differences in our whole-brain and sub-network data (Figures 3B,C). Unilateral brain silencing did not affect the rich-club effect at a whole-brain level; only the club members changed dramatically (Figure 2). On the other hand, the hemispheric sub-networks did reveal change in rich-club connectivity after hemispheric silencing. Apparently the level of network definition is critical for detecting changes in functional rich-club organization after functional brain lesioning. A better understanding of this
phenomenon may require the acquisition of structural diffusion tensor imaging data alongside rs-fMRI in future studies. This will allow comparison of functional networks and their rich-club organization with data from the intact structural networks.

Limitations in this proof-of-principle methods study should be acknowledged. First, experiments were conducted at 1.0% isoflurane anesthesia. We do not know how this baseline anesthesia may have interacted with the pentobarbital-induced brain silencing. We expect this effect to be relatively small as cerebral glucose utilization, cerebral blood flow coupling (Maekawa et al., 1986; Lenz et al., 1998) and FC, including the long distance connections (Liang et al., 2012), are largely preserved at 1.0% isoflurane anesthesia (Wang et al., 2011). We also know that the neurovascular coupling in rat brain is similar across distinct types of anesthetics including pentobarbital and isoflurane (Franceschini et al., 2010), however further validation studies, including robust dose-response experiments, are needed. Rs-fMRI acquisition may be ideally executed in awake rats, however, this is complicated due to motion artifacts related to post-surgical pain, stress and discomfort.

Second, anesthesia-based functional brain disruption by unilateral injection through the carotid artery may not be exclusively unilateral due to presence of the circle of Willis and recirculation effects. Nonetheless, the small injection volume used in our study limited this effect, which was confirmed by the strongly lateralized results for hemispheric FCs.

Third, the likelihood that a statistically significant result reflects a true effect is reduced in studies with a small sample size (Button et al., 2013). The large changes in functional network connectivity and organization that we observed in this rat model alleviate this problem. Nonetheless, firm conclusions on directionality of network changes and size of effect require additional studies with larger groups of animals. This is emphasized by the variation that we found between FC correlation coefficients obtained from the individual networks (Figure 4). This variation is however comparable to the variation found between individual interhemispheric FC in rats recovering from a unilateral stroke, as studied with rs-fMRI with a similar experimental setup and design (Van Meer et al., 2012).

In conclusion, our pilot study suggests that changes in rich-club organization following functional network disruption might affect the capacity to efficiently segregate and integrate multisensory information. These changes are unlikely to depend on the presence of structural damage. We speculate that functional rich-club alterations may reflect an underlying mechanism of neurological impairments in patients with no or limited structural brain damage, such as after transient ischemic attack.
Collin, G., Sporns, O., Mandl, R. C., and Van Den Heuvel, M. P. (2013). Structural and functional aspects relating to cost and benefit of rich club organization in the human cerebral cortex. *Cereb. Cortex*. doi: 10.1093/cercor/bht064. [Epub ahead of print].

Douv, L., Baayen, J. C., Klein, M., Velis, D., Alpherts, W. C., Bot, J., et al. (2009). Functional connectivity in the brain before and during intraarterial amobarbital injection (Wada test). *Neuroimage* 46, 584–588. doi: 10.1016/j.neuroimage.2009.02.034

Douv, L., Van Dellen, E., Baayen, J. C., Klein, M., Velis, D. N., Alpherts, W. C., et al. (2010). The lesioned brain: still a small-world? *Front. Hum. Neurosci.* 4:174. doi: 10.3389/fnhum.2010.00174

Fornito, A., Zalesky, A., and Breakspear, M. (2013). Graph analysis of the human connectome: promise, progress, and pitfalls. *Neuroimage* 80C, 426–444. doi: 10.1016/j.neuroimage.2013.04.087

Franceschini, M. A., Radhakrishnan, H., Thakur, K., Wu, W., Ruvinskaya, S., Carp, S., et al. (2010). The effect of different anesthetics on neurovascular coupling. *Neuroimage* 51, 1367–1377. doi: 10.1016/j.neuroimage.2010.03.060

Glick, S. D., and Ross, D. A. (1981). Right-sided population bias and lateralization of activity in normal rats. *Brain Res.* 205, 222–225. doi: 10.1016/0006-8993(81)90737-X

Grekle, C., and Fink, G. R. (2011). Reorganization of cerebral networks after stroke: new insights from neuroimaging with connectivity approaches. *Brain* 134, 1264–1276. doi: 10.1093/brain/awt033

Hagmann, P., Cammoun, L., Gigandet, X., Meuli, R., Honey, C. J., Wedeen, V. J., et al. (2008). Mapping the structural core of human cerebral cortex. *PLoS Biol.* 6:e159. doi: 10.1371/journal.pbio.0060159

Harriger, L., Van Den Heuvel, M. P., and Sporns, O. (2012). Rich club organization of macaque cerebral cortex and its role in network communication. *PLoS ONE* 7:e46497. doi: 10.1371/journal.pone.0046497

Honey, C. J., Sporns, O., Cammoun, L., Gigandet, X., Thiran, J. P., Meuli, R., et al. (2009). Predicting human resting-state functional connectivity from structural connectivity. *Proc. Natl. Acad. Sci. U.S.A.* 106, 2035–2040. doi: 10.1073/pnas.0811168106

Hudetz, A. G. (2012). General anesthesia and human brain connectivity. *Brain Connect.* 2, 289–302. doi: 10.1089.brain.2012.0107

Langhorne, P., Coupar, F., and Pollock, A. (2009). Motor recovery after stroke: a systematic review. *Lancet Neurol.* 8, 741–754. doi: 10.1016/S1474-4222(09)70150-4

Lenz, C., Rebel, A., Van Ackern, K., Kuschinsky, W., and Waschke, K. F. (1998). Local cerebral blood flow, local cerebral glucose utilization, and flow-metabolism coupling during sevoflurane versus isoflurane anesthesia in rats. *Anesthesiology* 89, 1480–1488. doi: 10.1097/00000542-199812000-00026

Liang, Z., King, J., and Zhang, N. (2012). Intrinsic organization of the anesthetized brain. *J. Neurosci.* 32, 10183–10191. doi: 10.1523/JNEUROSCI.1020-12.2012

Mohankumar, T., Tommasino, C., Shapiro, H. M., Keffer-Goodman, J. L., and Kohlenberger, R. W. (1986). Local cerebral blood flow and glucose utilization during intravenous anesthesia in the rat. *Anesthesiology* 65, 144–151. doi: 10.1097/00000542-198608000-00003

Motamedi, G., and Sporns, O. (2010). Complex network measures of brain connectivity: uses and interpretations. *Epilepsia* 52, S25–S38. doi: 10.1111/j.1528-1167.2009.02225.x

Opsahl, T., Colizza, V., Panzarasa, P., and Ramasco, J. J. (2008). Prominence and control: the weighted rich-club effect. *Phys. Rev. Lett.* 101:168702. doi: 10.1103/PhysRevLett.101.168702

Paxinos, G., and Watson, C. (2006). *The Rat Brain in Stereotaxic Coordinates.* London: Elsevier.

Rubinov, M., and Sporns, O. (2010). Complex network measures of brain connectivity: uses and interpretations. *Neuroimage* 52, 1059–1069. doi: 10.1016/j.neuroimage.2009.10.003

Stinear, C. (2010). Prediction of recovery of motor function after stroke. *Lancet Neurol.* 9, 1228–1232. doi: 10.1016/S1474-4222(10)70247-7

Tatemichi, T. K., Desmond, D. W., Stern, Y., Paik, M., Sano, M., and Bagiella, E. (2009). Cognitive impairment after stroke: frequency, patterns, and relationship to functional abilities. *J. Neurol. Neurosurg. Psychiatry* 80, 202–207. doi: 10.1136/jepp.57.2.202

Towndrow, E. K., Vertes, P. E., Ahnert, E. S., Schafer, W. R., and Bullmore, E. T. (2013). The rich club of the C. elegans neuronal connectome. *J. Neurosci.* 33, 6380–6387. doi: 10.1523/JNEUROSCI.3784-12.2013

Van Den Heuvel, M. P., Kahn, R. S., Goni, J., and Sporns, O. (2012). High-cost, high-capacity backbone for global brain communication. *Proc. Natl. Acad. Sci. U.S.A.* 109, 11372–11377. doi: 10.1073/pnas.1203593109

**ACKNOWLEDGMENTS**

Willem M. Otte is supported by the Dutch National Epilepsy Fund (NEF 12-05).

**REFERENCES**

Achard, S., Bassett, D. S., Meyer-Lindenberg, A., and Bullmore, E. (2008). Fractal connectivity of long-memory networks. *Phys. Rev. E Stat. Nonlin. Soft Matter Phys.* 77, 036104. doi: 10.1103/PhysRevE.77.036104

Alonso, J., Castellano, M. A., and Rodriguez, M. (1991). Behavioral lateralization in rat prenatal stress effects on sex differences. *Brain Res.* 539, 45–50. doi: 10.1016/0006-8993(91)90684-N

Barrat, A., Barthelemy, M., Pastor-Satorras, R., and Vespignani, A. (2004). The architecture of complex weighted networks. *Proc. Natl. Acad. Sci. U.S.A.* 101, 3747–3752. doi: 10.1073/pnas.0400087101

Bullmore, E., and Sporns, O. (2012). The economy of brain network organization. *Nat. Rev. Neurosci.* 13, 336–349. doi: 10.1038/nrn3214

Button, K. S., Ioannidis, J. P., Mokrysz, C., Nosek, B. A., Flint, J., Robinson, E. S., et al. (2013). Power failure: why small sample size undermines the reliability of neuroscience. *Nat. Rev. Neurosci.* 14, 365–376. doi: 10.1038/nrn3475

Carter, A. R., Aastafiev, S. V., Lang, C. E., Connor, L. T., Rengachary, J., Strube, M. J., et al. (2010). Resting interhemispheric functional magnetic resonance imaging connectivity predicts performance after stroke. *Ann. Neurol.* 67, 365–375. doi: 10.1002/ana.21905

FIGURE 5 | The interhemispheric somatosensory functional EEG connectivities (measured as correlation coefficients (r)) for consecutive epochs, each 30 s, plotted over time (15 min) for the distinct frequency bands (A). The corresponding moment of pentobarbital injection and periods of resting-state fMRI used to construct functional networks are shown below (B). Note that functional imaging was acquired prior to EEG, but was identical in approach and timing of pentobarbital injection.
Van Den Heuvel, M. P., and Sporns, O. (2011). Rich-club organization of the human connectome. *J. Neurosci.* 31, 15775–15786. doi: 10.1523/JNEUROSCI.3539-11.2011

Van Meer, M. P., Otte, W. M., Van Der Marel, K., Nijboer, C. H., Kavelaars, A., Van Der Sprenkel, J. W., et al. (2012). Extent of bilateral neuronal network reorganization and functional recovery in relation to stroke severity. *J. Neurosci.* 32, 4495–4507. doi: 10.1523/JNEUROSCI.3662-11.2012

Wada, J., and Rasmussen, T. (2007). Intracarotid injection of sodium amytal for the lateralization of cerebral speech dominance. 1960. *J. Neurosurg.* 106, 1117–1133. doi: 10.3171/jns.2007.106.6.1117

Wang, K., Van Meer, M. P., Van Der Marel, K., Van Der Toorn, A., Xu, L., Liu, Y., et al. (2011). Temporal scaling properties and spatial synchronization of spontaneous blood oxygenation level-dependent (BOLD) signal fluctuations in rat sensorimotor network at different levels of isoflurane anesthesia. *NMR Biomed.* 24, 61–67. doi: 10.1002/nbm.1556

**Conflict of Interest Statement:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Received: 13 September 2013; accepted: 06 March 2014; published online: 21 March 2014.

Citation: Otte WM, van der Marel K, Braun KP and Dijkhuizen RM (2014) Effects of transient unilateral functional brain disruption on global neural network status in rats: a methods paper. *Front. Syst. Neurosci.* 8:40. doi: 10.3389/fnsys.2014.00040

This article was submitted to the journal Frontiers in Systems Neuroscience. Copyright © 2014 Otte, van der Marel, Braun and Dijkhuizen. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) or licensor are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.