Parietofrontal network upregulation after motor stroke

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ABSTRACT

Objective: Motor recovery after stroke shows a high inter-subject variability. The brain's potential to form new connections determines individual levels of recovery of motor function. Most of our daily activities require visuomotor integration, which engages parietal areas. Compared to the frontal motor system, less is known about the parietal motor system's reconfiguration related to stroke recovery. Here, we tested if functional connectivity among parietal and frontal motor areas undergoes plastic changes after stroke and assessed the behavioral relevance for motor function after stroke.

Methods: We investigated stroke lesion-induced changes in functional connectivity by measuring high-density electroencephalography (EEG) and assessing task-related changes in coherence during a visually guided grip task with the paretic hand in 30 chronic stroke patients with variable motor deficits and 19 healthy control subjects. Quantitative changes in task-related coherence in sensorimotor rhythms were compared to the residual motor deficit.

Results: Parietofrontal coupling was significantly stronger in patients compared to controls. Whereas motor network coupling generally increased during the task in both groups, the task-related coherence between the parietal and primary motor cortex in the stroke lesioned hemisphere showed increased connectivity across a broad range of sensorimotor rhythms. Particularly the parietofrontal task-induced coupling pattern was significantly and positively related to residual impairment in the Nine-Hole Peg Test performance and grip force.

Interpretation: These results demonstrate that parietofrontal motor system integration during visually guided movements is stronger in the stroke-lesioned brain. The correlation with the residual motor deficit could either indicate an unspecific marker of motor network damage or it might indicate that upregulated parietofrontal connectivity has some impact on post-stroke motor function.

1. Introduction

The basis of spontaneous recovery from a motor stroke in humans is unclear. Given a window of heightened microstructural plasticity (Rossini et al., 2003; Ward, 2017) and evidence for dynamic reconfiguration of cortical areas engaged in motor activity after stroke (Grefkes and Fink, 2011), it is conceivable that interaction patterns between cortical areas reorganize after a lesion. Unravelling common patterns of neuropsychiatric processes that lead to regain of motor function is a major challenge of neurorehabilitative medicine and a prerequisite for a mechanistic approach of interventional therapies.

During physiologic motor activity, the primary and secondary motor areas engage in balanced facilitatory and inhibitory interactions (Bonstrup et al., 2016; Grefkes et al., 2008a). After a motor stroke, in the acute phase, the facilitatory coupling among the frontal motor areas, i.e. ventral premotor cortex (PMv), the supplementary motor area (SMA) and the primary motor cortex (M1) in the contralateral (ipsilesional) hemisphere appears to be disrupted; and to normalize along with recovery (Rehme et al., 2011b). Most of our daily activities require precise interaction between visual perception and the motor system. For higher order movements, that require visuomotor and sensorimotor integration (Vingerhoets, 2014) as well as for reaching and grasping (Bernier et al., 2017; Corbetta and Shulman, 2002; Grefkes et al., 2004; Konen et al., 2013), parietal motor areas are...
specifically engaged. However, their relative role in motor system re-
configuration after stroke is less well understood compared to the
frontal brain. Conceivably, parietofrontal pathways are of high im-
portance for post-stroke rehabilitation (Wu et al., 2014). Recently, it
was shown that lesion-induced network plasticity involves parieto-
frontal motor pathways connecting PMv with the anterior intraparietal
sulcus (aIPS) (Schulz et al., 2015). And in functional magnetic re-
sonance imaging (fMRI), it was demonstrated that reciprocal facili-
tatory connectivity between the aIPS and M1 in the ipsilesional
hemisphere is enhanced in well-recovered chronic stroke survivors
compared with healthy participants during a visually guided grip task
(Schulz et al., 2016). However, the functional relevance of parieto-
frontal network upregulation for motor function has not been char-
acterized yet. Whether the abnormally increased parietofrontal con-
nectivity in stroke survivors is systematically varying with the degree of
paresis, is a highly intriguing question not only for deeper pathophy-
siologic understanding, but also to substantiate the validity and im-
portance of this connection as a tentative target for non-invasive brain
stimulation protocols.

We recorded high-density EEG during a visually guided grip task
with the paretic hand in 30 chronic stroke patients (we use the term
chronic in agreement with previous de-
cit. We first set out to reproduce the previous finding of enhanced fa-
cilitatory coupling between the parietal and motor cortex in a larger
patient cohort and with a different method to detect brain connectivity:
Across perceptual, cognitive and motor systems, synchronization of
oscillations has been detected as a key mechanism of how transient
coalitions of neural populations at small- and large spatial scales
commit to a common task (communication through coherence concept)
(Bonnefond et al., 2017; Siegel et al., 2012). We hypothesized that
parietal and frontal motor areas would show higher coherence during
the task for patients than for control participants. Secondly, we hy-
pothesized that higher coherence parameters are found in patients with
a stronger residual motor deficit.

2. Participants and methods

2.1. Participants

30 patients (19 male, 1 left-handed, aged 65 ± 13 years, mean ±
std) were included three months (104 ± 17 days) after first-ever is-
chemic stroke causing a motor deficit involving hand function (five
subcortical, 25 cortical/cortico-subcortical). 14 patients had lesions
in the dominant hemisphere. Initial and residual motor impairment
was determined by means of grip force, the Nine-Hole Peg Test performance
(NHP), and the Fugl–Meyer score for the upper extremity (UEFM)
3–5 days after stroke and 3 months after stroke. For the grip force and
the NHP, behavioral scores were calculated as proportional values
(affected/unaffected hand), whereas in case of the NHP, prior to nor-
malization the score for each hand was expressed as pegs/s to give a
performance value. Individual motor recovery values were obtained by
calculating the difference between initial and residual behavioral
scores. A group similar in age and gender (n = 19, 10 males, one left-

handed, aged 64.8 ± 11.1 years) served as controls. The study design
was approved by the local ethical committee. All participants gave their
written informed consent according to the ethical declaration of
Helsinki.

2.2. Motor task

Participants underwent EEG during a simple motor task which re-
quired them to perform isometric visually guided whole hand grips
with the paretic hand using a grip-force device (Grip Force Bimanual,
Current Designs, Inc., Philadelphia, USA). The control participants were
randomly assigned to use the left or right hand in a distribution
matching the lesion-side of the patient group (nine right hand).
Participants were seated comfortably in an armchair with the right
and left arm relaxed positioned in their lap, each holding one of the bi-
manual grip-force devices. We compared two conditions of varying
target grip force, one keeping the force constant across the group
(constant output of 5 kg) and the other keeping the task effort constant
across the group (constant effort of 20% MVC). Each condition was
recorded with 20 repetitions of a 9 s constant grip hold phase. The begin-
ning of each grip as well as continuous feedback about the applied force
were provided visually by the appearance and vertical level of a hori-
zontal bar on a screen. The participants were instructed to lift the bar
into the target zone (paralleling the target force of either 5 kg (=con-
stant-output) or 20% of maximal force (= constant-effort)) and hold it
constant until it disappeared (after 9 s, Supplementary Fig. 1). Participants
were instructed to avoid eye movements and fixate the bar,
whose level was within a small visual angle of ± 5°, thus not requiring
large amplitude eye movements during the force build-up. During the
inter-trial interval of 12 ± 2 s, participants were instructed to fixate
a cross in the center of the screen and relax. To assess bilateral move-
ments, the force applied with the (unaffected) non-active hand was
continuously monitored throughout the hand grip as the patient held a
grip force device in both hands. If necessary, breaks were introduced
depending on the participant’s needs. The task was described in the
previous report on functional MRI (fMRI) derived effective coupling
(Schulz et al., 2016) and in (Bonstrup et al., 2015).

2.3. Data acquisition

2.3.1. Electroencephalography

The EEG was recorded from 63 cephalic active surface electrodes,
referenced to a nose-tip or Cz-electrode during recording (interim re-
placement of recording setup). One electrode was mounted below the
left eye for electrooculogram recording. Before and after each experi-
mental session, a resting state was recorded for 4–5 min with eyes fixed
on a cross in the center of the screen. See Supplementary material
section 1.1 for details on the recording setup.

2.4. Data analysis

2.4.1. EEG data preprocessing

The continuous EEG was offline down sampled to 125 Hz, detrended
and subjected to an independent component analysis (logistic infomax
ICA; (Makeig et al., 1996)) to remove eye-blink artifacts. The 20 trials
of each grip condition were segmented in epochs of 1 s duration cov-
ering the hold phase, starting 1 s after the beginning of each trial until the
end (20 × 8 s). The resting state condition was likewise segmented
into epochs of 1 s. Trials were then visually inspected to reject re-
mainig artifacts (number of 1 s long trials after artifact rejection
(mean ± std): task 119 ± 14, rest 235 ± 29). See Supplementary
material section 1.2 for details on the preprocessing.

2.4.2. Source reconstruction and spectral power and coherence analysis

We reconstructed source space activity and connectivity in the
parietofrontal motor network using spatial filtering. The network con-
sisted of five ipsilesional regions of interest (ROIs) contralateral to the
(affected) active hand, consisting of M1, PMv, SMA, and the aIPS and
caudal part of the intraparietal sulcus (cIPS). Coordinates were pre-
deefined as reported previously ((Schulz et al., 2016), Suppl. Table 2).
For each location, a linear constrained minimum variance (LCMV)
beamforming filter was computed based on an individual forward
model and a covariance matrix of sensor space time series. The choice of
a beamforming approach for spatial filtering was based on previous
publications in the field of motor stroke research using EEG or mag-
netoencephalography (MEG). Cross-spectra between each pair of sen-
sors were calculated at frequency bands of interest using the continuous
wavelet transformation with a width of 5 cycles. The cross-spectra were
2.4.3. Task-related spectral power and coherence analysis

For normalization of the underlying distribution, spectral power estimates were log-transformed and coherence estimates were subjected to a hyperbolic inverse tangent (tanh⁻¹) transformation. As a second step, to reduce inter-subject variability, the source spectral power and coherence estimates recorded during task execution were normalized with the spectral estimates during rest (Gerloff et al., 2006). Together, task-related spectral power (TR-Pow) and task-related coherence (TR-Coh) were derived using the following formula:

\[
TR-Pow = \log(\text{Pow}_{\text{activation}}) - \log(\text{Pow}_{\text{rest}})
\]

\[
TR-Coh = \tanh^{-1}(\text{Coh}_{\text{activation}}) - \tanh^{-1}(\text{Coh}_{\text{rest}})
\]

2.5. Statistics

The statistical analyses were done with R Version 3.2.5 (R Team, 2015) and MATLAB Version 2011a. To assess differences in TR-Pow and TR-Coh in the parietofrontal motor network between chronic stroke patients and the control group, we iteratively ran a linear mixed effects (LME) model (Bonstrup et al., 2015) using the lmer function provided in the lmerTest package (Kuznetsova et al., 2015) for TR-Pow at each region (five ROIs) or TR-Coh between all pairs of regions (10 connections) at each of the predefined frequency bands within the parietofrontal motor network (15 and 30 models, respectively). The models were designed to explain variance in TR-Pow and TR-Coh with the fixed effect GROUP (two level factor: patient and control), grip TASK (two level factor: constant grip force of 5 kg or 20% of MVC) and subject ID as a random intercept. Each factors' predictive power was assessed using analysis of variance and if insignificant, the factor was dropped. The same statistical framework was used to assess the relationship between TR-Coh and motor function in each group individually, by explaining variance in TR-Coh with the fixed effect error (affected/unaffected side) grip force (GRIFF), NHP or UEFM score, alone and in interaction with the individual motor recovery values. Each model's validity was assessed via diagnostic plots of the model's residuals to assess normal distribution and homoscedasticity. The whole brain source spectral power maps in Supplementary Fig. 3 were thresholded using one-sample t-tests at each frequency and source location \((p < 0.05, \text{corrected for multiple comparisons using False discovery rate (FDR) correction according to Benjamini and Hochberg for 8214 tests at an alpha-level of 0.05})\) (Benjamini and Hochberg, 1995).

3. Results

3.1. Clinical data

The patient group \((n = 30)\) mostly consisted of well-recovered (UEFM > 60, \(n = 24)\) chronic stroke patients (Di Pino et al., 2014; Rehme et al., 2012), with a few moderate (UEFMA > 40, \(n = 3)\) to poorly-recovered (UEFMA ≤ 40, \(n = 3)\) patients. Of the well-recovered patients, five initially had a severe motor impairment (UEFMA ≤ 40), while another five had a moderate motor impairment (UEFMA > 40 & ≤ 60). The clinical data are given in Table 1, and a stroke lesion map is plotted in Fig. 1. Four out of the 30 patients were unable to fully reach the target force of 5 kg due to residual paresis and one patient was unable to generate any force at all. In these cases, patients were instructed to build up as much force as possible to lift the bar. During the motor task, four of the patients showed co-contraction of the unaffected hand in the constant-output task, and two patients in the constant-effort task, at a level within the detection sensitivity of the grip device (0.026–10 kg). Supplementary Fig. 2 depicts the time course of exerted grip force in the active and contralateral hand during each condition for each patient.

3.2. Task-related spectral power

In a whole brain spectral power analysis, there were significant TR-Pow decreases in the lower and upper alpha as well as beta bands over bilateral primary somatosensory cortices, premotor cortices and SMA. Supplementary Fig. 3 depicts whole brain group averaged spectral power maps rendered on the cortical surface for each group and frequency range. In the alpha frequency range, the TR-Pow decrease also involved the parietal cortices, as in the blood-oxygen-level dependent (BOLD)-activation maps reported in our previous study (Schulz et al., 2016). At the ROI within the parietofrontal motor network, both groups showed significant TR-Pow decreases at parietal and frontal motor areas. Regarding differences in TR-Pow reductions between patients and control group, there were no significant regions in the whole-brain topographical maps (two-sample t-test \(p < 0.05, \text{FDR-corrected for 8214 comparisons})\). Looking specifically at the parietofrontal ROI, a statistical group difference in the beta band was only found at the aIPS and cIPS (LME, factor GROUP \(p = 0.03\) at both ROI), whereas in both alpha bands, no statistical differences were found (Supplementary Table 1). Of note, this absence of systematic differences in spectral power between the two groups renders it unlikely that the observed differences in connectivity (see below) resulted from different signal-to-noise ratios between groups (Siegel et al., 2012). We found no significant difference between the TR-Pow decreases in both force levels (constant output of 5 kg and constant effort of 20% MVC across the group, factor TASK, (Supplementary Table 2)). This reproduces a previous finding in a smaller but overlapping patient group (Bonstrup et al., 2015).

3.3. Parietofrontal functional connectivity

In general, functional coupling within the parietofrontal motor network increased during the grip task. Both groups showed significant parietofrontal coupling predominantly in the alpha bands (aIPS-M1, cIPS-M1, cIPS-PMv, aIPS/cIPS-SMA) as well as frontomesiocentral coupling in the alpha bands (PMv-SMA). In both groups, less coupling was evident in the beta band. The most consistent finding over all frequency bands and strongest numerical difference in coupling between stroke patients and control participants was the connection between aIPS and M1. In stroke patients, we detected an increase in coupling between the aIPS and M1 which was significantly larger than in the control group and generalized over frequency bands (lower alpha: TR-Coh difference Stroke-Controls 0.2, \(p = 0.003\), upper alpha: 0.2, \(p = 0.008\), beta: 0.19, \(p = 0.010\), LME factor GROUP). Since coherence quantifies synchronization based on phase and amplitude and we detected large spectral power reductions in both groups, we confirmed the result by computing the phase locking value as a synchronization measure (see Supplementary material 1.3). We likewise found that the coupling between aIPS and M1 was exclusively (within the parietofrontal network) and consistently (over all frequency bands) increased and stronger in the patient group (Supplementary Table 5). Fig. 2 topographically illustrates significant coupling changes of the network for each group and the statistical comparison between groups for the lower alpha band (8–10 Hz, see Supplementary material Fig. 4 for topographical plots of coupling in the upper alpha and beta rhythm). Although obtained with a completely different method and based on a very different concept of neuronal coupling, these findings replicate previously reported coupling estimates derived from a DCM.
| ID | Sex | Age  | Handedness | Stroke | Stroke location | Initial NIHSS | Initial grip force | Initial NHP | Initial UEFM | TAS (days) | Residual grip force | Residual NHP | Residual UEFM |
|----|-----|------|-------------|--------|----------------|--------------|-------------------|--------------|--------------|------------|---------------------|--------------|--------------|
| 1  | M   | 62   | Right       | Left   | CI, CR         | 3            | 0.2               | 0.2          | 37           | 95         | 0.7                 | 1.0          | 62           |
| 2  | M   | 68   | Right       | Right  | Left           | 65           | 0.5               | 0.5          | 102          | 109        | 0.9                 | 1.0          | 62           |
| 3  | F   | 63   | Right       | Right  | CR, CI, INS    | 3            | 0.42              | 0.4           | 62           | 90         | 0.71                | 1.0          | 65           |
| 4  | M   | 70   | Right       | Right  | CR              | 1            | 0.49              | 0.42          | 63           | 104        | 1.0                 | 1.1          | 68           |
| 5  | M   | 70   | Right       | Right  | CR, CI, INS    | 3            | 0.49              | 0.42          | 63           | 104        | 1.0                 | 1.1          | 68           |
| 6  | M   | 58   | Right       | Right  | CR, CI, INS    | 7            | 0.53              | 0.53          | 89           | 89         | 0.78                | 0.85         | 30           |
| 7  | F   | 49   | Right       | Left   | CI, CR, BG, INS| 10           | 0               | 0.2           | 4            | 168        | 0.12                | 0          | 66           |
| 8  | M   | 68   | Left        | Left   | CI, CR, INS    | 5           | 0.8               | 0.8           | 65           | 124        | 0.8                 | 0.7          | 65           |
| 9  | F   | 73   | Right       | Right  | CR, CI, BG     | 3            | 0.8               | 0.8           | 62           | 124        | 0.8                 | 0.7          | 65           |
| 10 | M   | 58   | Right       | Left   | PON            | 7            | 0               | 0.2           | 7            | 146        | 0.6                 | 0.8          | 66           |
| 11 | F   | 53   | Right       | Left   | CR              | 4            | 0.9               | 0.9           | 65           | 90         | 1.5                 | 0.7          | 66           |
| 12 | M   | 68   | Left        | Left   | CI, CR, INS    | 3            | 0.68              | 0.68          | 41           | 93         | 1.19                | 0.7          | 66           |
| 13 | M   | 48   | Right       | Right  | CR              | 3            | 0.53              | 0.53          | 7            | 101        | 1.06                | 1.0          | 66           |
| 14 | F   | 70   | Right       | Left   | CR              | 3            | 0.4               | 0.4           | 63           | 96         | 0.7                 | 0.6          | 66           |
| 15 | M   | 63   | Right       | Right  | CR, CI, INS    | 4            | 0.9               | 0.9           | 65           | 90         | 0.7                 | 0.6          | 66           |
| 16 | F   | 55   | Right       | Right  | PON            | 1            | 0.8               | 0.8           | 63           | 96         | 1.08                | 1.0          | 66           |
| 17 | M   | 48   | Left        | Left   | CR, SPL, MFG   | 3            | 0.8               | 0.8           | 56           | 97         | 0.7                | 1.06         | 66           |
| 18 | M   | 63   | Right       | Left   | CR              | 3            | 0.3               | 0.2           | 42           | 101        | 0.7                 | 0.6          | 63           |
| 19 | M   | 58   | Right       | Left   | CR, BG, INS    | 7            | 0.53              | 0.53          | 25           | 99         | 1.3                 | 1.0          | 64           |
| 20 | M   | 58   | Right       | Left   | CR, PON, CI, INS| 4            | 0.53              | 0.53          | 25           | 99         | 1.3                 | 1.0          | 64           |
| 21 | M   | 53   | Right       | Right  | CR              | 4            | 0.9               | 0.9           | 65           | 93         | 1.19                | 0.7          | 66           |
| 22 | M   | 68   | Right       | Right  | CR              | 4            | 0.62              | 0.62          | 57           | 96         | 1.14                | 0.8          | 66           |
| 23 | F   | 78   | Right       | Right  | CR              | 4            | 0.62              | 0.62          | 57           | 96         | 1.14                | 0.8          | 66           |
| 24 | F   | 47   | Right       | Right  | CR              | 3            | 0.9               | 0.9           | 66           | 93         | 0.8                 | 0.8          | 66           |
| 25 | M   | 50   | Right       | Right  | CR              | 7            | 0.7               | 0.7           | 65           | 99         | 0.7                 | 1.0          | 66           |
| 26 | M   | 65   | Right       | Left   | BG              | 5            | 0.15              | 0.15          | 26           | 108        | 1.14                | 0.96         | 66           |
| 27 | M   | 55   | Right       | Left   | MED             | 4            | 0.53              | 0.53          | 93           | 90         | 0.8                 | 0.95         | 30           |
| 28 | M   | 81   | Right       | Left   | PONS            | 5            | 0.15              | 0.15          | 20           | 85         | 0.72                | 0.7          | 60           |
| 29 | M   | 30   | Right       | Left   | CR, STG, INS   | 0            | 0.9               | 0.9           | 64           | 90         | 0.88                | 0.91         | 64           |

Mean ± std: 19 M, 11 F, 65 ± 13, 29 R, 17 L, 4 ± 3, 0.5 ± 0.34, 0.36 ± 0.26, 46 ± 21, 104 ± 17, 0.84 ± 0.3, 0.77 ± 0.3, 60 ± 14.5

Age is given in years. Grip force and NHP values are given in proportional values (affected/unaffected hand). Initial NIHSS, grip force and NHP taken 3–5 days post stroke. Residual NIHSS, grip force and NHP after passage of TAS, time after stroke in days. BG indicates basal ganglia. CI, internal capsule. CR, corona radiata. F, female. FP, insular cortex. FTG, lateral temporal gyrus. ITG, inferior temporal gyrus. MFG, middle frontal gyrus. MFG, middle temporal gyrus. POC, post-central gyrus. PRE, precuneus. PRC, pre-central gyrus. SFG, superior frontal gyrus. SPL, superior parietal lobule. STG, superior temporal gyrus. SFG, superior frontal gyrus. UEFM, upper extremity Fugl–Meyer score.
study based on fMRI data collected in an overlapping but smaller study population implementing the same grip task (Schulz et al., 2016). Excluding patients that performed the task but were unable to reach the target grip force did not change the resulting coupling patterns. The task effort (constant grip force of 5 kg or 20% of MVC) had no additional explanatory value for the TR-Coh of any connection within the network (Supplementary Table 3). For a comprehensive tabulation of mean coupling estimates for all frequency bands, see Table 2.

3.4. Prediction of parietofrontal connectivity by residual motor deficit

If the functional connectivity between aIPS and M1 of the ipsilesional hemisphere is enhanced in stroke patients compared with controls, we postulated that the strength of this connection should be inversely related to individual motor impairment. To test this, we used the relative (affected/unaffected side) grip force, NHP as well as the UEFM score to explain variance in parietofrontal connectivity in the stroke patient group. Each of these measures reflects different motor skills and visuomotor integration demands: The NHP relies on fine motor skills and dexterity, the grip force reflects muscle strength and the UEFM indicates active movement range and synergies of proximal and distal muscles. We found that the strength of TR-Coh between the aIPS and M1 in the alpha bands of the lesioned hemisphere could be predicted by the variance in residual fine motor skills as measured by NHP (LME, factor NHP lower alpha: \( p = 0.022 \), upper alpha: \( p = 0.044 \)) and in the grip force in the upper alpha and beta bands (LME, factor GRIP upper alpha: \( p = 0.045 \), beta: \( p = 0.007 \), Table 3). An increase in 10% of grip force was related to a decrease in TR-Coh of 0.025 (lower alpha), 0.033 (upper alpha) or 0.043 (beta). Additionally, an increase in 10% of NHP was related to a decrease in TR-Coh of 0.031 (lower alpha) and 0.033 (upper alpha) as seen in Fig. 3. Including the task effort (constant grip force of 5 kg or 20% of MVC) into the statistical model of grip force and TR-Coh in the upper alpha and beta bands led to no significant model improvement (LME, interaction GRIP x TASK upper alpha: \( p = 0.47 \), beta: \( p = 0.84 \)). In the control group, we found no significant relationship between grip force and TR-Coh between aIPS and M1 (Supplementary Table 4).

Patients with a high grip force or NHP three months post stroke are either mildly affected by the infarction (high initial motor function, Table 1) or well recovered (low initial motor function, Table 1). To learn about the role of the detected relationship between parietofrontal connectivity and the residual motor deficit for motor recovery, we tested whether motor recovery (quantified by the improvement in motor function from the initial assessment 3–5 days post stroke to the late assessment three months after stroke), would show a significant interaction with the motor performance values in explaining TR-Coh (Fig. 3). Such an interaction would be reflected in a different slope in the relationship between TR-Coh and residual motor function for patients with a high and low motor recovery. However, including motor recovery in the model for any of the frequency bands or motor performance tests did not explain additional variance, indicating that well recovered and mildly affected patients show low parietofrontal connectivity, as the control participants.

4. Discussion

We investigated connectivity between parietal and frontal motor areas during a grip task in 30 chronic stroke patients and 19 healthy control subjects and found a significantly stronger interaction between the aIPS and M1 in the stroke patients. Although parietofrontal coupling generally increased during the task in both groups, only the connection between the aIPS and M1 proved to be significantly enhanced in the ipsilesional hemisphere in patients, that is, contralateral to the affected upper extremity; this pattern generalized across the alpha and beta rhythms. Having utilized a completely different imaging modality, this finding successfully replicates results of previous investigations in which parietofrontal motor connectivity was assessed using fMRI and DCM, thereby substantiating the physiologic importance and validity of this connection independent of neuroimaging modality or choice of data analysis tools (Schulz et al., 2016). Furthermore, and importantly, in this larger patient group we found that the increased task-induced coupling was significantly related to residual motor deficit. A possible interpretation is that this intensified cross-talk between parietal and frontal motor areas subserves for post-
Table 2
Mean TR-Coh changes during the hand grip relative to rest with corresponding 95% confidence interval for stroke patients and control participants (A), as well as the group difference (TR-Coh stroke–controls) and corresponding p-values (LME fixed effect GROUP).

|                  | Stroke          | Control         |
|------------------|-----------------|-----------------|
|                  | Lower alpha     | Upper alpha     | Beta             | Lower alpha     | Upper alpha     | Beta             |
|                  | Mean (95% conf.)| Mean (95% conf.)| Mean (95% conf.)| Mean (95% conf.)| Mean (95% conf.)| Mean (95% conf.)|
| Lower alpha      |                |                |                |                |                |                |
| Upper alpha      |                |                |                |                |                |                |
| Beta             |                |                |                |                |                |                |
| cIPS-aIPS        | 0.00           | 0.04           | 0.00           | 0.04           | 0.06           | 0.02           |
| aIPS-M1          | 0.13***        | 0.07           | 0.30           | 0.10*          | 0.08           | 0.01           |
| cIPS-M1          | 0.05**         | 0.03           | 0.10           | 0.04**         | 0.03           | 0.08           |
| aIPS-PMV         | 0.06***        | 0.03           | 0.10           | 0.06           | 0.06           | 0.06**         |
| cIPS-PMV         | 0.08***        | 0.03           | 0.20           | 0.07***        | 0.03           | 0.08           |
| M1-PMV           | 0.02           | 0.04           | 0.10           | 0.09           | 0.06           | 0.00           |
| aIPS-SMA         | 0.06***        | 0.02           | 0.10           | 0.08***        | 0.03           | 0.03           |
| cIPS-SMA         | 0.03           | 0.04           | 0.10           | 0.02           | 0.04           | 0.08           |
| M1-SMA           | 0.03           | 0.04           | 0.10           | 0.02           | 0.02           | 0.04           |
| PMV-SMA          | 0.06***        | 0.03           | 0.10           | 0.01           | 0.02           | 0.01           |

Stars indicate significant changes within the group (one-sample t-test, *p < 0.05, **p < 0.01, ***p < 0.001).

|                  | Diff. | p-Value | Diff. | p-Value | Diff. | p-Value |
|------------------|-------|---------|-------|---------|-------|---------|
| cIPS-aIPS        | −0.02 | 0.60    | −0.02 | 0.50    | −0.03 | 0.60    |
| aIPS-M1          | 0.2** | 0.00    | 0.2** | 0.01    | 0.19**| 0.01    |
| cIPS-M1          | 0.01  | 0.70    | −0.02 | 0.40    | 0.01  | 0.90    |
| aIPS-PMV         | 0.00  | 1.00    | 0.00  | 1.00    | −0.02 | 0.40    |
| cIPS-PMV         | 0.02  | 0.50    | 0.03  | 0.30    | 0.01  | 0.70    |
| M1-PMV           | −0.08 | 0.10    | −0.07 | 0.10    | −0.05 | 0.20    |
| aIPS-SMA         | 0.01  | 0.80    | 0.03  | 0.20    | 0.05  | 0.06    |
| cIPS-SMA         | 0.04  | 0.09    | 0.04  | 0.05    | 0.04  | 0.10    |
| M1-SMA           | 0.04  | 0.30    | 0.02  | 0.60    | 0.06  | 0.20    |
| PMV-SMA          | 0.00  | 0.90    | −0.03 | 0.40    | 0.01  | 0.50    |

Stars indicate significant changes across the group (LME, factor GROUP, **p < 0.01).
stroke motor function.

### 4.1. Relevance to neurorehabilitation research

This finding is of relevance for motor rehabilitation research for several reasons: First, the target identification process for brain stimulation protocols used in a neurorehabilitative setting requires robust pathophysiologic findings that become apparent independent of the specific methodology used in data acquisition and analysis. For detected ‘abnormal’ states in the stroke lesioned brain to be successfully translated from an exploratory study to a clinical interventional trial, a functionally relevant role of that state for therapeutic goals is mandatory. Second, the EEG provides direct information about neural activity as it reflects an aggregate measure of synaptic potential of cortical neurons. There is recent converging evidence from animal and human invasive recordings that low-frequency oscillations in the electric field of the motor area (alpha and low-beta range) are directly linked to pyramidal neuron spiking activity (Haegens et al., 2011; Miller et al., 2012). Within the oscillatory code, information processing capacities of the brain are multiplexed through nested oscillations, and communication between neuronal populations is facilitated. Fries suggested a mechanism for neuronal communication through coherence (Fries, 2005). In this framework, coherence among neuronal groups ensures that they can interact effectively via the opening and closing of communication windows for input and output at the same time. The behavioral relevance for a synchronization of the low-frequency temporal reference frame among interacting regions has been abundantly reported across species, regions and cognitive systems (Arce-McShane et al., 2020).

### Table 3

Regression coefficients with 95% Confidence Interval of fixed effect clinical score (grip force, NHP, UEFM) for coupling strength at each ROI and frequency band in stroke patients. Grip force and NHP performance were both modeled as proportional values (affected/unaffected hand).

| Coef. | Lower | Upper | Coef. | Lower | Upper | Coef. | Lower | Upper |
|-------|-------|-------|-------|-------|-------|-------|-------|-------|
| GRIP  |       |       |       |       |       |       |       |       |
| cIPS aIPS | 0.00 | -0.19 | 0.18 | -0.05 | -0.20 | 0.14 | -0.23 | -0.50 | 0.02 |
| aIPS M1 | -0.25 | -0.52 | 0.02 | -0.33* | -0.60 | -0.01 | -0.43** | -0.70 | -0.12 |
| cIPS M1 | 0.01 | -0.12 | 0.13 | 0.01 | -0.10 | 0.15 | -0.04 | -0.20 | 0.07 |
| cIPS PMV | 0.00 | -0.13 | 0.12 | -0.03 | -0.10 | 0.09 | 0.08 | -0.08 | 0.24 |
| cIPS PMV | -0.02 | -0.13 | 0.10 | -0.09 | -0.20 | 0.03 | -0.10* | -0.20 | -0.02 |
| M1 PMV | -0.02 | -0.15 | 0.11 | 0.01 | -0.10 | 0.13 | 0.04 | -0.09 | 0.18 |
| aIPS SMA | 0.01 | -0.11 | 0.13 | -0.06 | -0.20 | 0.05 | 0.06 | -0.08 | 0.19 |
| cIPS SMA | 0.01 | -0.08 | 0.10 | 0.02 | -0.07 | 0.11 | -0.02 | -0.10 | 0.11 |
| M1 SMA | -0.06 | -0.21 | 0.08 | -0.09 | -0.20 | 0.05 | 0.02 | -0.10 | 0.17 |
| PMV SMA | -0.03 | -0.13 | 0.08 | 0.01 | -0.08 | 0.11 | 0.01 | -0.09 | 0.12 |
| Nine Hole Peg |       |       |       |       |       |       |       |       |
| cIPS aIPS | -0.01 | -0.19 | 0.17 | -0.02 | -0.21 | 0.16 | -0.15 | -0.40 | 0.10 |
| aIPS M1 | -0.31* | -0.57 | -0.05 | -0.33* | -0.65 | -0.01 | -0.20 | -0.52 | 0.14 |
| cIPS M1 | -0.05 | -0.18 | 0.07 | -0.07 | -0.21 | 0.07 | -0.05 | -0.16 | 0.07 |
| aIPS PMV | -0.04 | -0.14 | 0.11 | -0.08 | -0.20 | 0.03 | 0.13 | -0.02 | 0.28 |
| cIPS PMV | -0.04 | -0.15 | 0.08 | -0.09 | -0.21 | 0.02 | -0.02 | -0.11 | 0.06 |
| M1 PMV | -0.03 | -0.15 | 0.10 | -0.01 | -0.13 | 0.11 | 0.04 | -0.10 | 0.17 |
| aIPS SMA | -0.01 | -0.14 | 0.11 | -0.02 | -0.14 | 0.10 | 0.04 | -0.09 | 0.17 |
| cIPS SMA | 0.03 | -0.06 | 0.12 | 0.03 | -0.06 | 0.13 | 0.01 | -0.11 | 0.14 |
| M1 SMA | -0.08 | -0.22 | 0.07 | -0.02 | -0.16 | 0.12 | 0.10 | -0.05 | 0.25 |
| PMV SMA | -0.01 | -0.11 | 0.11 | 0.01 | -0.08 | 0.11 | 0.03 | -0.07 | 0.14 |
| UEFM  |       |       |       |       |       |       |       |       |
| cIPS aIPS | 0.00 | 0.00 | 0.01 | 0.00 | 0.00 | 0.01 | 0.00 | -0.01 | 0.01 |
| aIPS M1 | 0.00 | -0.01 | 0.00 | 0.00 | -0.01 | 0.00 | -0.01 | -0.01 | 0.00 |
| cIPS M1 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 |
| aIPS PMV | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 |
| cIPS PMV | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 |
| M1 PMV | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 |
| aIPS SMA | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 |
| cIPS SMA | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 |
| M1 SMA | 0.00 | -0.01 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.01 |
| PMV SMA | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 |

Stars indicate significant fixed effect of motor value (GRIP, NHP or UEFM) on TR-Coh (LME, "p < 0.05, **p < 0.01, uncorrected).
et al., 2016; Gross et al., 2001; Hummel and Gerloff, 2005). And also in the course of recovery from stroke, neuroplastic processes have shown to be paralleled by abnormal coherence within and between cortical sites (Nicolo et al., 2015; Wu et al., 2014).

4.2. Why do stroke patients show increased coupling between parietal and frontal areas during a grip task?

It is conceivable that connectivity patterns vary over time and depend both on the severity of stroke and the level of recovery thereafter. Previous studies on stroke recovery have found that in the acute stage, stroke patients have reduced facilitatory coupling from secondary motor areas to the contralateral (ipsilesional) M1; additionally, they experience reduced inhibitory coupling from secondary motor areas to the ipsilateral (contralesional) M1 (Rehme et al., 2011b). Recovery is accompanied by a normalization of this abnormal pattern but in the chronic stage, conditional on the degree of deficit, an enhanced inhibitory coupling from contralesional to ipsilesional M1 can be detected (Grefkes et al., 2008b; Rehme et al., 2011a). Albeit these abnormal connectivity patterns which help understand variable recovery patterns between stroke patients, we currently lack a precise neurophysiological framework. In the present study, we found that in a simple visuomotor grip task, stroke patients show an increased synchrony between the aIPS and M1 (Table 2), and with a higher residual motor deficit, this synchrony was also higher (Table 3, Fig. 3). Given the established importance of the parietal cortex for visuomotor and sensorimotor integration, both essential components in human everyday motor behaviour, this finding might add an important and conceptually plausible missing link.

It is likely that in the post-stroke brain, higher integrative demands are required for task performance that rely on specific features such as visuomotor translation of the cue position, somatosensory integration, hand shaping and visual feedback information processing into force adaptation, all of which have been related to parietal motor area activity also in the healthy human brain (Bernier et al., 2017; Klaes et al., 2015; Konen et al., 2013; Murata et al., 2000; Sakata and Kusunoki, 1992; Sakata et al., 1997; Taira et al., 1990; Vingerhoets, 2014). The visuomotor integrative role of the dorsal parietal cortex was further differentiated to provide corrective movement plans to goal perturbation (Tunik et al., 2008). Such on-line adaptive adjustments of force output to small changes in the bar height were likely stronger in the stroke patients (Supplemental Fig. 2).

Direct projections to distal hand motoneurons which provide the posterior parietal cortex with the potential to control motoneuron activity directly at the spinal level, have recently been detected in monkeys (Rathelot et al., 2017), undermining the parietal cortex as a ‘command apparatus’ for hand movements. The specific information is more effectively relayed if the windows for communication between the parietal cortex and primary motor areas are opening and closing at the same points in time, indicated by higher synchrony between the oscillations at aIPS and M1.

The plastic reorganization and remodeling process could likewise lead to a greater contribution of alternative motor tracts arising from frontal and parietal sites to the corticospinal tract (McNeal et al., 2010; Newton et al., 2006; Puig et al., 2017; Schulz et al., 2015; Schulz et al., 2012). We propose that by synchronizing independent neural computations across task-involved regions, which together give rise to corticospinal projections, the summation of spikes becomes more effective at driving postsynaptic neurons at lower spinal levels (Salinas and Sejnowski, 2001). Thereby, a lesion-induced loss of cortical signal generating areas and conducting fibers is compensated. Importantly, the above interpretations regarding synchronization as a mean for task-specific information relay across cortical sites and synchronization as a mean for maximizing effect of presynaptic spiking activity on lower motor neurons, are not mutually exclusive but could have overlapping and synergistic function.

4.3. What is the relevance of the frequency bands of coupled oscillations?

We a priori focused our analysis on motor-relevant frequency bands based on existing studies pertaining to task-induced changes in the power or coherence spectrogram in stroke patients (Gerloff et al., 2006). The crucial characteristic of alpha is a functional inhibition or engagement of task-irrelevant areas by amplitude down- and upregulation, a view that is supported by ample experimental evidence across cognitive systems, species and brain locations. A higher alpha rhythm over the sensorimotor cortexes is specifically reactive during motor tasks (termed the mu rhythm), whereas the lower, or classic alpha rhythm, is reactive in the visual and general attentive system. In the motor system, beta generally shows a high conformity and/or overlap with alpha rhythms, but a specific role has been carved out for corticospinal coherence especially during hold periods of motor tasks (Chen et al., 2013). Thus, a status quo view of beta signaling emerged, i.e. the signaling responsible for the maintenance of current sensorimotor or cognitive states (Engel and Fries, 2010). However, it was shown for both rhythms that local spiking activity is structured by the oscillation of peaks and troughs in a pulsed manner (Haegens et al., 2011; Miller et al., 2012), see above, compatible with the view that both play predominantly top-down directing roles, although clearly distinctive functions for each rhythm are not extractable from the literature. Rossiter et al. found reduced movement-related beta desynchronization (15–30 Hz) in contralateral (ipsilesional) M1 during a visually guided grip task in patients compared with control subjects which were also related to the degree of motor deficit (Rossiter et al., 2014). In our results, we found a similar qualitative relation, but the group difference was not statistically significant (mean TR-Pow 20 Hz M1 patients: −0.42, controls: −0.44; LME, p = 0.67). A few studies have investigated resting-state oscillatory phase coupling networks in stroke patients which revealed a predictive value of ipsi- and contralateral connectivity for motor recovery in the alpha band (8–12 Hz), (Wempe et al., 2012; Wu et al., 2011), low beta band (13–16 Hz) (Nicolo et al., 2015), high beta band (20–30 Hz (Wu et al., 2014), 24–33 Hz (Pellegrino et al., 2012)) or broadband beta (13–30 Hz) (De Vico Fallani et al., 2013).

Taken together, the similarity between the connection profiles across frequency bands is in accordance with current concepts of the rhythms’ functional roles: disinhibition of local circuitry in motor-relevant areas, signal transmission for the maintenance of the constant force output throughout the hold phase of the grip task and constant integration of visual feedback and integration in the motor command.

4.4. Predictive value of parietofrontal connectivity for residual motor deficit

An additional finding of this study was that apart from aIPS-M1 functional coupling being higher in stroke patients compared with controls, the degree of coupling correlated with the residual motor deficit. This finding is consistent with aforementioned fMRI findings in frontal motor network architecture, where increased inhibitory coupling from contralesional to ipsilesional M1 was significantly and positively related to residual motor deficit (Rehme et al., 2011a). The increased aIPS-M1 coupling could be an expression of a task-specific network adaptation, meaning that the coupling is directly related to the visuomotor demands of the task. Alternatively, it could be an expression of post-stroke network reorganization that is incidentally revealed by this task. A causal involvement of aIPS-M1 coupling in meeting task demands is difficult to assess with pure observational techniques. However, since the experimental design included a task in which the applied grip force was 20% of the maximal grip force, we can conclude, that TR-Coh is not only related to the clinical impairment as measured by grip force and the NHP, but likewise related to the task specific requirement of force generation. However, this does not rule out the possibility that the M1-aIPS coupling is an unspecific property of the stroke-lesioned brain. Furthermore, the qualitative nature of high
parietofrontal connectivity remains to be elucidated: Is the high parietofrontal connectivity causing a functional impairment (maladaptive) or is it an (unsuccessful) attempt to generate motor output (adaptive). Such causal links can only be elucidated with brain stimulation (Di Pino et al., 2014) or neurofeedback (Enriquez-Geppert et al., 2017) protocols.

In our results, a low motor function was associated with high parietofrontal connectivity and a high motor function was associated to low parietofrontal connectivity. A high motor function in the chronic stage can either be attributed to a mild initial impact of the stroke or a good motor recovery. Therefore, we analyzed if the group of patients with a high motor function could be dissociated into well recovered (high motor recovery) and mildly affected (low motor recovery) patients based on their TR-Coh values. Such a pattern would be reflected in a different slope in the relationship between TR-Coh and residual motor function for patients with a high and low motor recovery. We did not find a significant relationship between motor recovery and residual motor function in explaining variance in TR-Coh three months after stroke. From this data it is attractive to speculate, that the downregulation of initially upregulated parietofrontal connectivity is a part of successful motor recovery. However, since we lack information on initial TR-Coh values, we are unable to confirm this point with the present study.

A longitudinal recording of EEG during the course of recovery could further explain if a) successful recovery goes along with a reduction of initially high parietofrontal connectivity or b) patients who do not recover from severe initial impairment upregulate parietofrontal connectivity over time.

Apart from those conceptual limitations regarding the interpretational scope of our study, another important limitation affects spatial accuracy of the source reconstruction (Schulz et al., 2016). This work was supported by the German Research Foundation (SFB 936 C1 to CG, C2 to GT) and German National Academy of Sciences Leopoldina (Fellowship programme grant number SFB43032 to MB).
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