Localization of cortico-peripheral coherence with electroencephalography

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Abstract

BACKGROUND: The analysis of coherent networks from continuous recordings of neural activity with functional MRI or magnetoencephalography has provided important new insights into brain physiology and pathology. Here we assess whether valid localizations of coherent cortical networks can also be obtained from high-resolution electroencephalography (EEG) recordings.

METHODS: EEG was recorded from healthy subjects and from patients with ischemic brain lesions during a tonic hand muscle contraction task and during continuous visual stimulation with an alternating checkerboard. These tasks induce oscillations in the primary hand motor area or in the primary visual cortex, respectively, which are coherent with extracerebral signals (hand muscle electromyogram or visual stimulation frequency). Cortical oscillations were reconstructed with different inverse solutions and the coherence between oscillations at each cortical voxel and the extracerebral signals was calculated. Moreover, simulations of coherent point sources were performed.

RESULTS: Cortico-muscular coherence was correctly localized to the primary hand motor area and the steady-state visual evoked potentials to the primary visual cortex in all subjects and patients. Sophisticated head models tended to yield better localization accuracy than a single sphere model. A Minimum Variance Beamformer (MVBF) provided more accurate and focal localizations of simulated point sources than an $L_2$ Minimum Norm (MN) inverse solution. In the real datasets, the MN maps had less localization error but were less focal than MVBF maps.

CONCLUSIONS: EEG can localize coherent cortical networks with sufficient accuracy.

Keywords: EEG, beamformer, minimum-norm, functional connectivity, imaginary coherence

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1. Introduction

The brain is a complex network of dynamic systems with abundant functional interactions between local and more remote brain areas (Varela et al., 2001). Interregional neural communication is thought to be accompanied by a synchronization of oscillations between different brain regions (Aertsen et al., 1989; Engel et al., 1992; Gray et al., 1989; Gray and Singer, 1989). This interregional synchronization can be quantified with the concept of “functional connectivity” which is a measure of similarity between activity in different brain regions and is therefore considered to be an index of functional interaction (Gross et al., 2001; Lachaux et al., 1999; Nolte et al., 2004; Nunez et al., 1997; Stam et al., 2003). Studies using functional magnetic resonance imaging (fMRI) in healthy humans have shown that spontaneous fluctuations of brain activity at rest are highly organized and coherent within specific neuro-anatomical systems (Fox et al., 2005; Fransson, 2005). Thus, a careful analysis of coherence between brain regions gives access to the functional brain organization. Furthermore, the pattern of coherence between brain regions observed with fMRI at rest corresponds to the pattern of brain activation induced by corresponding tasks (Vincent et al., 2007), thus showing that techniques relying on functional connectivity do not depend on a correct task participation of the patients. Functional connectivity therefore opens an exciting and accessible window for a non-invasive assessment of brain function.

The concept of functional connectivity can be applied to any measurement of brain activity, such as fMRI, magnetoencephalography (MEG), or electroencephalography (EEG). EEG was the first brain signal to be used for calculating functional connectivity (Lopes da Silva et al., 1973; Thatcher et al., 1986). However, unlike MEG, it has the disadvantage that the indices of functional interaction calculated between sensors cannot be attributed to brain regions of interest or to structural lesions. Field spread leads to a wide representation of sources in many sensors, which makes the interpretation of functional connectivity measures between sensor pairs difficult (Schoffelen and Gross, 2009; Srinivasan et al., 2007).

For MEG, it has been shown that this limitation can be overcome by combining measures of functional connectivity with source localization algorithms. For instance, such “functional connectivity maps” (FCMs) can correctly localize the cortical generators of cortico-muscular interaction to the primary motor cortex (Gross et al., 2001; Jerbi et al., 2007; Schoffelen et al., 2008). We have shown that MEG maps of resting state intracortical interaction can correctly localize dysfunctional cortex in patients with brain tumors and distinguish it from tumor-infiltrated areas that are still functional (Guggisberg et al., 2008a; Martino et al., 2011).

Given the limited availability and high costs of MEG, it would be desirable to use source localization combined with functional connectivity measurements also for EEG recordings (Astolfi et al., 2005; De Vico Fallani et al., 2007). In principle, the algorithms used in MEG studies can be applied also to EEG, but there are differences between EEG and MEG signals that need to be considered.

- Electric fields are distorted to a greater extent than magnetic fields by the inhomogeneous conductivities of the brain, skull, and scalp. EEG is therefore more dependent on accurate head models to resolve the forward problem than MEG. The analysis of patients with brain lesions could also be problematic because of different conductivity of lesion tissue compared to healthy tissue.

- When functional connectivity is assessed between EEG sensors, the choice of the reference electrode critically influences and potentially biases the results (Fein et al., 1988; Guevara et al., 2005). Projecting EEG sensor data through inverse solutions to the brain effectively eliminates the reference problem of EEG studies, since the data becomes reference-free (Fender, 1987; Geselowitz, 1998). However, previous studies have reported that the choice of the reference electrode can lead to minor variations in the output of minimum variance beamformers (MVBF) (Van Veen et al., 1997).

This study aimed to evaluate the feasibility of localizing cortico-peripheral coherence with EEG in healthy subjects and patients with brain lesions, and to compare the performance of different analysis parameters. More specifically, we assessed the influence of the head model, the inverse solution, the method for determination of dipole orientations, and (for beamformers) the reference electrode on the accuracy of functional connectivity localization.

To this end, we collected two sets of real data in which cortical nodes with known location oscillate in synchrony with extracerebral activity.

- Tonic muscle contraction is known to induce beta frequency (~15-30 Hz) oscillations in the contralateral primary motor cortex which are coherent with correlated electromyographic (EMG) activity (Mima and Hallett, 1999; Murthy and Fetz, 1992). Hence, the algorithms should localize cortical activity that is coherent with EMG activity to the contralateral primary motor cortex.

- Rhythmic visual stimulation is known to induce a steady state visual evoked potential (SSVEP) in the primary visual cortex, which oscillates in the frequency of the stimulation and harmonics (Bianciardi et al., 2009; Emir et al., 2008; Victor et al., 1994). Hence, the algorithms should localize cortical oscillations that are coherent with the stimulation frequency to the primary visual cortex.

In order to test whether the EEG localizations remain valid in patients with brain lesions, we obtained the data not
only from healthy subjects but also from patients with unilateral territorial ischemic stroke.

In addition, we carried out simulations of point sources to evaluate the performance of different inverse solutions.

2. Methods

2.1. Subjects and patients

Two healthy subjects and four patients with ischemic stroke participated in this study (mean age 44, age range 29-70, 3 women). The lesions were located in the territory of the left middle cerebral artery in 3 patients and of the right anterior cerebral artery in one patient (Figures 1 and 2, blue arrows).

2.2. Recordings

Recordings were obtained with a 128-channel Biosemi ActiveTwo system (Biosemi V.O.F. Amsterdam, Netherlands), using a sampling rate of 512 Hz. Impedances were kept below 5 kΩ.

Bad EEG channels as determined by visual inspection were excluded. The data was re-referenced to the average of all non-excluded channels. We additionally created a second version of EEG data by re-referencing to the vertex electrode (Cz), in order to assess the influence of the reference montage on beamformers (see below).

2.3. Corticomuscular coherence task

EEG and EMG of the first dorsal interosseus (FDI) muscle were recorded simultaneously in 1 healthy subject and 3 patients during a tonic muscle contraction task. The contractions were performed with the non-paretic hand ipsilateral to the lesioned hemisphere in 2 patients with hemiparesis, and with the hand contralateral to the lesioned hemisphere in one patient without hemiparesis. Bipolar surface EMG was recorded at 512 Hz with the active EMG electrode fixed on the FDI muscle belly and the reference electrode ~4 cm proximal over the volar part of the radius head. The participants were visually instructed by a hand symbol displayed on a monitor to exert about 50% of their maximum force during repeated periods of 10 s which were separated by pauses of 5 s. The EEG and EMG data were high-pass filtered at 10 Hz with a fifth-order ellipsoid IIR filter. Forward and backward filtering was applied to obtain zero-phase distortion. Segments with artifacts as well as the resting periods were excluded. A total of at least 3 min of artifact-free EEG and EMG signal during FDI contraction were obtained for each participant.

2.4. Steady state visual evoked potential (SSVEP) task

EEG recordings during continuous visual stimulation with a bilateral, alternating, black and white checkerboard pattern were obtained in 2 healthy subjects and 2 patients with ischemic stroke. All patients had a normal visual field as determined clinically. Participants were asked to
Figure 2: FCMs of steady-state visual evoked potentials (SSVEP) oscillating coherently with the visual stimulation frequency obtained from 2 healthy subjects and 2 patients with ischemic brain lesions (marked with blue arrows). The IC is indicated as % of the individual maximum value at the peak voxel. The peak absolute imaginary coherence magnitude was 0.49 ± 0.07 (mean ± SD). IC at the stimulation frequency was largest around the primary visual cortex in all participants. The same analysis parameters as in Figure 1 were used.

2.5. Structural imaging

A high-resolution, T1-weighted cerebral MRI covering the entire skull was obtained for each participant. In addition, a MNI standard volume consisting of the average of 152 healthy subjects was used for coregistration in order to assess the impact on localization accuracy.

2.6. Head models and lead-potential

Each subject’s brain was segmented into scalp, skull, gray and white matter with the software BrainSuite09 (http://www.loni.ucla.edu/~shattuck/brainsuite/) written by David Shattuck. The gray matter volume was divided into a regular 10 mm grid. Three different head models were used to calculate the lead-potential for each grid location.

Spherical model with anatomical constraints (SMAC)

The SMAC head models were created with the software SMAC Toolbox (Functional Brain Mapping Laboratory, University of Geneva) written by François Tavel. The algorithm of this head model is described elsewhere (Spinelli et al., 2000). In short, the individual T1 MRI was transformed to a best-fitting sphere. The solution points were defined in the gray matter tissue of this deformed MRI and the lead field was computed using an analytical solution for a 3-shell spherical head model (Ary et al., 1981) which assumes conductivities of 1:0.0125:1 and radii of 1:0.92:0.87 for scalp, skull, and brain, respectively (Rush and Driscoll, 1968).

Berg 3-shell single-sphere head model

This model was created with the NUTEEG plugin (Wong and Gordon, 2009) of NUTMEG (http://nutmeg.berkeley.edu) (Dalal et al., 2011). A single sphere with the best fit to the individual skull shape was defined and 3 concentric shells with radii of 1:0.93:0.85 relative to the fitted sphere were created (Berg and Scherg, 1994). The solution points were confined to the gray matter as defined by the segmented MRI of the subject. The conductivities for scalp, skull, and brain, were set to the same standard values as for SMAC.
A 3-shell BEM model was created with the Helsinki BEM library (http://peili.hut.fi/BEN/) (Stenroos et al., 2007) and the NUTEEG plugin of NUTMEG. The BEM approach constructs a realistic head model by defining discrete 3-D compartments comprising the brain, scalp, and skull. Each compartment is assumed to have uniform electrical conductivity. Together with the shape of each compartment, this information is then used to compute the voltage at the electrode positions expected to result from a source with a given position and orientation (Mosher et al., 1999). We created triangular meshes for scalp, and outer and inner skull surface based on the segmented T1 MRI (with 1000, 1500, and 2000 nodes, respectively). The lead-potential was calculated over the grid of gray matter solution points. The conductivities for each shell were set identically to the SMAC and Berg models.

The electrode positions were determined by fitting and projecting the spherical standard electrode coordinates to the individual scalp form.

2.7. Scalarization of the lead-potential

For the localization of functional connectivity in the brain, we require an estimation of neural network oscillations at each solution point. Vector weights obtained from the inverse solution allow only reconstructing squared power values for a given location; the reconstruction of neural oscillations requires a scalar weight matrix. In order to scalarize the lead-potential as input to scalar weights computation, we need to determine the dipole orientation at each grid location. Here, we compared two different techniques to achieve this.

(i) Computation of an optimal dipole orientation yielding maximum output SNR. For the MVBF, the optimum orientation at each dipole \( r \) is given by (Sekihara et al., 2004):

\[
\eta_{\text{opt}}(r) = \min_{v} \{ |L^T(r)R^{-1}L(r)|^{-1}|L^T(r)R^{-2}L(r)| \},
\]

where \( v_{\min} \) is the eigenvector corresponding to the minimum eigenvalues of the matrix in \( \{ \} \), \( L \) is the vector lead-potential, \( R \) the sensor covariance matrix, and superscript \( T \) denotes the matrix transpose.

For the MN inverse solution, it is obtained as (Pascual-Marqui et al., 2009):

\[
\eta_{\text{opt}}(r) = \max_{v} \{ L^T(r)G^{-1}RG^{-1}L(r) \},
\]

where \( v_{\max} \) the eigenvector corresponding to the maximum eigenvalues of the matrix in \( \{ \} \), and \( G = LL^T \) is the Gram matrix.

(ii) Anatomically constrained surface normal dipole orientations. A model of the individual cortical surface was created for each participant with BrainSuite09. The number of vertices of the cortical surface model was down-sampled to 8000. For each solution point, the dipole orientation was defined as normal to the orientation of the closest cortex surface with the NUTEEG plugin of NUTMEG.

The scalar lead-potential was then calculated as:

\[
l(r, \eta_{\text{opt}}) = L(r)\eta_{\text{opt}}(r)
\]

and will henceforth be designated \( l(r) \) for simplicity.

2.8. Inverse solutions

Inverse solutions were calculated with NUTMEG (Dalal et al., 2011). A spatial filter matrix \( w \) was calculated for estimation of the signal at each grid location. Two different inverse solutions were compared. The minimum variance beamformer (MVBF) (Van Veen et al., 1997; Vrba and Robinson, 2001; Sekihara et al., 2001, 2005) is an adaptive spatial filter which uses the sensor covariance matrix \( R \) of the entire recording to estimate the spatial filter matrix \( w \).

\[
w_{\text{MVBF}}(r) = \frac{R^{-1}l(r)}{l^T(r)R^{-1}l(r)}
\]

For our average-referenced EEG datasets, we used the Moore-Penrose pseudoinverse to estimate \( R^{-1} \) for the following reason. An average reference introduces a linear dependence between the channels, such that any given channel can be calculated as a sum of the remaining channels, reducing the rank of the data matrix by exactly 1. The sensor covariance matrix then becomes rank-deficient and invertible only with regularized inversion methods. An alternative solution to this problem is to apply a rotation to the rank-deficient covariance matrix such that the last column contains zeros, and the remaining columns contain independent data. The same rotation is then also applied to the lead-potential, to keep the contributions of each channel identical. Both the pseudoinverse and the matrix rotation approach produced identical beamformer output for the tested data.

The covariance can also become ill-conditioned when using especially short data lengths or bandwidths, in which case Tikhonov regularization procedures can be applied (Brookes et al., 2008). This was however not the case in our datasets.

In order to ensure that the MVBF performance was not sensitive to the choice of the data bandwidth used for calculation of the covariance matrix, we compared three different bandwidths for the analysis of corticomuscular coherence in 2 subjects. One subject had the individual corticomuscular coherence peak at 24 Hz, the second subject at 26 Hz. The frequency bands \( >10 \) Hz, 20-30 Hz, and peak coherence \( \pm 1 \) Hz were compared. The resulting FCMs were highly correlated among each other \( r > 0.78 \), and the coherence peaks located within the same 10 mm radius. For the main analyses of the corticomuscular dataset, the sensor covariance was calculated from the data that was high-pass filtered at 10 Hz. A bandwidth of 4 to 6 Hz was used for the SSVEP dataset.
An alternative version of a beamformer is “dynamic imaging of coherent sources (DICS)” (Gross et al., 2001). DICS uses correlations in the frequency domain (a cross-spectrum density matrix) for source estimation instead of temporal correlations (sensor covariance matrix) used for the MVBF. Given the similarity of the two approaches, DICS was not separately tested in this study.

The $L_2$ Minimum Norm (MN) is a non-adaptive inverse solution which tries to minimize the overall 3-D DICS was not separately tested in this study.

Given the similarity of the two approaches, DICS uses correlations in the frequency domain (a cross-

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J

d

ize the MN or WMN solution by the variance of the esti-

tivity (Dale et al., 2000; Lin et al., 2004). These variants

are not evaluated here.

The inverse solutions sLORETA (Pascual-Marqui, 2002) and dSPM (Dale et al., 2000) additionally standardize the MN or WMN solution by the variance of the estimated current density $J$:

$$w_{sMN}(r) = w_{MN}(r)J(r)$$

We show below that the standardization $J$ is canceled out during calculation of source space coherence. The sLORETA and dSPM algorithms therefore yield the same source coherence results as (W)MN algorithms.

### 2.9. Functional connectivity in source space

Localization of coherence in source space was done with the FCM plugin of NUTMEG written by AGG. The EEG, EMG and visual stimulation sinusoid were divided into non-overlapping data segments of 1 s duration (512 data points). A fast Fourier transform (FFT) was performed for each data segment and sensor. For the corticomuscular coherence, a multi-taper approach was used (Mitra and Pesaran, 1999). Each data segment was tapered using a set of discrete prolate spheroidal sequences (Slepian functions). The number of tapers used determines the amount of spectral smoothing. As described previously (Schoffelen et al., 2008), we maximized the sensitivity of subsequent analysis steps by determining the peak frequency and the amount of spectral smoothing individually for each subject. To this end, we computed the coherence between the EMG and the contralateral central electrode (C3 or C4) using different numbers of tapers. We then selected the number of tapers which produced a coherence peak with a width corresponding approximately to the smoothing bandwidth. The frequency was set at the coherence peak. For the SSVEP dataset, the Fourier coefficients were calculated for the frequency band between 4 and 6 Hz.

The complex Fourier coefficients $F$ of all electrodes at frequency $f$ were then projected through the spatial filter $w$ into the source space:

$$d(r, f) = w^T(r)F(f)$$

This is equivalent to first projecting the filtered data to source space and then calculating the FFT at each grid location (Guggisberg et al., 2008a), but is computationally more efficient.

The complex coherence between neural oscillations at each grid location and either the EMG or the visual stimulus sinusoid could then be computed as

$$CC(r, f) = \frac{\sum_{k=1}^{K} d_k(r, f)^* E_k(f)}{\sqrt{\sum_{k=1}^{K} |d_k(r, f)|^2 \sum_{k=1}^{K} |E_k(f)|^2}}$$

where $k = [1, ..., K]$ stands for the data segments, $*$ for the complex conjugate, and $E$ for the Fourier transformed extracerebral signals (EMG or visual stimulus sinusoid), averaged (in the case of EMG) across all tapers.

From the complex coherence, the absolute imaginary component (IC) (Nolte et al., 2004) as well as the magnitude squared coherence (MSC) were computed:

$$IC = |\Im{CC}|$$

$$MSC = |CC|^2$$

When calculating cortico-peripheral coherence with sLORETA or dSPM, we obtain from equations 6, 7 and 8 above:

$$CC(r, f) = \frac{\sum_{k=1}^{K} w_{MN}(r)J(r)F_k(f)^* E_k(f)}{\sqrt{\sum_{k=1}^{K} |w_{MN}(r)J(r)F_k(f)|^2 \sum_{k=1}^{K} |E_k(f)|^2}}$$

$$= \frac{J(r)}{\sqrt{|J(r)|^2}} \frac{\sum_{k=1}^{K} w_{MN}(r)F_k(f)^* E_k(f)}{\sqrt{\sum_{k=1}^{K} |w_{MN}(r)F_k(f)|^2 \sum_{k=1}^{K} |E_k(f)|^2}}$$

Since the elements of $J$ are positive real, $J$ is canceled out during calculation of cortico-peripheral coherence. The same can be shown for calculations of cortico-cortical coherence. Hence, standardized (W)MN variants such as sLORETA and dSPM do not provide additional advantage over (W)MN for the localization of coherence.

### 2.10. Visualization

3-D renderings of the FCMs were performed with Cartool (Brunet et al., 2011) (http://sites.google.com/site/cartoolcommunity/home).
2.11. Evaluation of Performance

The anatomical centers of the hand notch and of the interhemispheric junction of the calcarian sulcus were defined individually in the co-registered MRIs. The distance between the FCM peak coherence and these anatomical targets were used as an index of FCM accuracy.

2.12. Simulations

We simulated single cortical point sources with a continuous 10 Hz sinusoidal rhythm while the remaining cortical grid locations were set to Gaussian random oscillations. An additional extracerebral 10 Hz oscillation was defined with a radial phase lag of $\pi/2$ (= 25 ms) relative to the cortical point source. This phase difference leads to maximal values in the imaginary part of coherence. The dipole orientations were fixed to be normal to the cortical surface. The cortical sources were then projected to the EEG sensors by using a scalar lead-potential calculated with a SMAC head model. Four different levels of Gaussian white random noise were added to the sensors (SNR 1, 2, 5, or 10). The simulated sensor data was then bandpass filtered between 1 and 20 Hz, and projected back to all gray matter grid locations through a spatial filter matrix calculated with the MVBF and MN inverse solutions described above. The IC between the extracerebral sinusoid and all cortical grid locations was then computed for the 10 Hz frequency bin. The distance between the IC peak and the source coordinate was then calculated to determine the localization error. The spatial spread of the obtained maps was quantified as the full-width-half-maximum (FWHM) around the IC peak. The sources were placed randomly in the cortex of healthy subject 1. The results of 20 different locations distributed throughout the entire cortex were averaged.

2.13. Statistics

A three-way analysis of variance (ANOVA) was used to compare the peak-to-peak error and FWHM of FCM maps obtained with different head models (SMAC vs. single sphere vs. BEM), methods for determining the dipole orientation (maximum output SNR vs. surface normal), and inverse solutions (MVBF vs. MN).

3. Results

3.1. Real Data

Figures 1 and 2 demonstrate that EEG-based FCMs correctly localized the imaginary component of cortico-muscular coherence to the primary motor cortex and SSVEP oscillations to the primary visual cortex in all subjects and in all patients with brain lesions.

Figure 3 visualizes the peak-to-target error of FCMs obtained with different analysis parameters. The choice of the inverse solution had a significant impact on the accuracy of imaginary coherence localizations, with MN performing significantly better than the MVBF, both in the cortico-muscular coherence (overall mean ± SD: MN, 21±4 mm; MVBF, 37±12 mm; $F_{1,47}=10.2$, $p=0.0028$) as well as in the SSVEP dataset (MN, 22±7 mm; MVBF, 34±14 mm; $F_{1,47}=8.6$, $p=0.0057$). On the other hand, MVBF produced maps with less spatial spread than MN in both the corticomuscular coherence (44 vs. 65 mm, $F_{1,47}=10.5$, $p=0.0025$) and the SSVEP dataset (36 vs. 46 mm, $F_{1,47}=3.8$, $p=0.060$). The choice of the head model and of the method for determining the dipole orientation did not significantly affect the localization accuracy in either dataset ($p>0.28$), although Figure 3 shows a non-significant difference in performance in favour of SMAC and BEM head models as compared to a single Berg sphere model.
(overall mean of both datasets and both inverse solutions: SMAC, 26±3 mm; Berg, 32±8 mm; BEM, 27±3 mm). With an optimal choice of parameters (SMAC or BEM head model, MN inverse solution), the peak-to-target localization error was <20 mm for both datasets.

When comparing average-referenced to vertex (Cz)-referenced sensor data, we found a non-significant but non-vanishing difference in MVBF localization accuracy for either dataset (p≥0.33). The sources of this difference will be discussed below.

When a standard MNI template instead of the individual MRI was used for the creation of a SMAC head model, the peak error of localizations obtained with MN was larger for the corticomuscular coherence dataset (27 mm instead of 19 mm), but not for the SSVEP dataset (18 mm in both cases).

The performance of MSC maps is shown in Figure 4, and was not significantly different from the performance of imaginary coherence maps (F1,95=0.58, p=0.45). Again, the inverse solution had a significant influence on performance in both datasets (F1,47 ≥7.9, p<0.008), in favor of MN. The choice of the head model (F1,47 ≤2.5, p≥0.1) and the method for determination of the dipole orientations (F1,47 ≤0.48, p≥0.49) were not associated with significantly different localization errors.

3.2. Simulations

The MVBF localized the single point sources with significantly less error (F1,159=9.9, p=0.002) and less spatial spread (F1,159=25.2, p<0.0001) than MN (see Figure 5). As expected, the SNR only influenced the spatial spread of the MVBF (F1,79=29.2, p<0.0001), but not of the MN solution (F1,79=0.32, p=0.57)

4. Discussion

This study demonstrates that coherent sources can be localized with reasonable accuracy from EEG data. With optimal analysis parameters, we observed a mean distance between the center of the anatomical target regions and the FCM peaks of <20 mm. Given that both FCMs and anatomical targets had extensions of >20 mm, this corresponds to a good overlap (shown in Figures 1 and 2), which is sufficient for localization of large-scale networks across the entire cortex. Hence, the analysis of coherent functional networks which has proven to offer exciting new possibilities in fMRI (Carter et al., 2010; He et al., 2007; Raichle and Snyder, 2007; Vincent et al., 2007) and MEG studies (Castellanos et al., 2010; de Pasquale et al., 2010; Guggisberg et al., 2008a; Jerbi et al., 2007) is also possible with EEG data. In comparison with fMRI, EEG has the disadvantage of lower spatial resolution, but the advantage that it can assess actual neural oscillations at different spectral frequencies. The localization of corticomuscular coherence, for example, would not be possible with fMRI. Furthermore, EEG recordings are less costly and more widely available than MRI or MEG recordings. Unlike MEG, EEG may also be recorded simultaneously with fMRI.

A methodological limitation of our study is that we did not individually record the position of EEG electrodes, but merely fit standard coordinates to the individual head shape. The usage of digitized electrode positions may improve the localization accuracy reported here. However, the corresponding infrastructure is not generally available and our data shows that reasonably precise localizations can be obtained even with this approach.

4.1. Inverse solutions

The most important parameter for localization accuracy and distribution was the choice of the inverse solution. Non-adaptive algorithms provided significantly better localization accuracy in real datasets than adaptive algorithms. On the other hand, beamformers yielded more accurate localizations of point sources in the simulations. This apparent contradiction can be explained with the principle of operation of the inverse solutions. The MVBF uses the temporal covariance of the EEG data (in addition to the sensor geometry) to create a custom spatial filter depending on the signal characteristics. This en-
ables more precise and focal source localizations (Sekihara et al., 2005; Dalal et al., 2008; Guggisberg et al., 2008b) that we also observe here in the simulations. However, beamformers are especially sensitive to the accuracy of the head model; measured data that is inconsistent with the head model is liable to be rejected as noise (Steinsträter et al., 2010). Additionally, the beamformer approach assumes that reconstructed sources are temporally uncorrelated with each other. In case of spatially distant sources that are correlated with each other and coherent with peripheral oscillations, the algorithm may therefore yield erroneous localizations (Hadjipapas et al., 2005; Sekihara et al., 2002; Van Veen et al., 1997). In continuous recordings of real data, such correlations may occur and seem to outweigh the technical advantages of MVBF for the localization of coherent functional networks. Yet, the performance differences observed in real data must be interpreted with caution, because we do not know the true location of strongest cortico-peripheral coherence in real datasets. For instance, there is evidence not only for involvement of the primary motor cortex in corticomyocellular coherence but also of somatosensory and parietal areas (Witham et al., 2010; Baker, 2007). Premotor areas could also be involved. The peak of corticomyocellular coherence could therefore be shifted away from the hand notch and lead to biased performance estimates of the inverse solutions. The peak coherence location estimated by the MVBF was on average anterior to the hand notch and to the localizations of MN. Hence, if the true corticomyocellular coherence peak was anterior to the hand notch, this would reduce the inferior performance of MVBF compared to MN we observed. In any case, valid localizations of motor and visual coherent sources could be achieved in all participants also with MVBF, although the distance to the anatomical target used here was ~10 mm larger than with MN.

The spatial spread of FCMs has similar characteristics as the spatial spread of activation maps and follows the point-spread function of each algorithm (Sekihara et al., 2005). The MVBF provides significantly more focal localizations than MN both in simulations and real data.

Although to our knowledge no systematic comparison of adaptive vs. non-adaptive algorithms has been performed for FCMs from MEG data so far, it is probable that the differences found with EEG data in this study also apply to MEG data.

4.2. EEG-specific parameters

The parameters that are specific for EEG studies turned out to be of relatively minor importance for localization accuracy of coherence. We did not observe significant differences in performance between the different head models, and valid results could even be obtained without a co-registered individual structural MRI with head models based on MRI templates. However, more precise head models (SMAC and BEM) yielded better localization accuracy, and should therefore be preferred. The motor and visual cortex that were used as targets in this study are relatively superficial; the accuracy of the head model is probably more critical for the localization of deeper networks, e.g., in the hippocampus. Furthermore, studies using a larger number of participants and a smaller solution grid size will probably find significant performance differences even for superficial sources. Future studies could address the effect of different head models on coherence localization with simulations, e.g., by creating simulations with a finite element model (FEM) and by evaluating the performance of other head models such as the ones used in this manuscript.

Valid localizations of coherent cortical sources were not only obtained in healthy participants but also in all patients with ischemic brain lesions. Hence, it is feasible to use the EEG to map functional connectivity also in patients with brain lesions. However, our study was not designed to make a systematic comparison between patients and healthy controls, and does therefore not rule out a performance decrease in patients, especially in areas close to the lesions.

In sharp contrast with functional connectivity analyses between EEG sensors where the choice of the reference is crucial (Fein et al., 1988; Guevara et al., 2005), the reference montage has no significant influence on localization accuracy of source space connectivity maps, since the inverse solutions make the data reference-free (Fender, 1987; Geselowitz, 1998). However, in accordance with a previous report (Van Veen et al., 1997), we found non-vanishing differences between average- and vertex-referenced montages when using a beamformer. This may be initially surprising since the coordinate transform introduced by the reference is applied equally to the lead-potential and the data and therefore should be canceled. However, the observed differences arise for 2 reasons. First, the inversion of the sensor covariance matrix R introduces minor variances leading to reference dependent variations in the localizations. This can be avoided by using the pseudoinverse of R (or other constant regularization across the different reference choices), even when the matrix is well-conditioned, which yields reference-independent results for vector beamformers. Second, equation (1) used for determination of the dipole orientation with maximum SNR for the MVBF is not neutral to a coordinate transform introduced by the EEG reference and therefore explains the non-vanishing differences observed here with a scalar beamformer. Nevertheless, the differences are minor and did not reach significance in our datasets (p ≥ 0.33). By the way, the MN algorithm for calculation of dipole orientation used here (equation 2) is reference-independent.

4.3. Dipole orientation

Concerning dipole orientations, our results show that output maximization algorithms yield similar FCM performance as anatomically defined surface normal orientations, while being more convenient and requiring much
less human work load. This is in agreement with a previous MEG study which showed that surface normal dipole orientations are only advantageous if the MEG-MRI coregistration error is smaller than 2 mm and the error in the estimation of the cortical surface orientation is smaller than 10° (Hillebrand and Barnes, 2003). Such output maximization algorithms are valid not only for beamformers but also for non-adaptive algorithms. We can, however, not rule out minor performance differences which might have been masked due to the relatively low number of participants in this study. Future studies could use simulations of different dipole orientations to compare more systematically the performance of the two procedures for orientation determination.

4.4. Measures of functional connectivity

In this paper, we compared the traditional measure of MSC to the more recently introduced variant IC. Unlike MSC, IC ignores spectral similarities among 2 signals that occur with zero time delay (Nolte et al., 2004). This has the advantage that it is robust to volume conduction and crosstalk among voxels arising from spatial leakage of the inverse solutions, since these artificial similarities occur with zero time-delay and are therefore removed. IC can therefore not only be used for the localization of cortico-peripheral connectivity as done in this study, but also for the localization of cortico-cortical interactions (Guggisberg et al., 2008a). On the other hand, it ignores a portion of true interactions and could therefore lead to less reliable localizations of interactions than MSC. Yet, cortico-peripheral coherence is presumed to never occur at zero lag because of the time needed for neural transmission. Indeed, we observed a similar localization accuracy for IC as for MSC in our analyses of cortico-peripheral connectivity.

Many other methods for calculating cortical functional connectivity are available instead of coherence applied here (see e.g. Lachaux et al., 1999; Le Van Quyen et al., 2001; Stam et al., 2003, 2007). While some of them have been compared head-to-head for analyses on the sensor level (Dauwels et al., 2010; Guevara et al., 2005; Le Van Quyen et al., 2001), future studies will need to address performance differences for source connectivity analyses.

5. Conclusions

We show that coherent functional sources can be accurately localized and studied with high-resolution EEG systems. The best performance is achieved with precise models of the individual head shape and with inverse solutions that are not susceptible to interference from spatially distant coherent sources.

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