Extraction of Single-Trial Post-Movement MEG Beta Synchronization in Normal and Parkinson's Patient Using ICA-Based Spatiotemporal Approach

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1. Introduction

The human brain is a dynamic system that frequently changes functional mode (Lopes da Silva, 1991; Lopes da Silva, 1996). Spatiotemporal analysis of brain activities with regard to distinct spatial locations and frequency bands reveals task-specific brain activation which changes in a fraction of a second (Jensen & Vanni, 2002). At rest, Rolandic EEG and MEG rhythms are dominated by rhythmic activity around 10 (alpha band) and 20 (beta band) Hz. Electrocorticographic (Pfurtscheller et al., 1994) and neuromagnetic recordings have shown that the ~20-Hz rhythm mainly originates in the anterior bank of the central sulcus while the ~10-Hz rhythm is concentrated predominantly in the post-central cortex (Pfurtscheller & Lopes da Silva, 1999). These two frequency components appear to have different functional roles, with the ~20-Hz rhythm being more closely connected to movements and their termination and the ~10-Hz component behaving more like a classical “idling” rhythm (Salmelin et al., 1995). Voluntary movement is composed of three phases: planning, execution and recovery (Pfurtscheller et al., 1998a). It has been suggested that localized event-related alpha desynchronization (ERD) upon movement can be viewed as an EEG/MEG correlate of an activated cortical sensorimotor network, servicing planning and execution, while beta event-related synchronization (ERS) may reflect deactivation/inhibition during the recovery phase in the underlying cortical network (Pfurtscheller et al., 1996).

Movement-related ERD and ERS have been used as probes to study neurophysiology in normal brains and pathophysiology in the diseased (Tamas et al., 2003). It has been reported that the diagnostic features of patients with Parkinson’s disease, in comparison with controls, are a slowing and suppression of the post-movement beta ERS independent of the amount of beta activity in the reference period (Pfurtscheller et al., 1998a). These findings imply that slowed and reduced recovery after the motor act impedes cortical preparation of the next movement (Pfurtscheller et al., 1996). Patients with Unverricht-Lundborg type myoclonic epilepsy demonstrate little rebound of beta activities contingent upon median nerve stimulation (Silen et al., 2000). The diminished beta ERS indicates that the myoclonic
patients have sustained motor cortex reactivity which can be attributed to impaired cortical inhibition (Pfurtscheller & Lopes da Silva, 1999).

ERD and ERS activities are time-locked, but not phase-locked, to external stimuli or tasks (Andrew & Pfurtscheller, 1995; Kalcher & Pfurtscheller, 1995; Pfurtscheller & Lopes da Silva, 1999). Existing methods for extraction of ERD/ERS signals essentially measure power or amplitude changes of corresponding frequency bands as derived from the average of dozens or hundreds of trials. The band power method squares and averages filtered brain signals within a selected frequency band (Pfurtscheller & Aranibar, 1977), and an inter-trial variance method to remove the phase-locked portion in the band power method was reported by Klimesch et al. (1998). Likewise, autoregressive and spectral decomposition methods have been used to extract significant frequency components in rhythmic signals (Florian & Pfurtscheller, 1995). Salmelin's temporal-spectral evolution method rectifies and averages filtered MEG signals (Salmelin et al., 1995). To increase the temporal resolution of the ERD/ERS technique, Clochon et al. (1996) proposed an amplitude modulation (AM) method based on the Hilbert transform to detect the envelope of filtered signals by squaring and summing their real and imaginary parts. All these approaches presume stereotypical frequency and temporal characteristics across trials and require an average of many trials for the ERD/ERS using a preset frequency filter and time window to preprocess every trial. However, non-phase-locked rhythmic signals can vary from trial-to-trial contingent upon variations in a subject’s performance and state, which may be linked to fluctuations in expectation, attention, arousal, and task strategy (Bastiaansen et al., 2001; Bastiaansen et al., 1999; Earle, 1988; Haig et al., 1995; Hoffman et al., 1991; Yabe et al., 1993). Since trial-to-trial variability in amplitude, latencies, or scalp distribution might carry important information on cognitive and physiological states (Jung et al., 2001), a method that permits the extraction and analysis of the oscillatory signal on a single-trial base is crucial for the study of subtle brain dynamics. Furthermore, such a method should require fewer trials for analysis and hence shorter experiment time, which is beneficial for patients with impairment of motor and/or cognitive performance (Muller-Gerking et al., 1999).

Single-trial multi-channel EEG analysis has been developed for time-locked, phase-locked, evoked brain activities (Jung et al., 2001; Tang et al., 2002). However, approaches to single-trial movement-related oscillatory changes are less explored. Independent component analysis (ICA), a data-driven method for multivariate data analysis, has been used to reveal temporally-independent neuronal activities of EEG measurements (Jung et al., 2001; Makeig et al, 1997; McKeown et al., 1998), MEG measurements (Wu et al., 2002; Wu et al., 2003; Tang et al., 2002), fMRI (Duann et al., 2002; McKeown et al., 1998) and recently perfusion MRI (Kao et al., 2003). The present study proposes a new approach using ICA and the Hilbert transformation for the single-trial detection of movement-related beta rhythmic activity during a self-paced right finger lifting task. This study focuses on beta activity and beta ERS, centered around 20 Hz, because it has been demonstrated that the movement-related short bursts of beta oscillation have higher task and movement specificity than alpha ERD (Pfurtscheller & Aranibar, 1979b; Pfurtscheller et al., 1996).

Since brain oscillation may be expressed alone in a specific frequency band independent of artifacts (Ermer et al., 2000; Lins et al., 1993a; Lins et al., 1993b; Mosher et al., 1992), ICA is applied to transform brain signals across all channels (in a single trial) into mutually independent components by means of an unmixing matrix in which each column represents a spatial map tailoring the weights of the corresponding temporal component at each MEG sensor. The spatial maps and temporal waveforms of decomposed independent components
are categorized into task-related and task-unrelated groups respectively, based on temporal and spatial characteristics. This temporal template is the grand average of hundreds of vector-norm envelopes of the band-pass filtered, single-trial MEG measurements obtained from right index finger lifting. The spatial template can be derived from the spatial distribution at beta rebound activity either from the grand average of the generation group (for signal extraction) or from each individual (for verification). Correlations between the temporal template and component waveforms, as well as between the spatial template and spatial maps, are computed, and coupled component waveforms and spatial maps that conjointly survive with high correlation values are taken as task-related information and subjected to data reconstruction. In this way the phase and amplitude information of noise-free MEG beta activities can be preserved for profound studies of temporal and spectral variation across trials. Due to the high signal-to-noise ratio (SNR) in beta activities extracted through ICA, trial-specific reactive frequency ranges can be determined by means of the comparisons of two short time spectra between the reference and post-movement periods. Beta reactivity per single trial can be quantified using the amplitude modulation (AM) method (Clochon et al., 1996), and insignificant epochs can be determined using a nonparametric sign test (Brovelli et al., 2002). Source estimation and localization techniques can be successfully applied to single-trial epoch to estimate the source locations of beta modulation.

The current study presents: 1) a novel ICA-based spatiotemporal approach for single-trial analysis of event-related beta oscillatory modulations with a high extraction rate; 2) the prospect of trial-specific frequency bandpass filtering that takes into account subtle trial-by-trial brain dynamics; 3) the feasibility of using sophisticated source estimation/localization methods demanding high signal-to-noise ratio (SNR) on single trial data; and 4) a common template approach permitting an effective alternative in cases where lengthy procedures cannot be endured by participants or in clinical settings where patients have attention problems or are incapable of sustaining long experiments. The proposed ICA-based approach was applied to discover the mechanisms of beta ERS in one Parkinson’s patient. It is helpful to investigate the reasoning of ERS vanishment due to suppression of post-movement beta rebound in each single-trial, rather than the cause of temporal jittering and/or loss of synchronization in Parkinson’s disease.

2. Materials and methods

2.1 Subjects and task

The present study examined six healthy right-handed subjects (gender balanced), aged 24-30 years. Five of the healthy subjects were used in the model generation group, and MEG data from the last healthy subject were used for validation. Subjects performed self-paced lifting of the right index finger approximately once every 8 sec. Subjects were trained to perform the movement briskly for a duration of 200 to 300 ms, as monitored by surface electromyogram (EMG) on extensor digitorum communis, with a range of finger movement around 35~40°, while keeping their eyes open in order to suppress the occipital alpha rhythm. In addition, somatosensory evoked fields (SEFs) for right median nerve stimulation were measured to locate the primary sensorimotor area (SMI) in each subject as part of the procedure for the generation of a temporal template (see below). Informed written consent was obtained from all subjects. This study was approved by the Institutional Review Board of Taipei Veterans General Hospital. In addition, one 56-year-old patient with idiopathic Parkinson’s disease in Hoehn and Yahr stage 1 was also recruited as a demonstration in this study.
2.2 Data recording

Cortical magnetic signals were recorded with a 306-channel (102 sensor unit) whole-head neuromagnetometer (band-pass, 0.05-250 Hz; digitized at 1kHz; Vectorview; Neuromag Ltd., Helsinki, Finland) with subjects in sitting position. Each sensor unit was composed of a pair of planar gradiometers and a magnetometer. The magnetometer measured magnetic flux \( B_z \), normal to the sensor unit, while the gradiometers measured two tangential derivatives of \( B_z \) (\( \partial B_z / \partial x \) and \( \partial B_z / \partial y \), mutually orthogonal). Only magnetic signals measured by the gradiometers were used in this study. Bipolar horizontal and vertical electro-oculograms (EOG) were recorded using electrodes placed below and above the left eye and at the bilateral outer canthi to monitor eye movement and blinks. The exact position of the head with respect to the sensor array was determined by measuring magnetic signals from four head position indicator (HPI) coils placed on the scalp. Coil positions were identified with a three-dimensional digitizer with respect to three predetermined landmarks (nasion and bilateral preauricular points) on the scalp, and this data used to superimpose MEG source signals on individual MRI images obtained with a 3.0 T Bruker MedSpec 5300 system (Bruker, Kalsrube, Germany). The anatomical image was acquired using a high-resolution T1-weighted, 3D gradient-echo pulse sequence (MDEFT: Modified Driven Equilibrium Fourier Transform; TR/TE/TI= 88.1ms/4.12ms/650ms, 128*128*128 matrix, FOV=250mm).

Empty room measurements were recorded for 3 minutes. Approximately 100 EOG-free trials of right index finger lifting were acquired and analyzed off-line. Since the focus was on beta-activities, the signals were further band-pass-filtered between 6-50 Hz (zero-phase, tenth-order, IIR Butterworth filter) to remove dc drifts and 60 Hz noise. The initial finger movement (movement onset; zero time) was registered with an optical switch (Taniguchi et al., 2000). Electromyographic (EMG) activity from the extensor digitorum communis (digitized at 1 KHz) was continuously recorded to monitor performance (see above). Each epoch comprised data points from -4s to 3s relative to the movement onset (Salmelin et al., 1995; Salmelin and Hari, 1994a) and epochs were subjected to further single-trial ICA analysis.

For SEF measurement, the right median nerve was electrically stimulated every 2 sec with constant current pulses (0.3 msec in duration) exceeding the motor threshold. Approximately 100 EOG-free trials were acquired and digitized at 1 kHz for off-line analysis.

2.3 Data analysis

2.3.1 Independent Component Analysis of the single-trial MEG epoch

We take the advantages of sensitivity and localizing power of superficial sources by planar gradiometers (Rosell et al., 2001; Kajola et al., 1991). Each single-trial MEG epoch contains m channels (m = 204, 102 pairs of gradiometers) and n time points (usually m < n). The paired gradiometer signals (\( \partial B_z / \partial x \) and \( \partial B_z / \partial y \)) are arranged into two \( \frac{m}{2} \times n \) sub-matrices \( B_1 \) and \( B_2 \) and concatenated into an \( m \times n \) matrix \( B \). The \( i \)th rows (\( i \leq 102 \)) of \( B_1 \) and \( B_2 \) contain the measured gradiometer signals from the \( i \)th sensor location, and the \( j \)th column in \( B \) contains the measured data at the \( j \)th time point across all gradiometer channels.
Mathematically, we can consider each row of $\mathbf{B}$ as samples generated from one random variable $b_i$, $i = 1, 2, \ldots, m$. In other words, matrix $\mathbf{B}$ is a realization of a random vector $b = [b_1 \ b_2 \ \cdots \ b_m]^T$.

The ICA techniques (Jung et al., 2001; Hyvarinen et al., 2001) seek to find a $p \times m$ ($p \leq m$) matrix, $\mathbf{W}$, which converts the random vector $b$ into another vector variable, $s$, consisting of $p$ mutually independent random variables, thus:

$$
\mathbf{s} = \begin{bmatrix}
    s_1 \\
    s_2 \\
    \vdots \\
    s_p
\end{bmatrix} = \mathbf{W} \mathbf{b}
$$

(1)

The mutual independence of $s_i$, for $i = 1, \ldots, p$, implies that if $P(s_i)$ represents the probability distribution of the $i^{th}$ component, the joint probability distribution for all components can be factorized as:

$$
P(s_1, s_2, \ldots, s_p) = P(s_1)P(s_2)\ldots P(s_p)
$$

(2)

The ICA techniques use this assumption of mutual independence to find the un-mixing matrix $\mathbf{W}$.

All calculations in the present study were carried out using the FastICA algorithm which features high speed calculation (cubic convergence) and does not require selection of step size parameters or learning rate, unlike the gradient-based algorithm (Hyvarinen et al., 1997, 2001). The FastICA technique first removes means of row vectors in the $\mathbf{B}$ sample matrix such that each random variable $b_i$ has a zero mean, and then employs a whitening process using principal component analysis. After whitening, the covariance matrix of the whitened data becomes an identity matrix, and only the first $p$ ($p \leq m$) most significant principal components are preserved in the FastICA calculation.

The next step is to look for a matrix that transforms the whitened data into a set of components as mutually independent as possible. Mutual information, as a measure of the independence of random variables, is used as the criterion for finding such a transformation. Mutual information can be expressed in terms of negentropy, an important measure of non-Gaussianity (Hyvarinen et al., 1997, 2001). Therefore, the problem of finding the independent components ($s$) and the transform matrix ($\mathbf{W}$) can be translated into a search for linear combinations of the whitened data that maximize the negentropy of the distributions of $s_i$, for $i = 1, \ldots, p$.

After applying FastICA to the pre-processed single-trial MEG epochs, matrix $\mathbf{B}$ can be factored into a (mixing) matrix $\mathbf{U}$ and an (independent source) matrix $\mathbf{S}$ as follows:
\[ B_{mn} = \begin{bmatrix} B_1 \\ B_2 \end{bmatrix} = U_{mp}S_{pn} \]

in which each row \( s_i \) of matrix \( S \in \mathbb{R}^{p \times n} \) represents samples of an independent component (IC) \( s_i \) for \( i = 1, \ldots, p \) and \( U \in \mathbb{R}^{mp \times p} \) is the pseudo-inverse of matrix \( W \) whose column vectors represent the weight distribution values of the corresponding ICs in \( S \) across all MEG gradiometer channels. In fact, matrix \( U \) is the “mixing matrix” that combines the \( p \) ICs to reconstruct signal \( B \). These temporal ICs can be categorized into task-related ICs and task-unrelated ICs. Since the elicited brain activities or artifacts can be distributed over multiple ICs, no one-to-one correspondence between IC and source information is projected (Makeig et al., 1997). To facilitate the selection of task-related ICs, a temporal and spatial template pair was constructed prior to selection (see below). Spatial map \( \tilde{x}_j \) of the \( j^{th} \) IC was defined as the topographic display of all vector norms for weights of 102 gradiometer pairs in the \( j^{th} \) column vector of \( U \),

\[ \tilde{x}_j = \begin{bmatrix} u_{1,j}^2 + u_{m,1,j}^2 \\ u_{2,j}^2 + u_{m,2,j}^2 \\ \vdots \\ u_{m,j}^2 + u_{m,m,j}^2 \end{bmatrix}^T \]

in which \( u_{i,j} \) is the entry in the \( i^{th} \) row and \( j^{th} \) column of \( U \) in Eq. (3). The spatial map is intended for component selection (see below).

### 2.3.2 Creation of a temporal template (VAMW-template) using amplitude modulation (envelope) of the MEG data

The recorded MEG signals at each gradiometer are filtered in the task-specific frequency band (Pfurtscheller & Lopes da Silva, 1999) and rectified by computing the AM waveform (envelope) using the amplitude modulation (AM) method (Clochon et al., 1996) as follows:

\[ m(t) = \sqrt{M_{BP}(t)^2 + H(M_{BP}(t))^2} \]

in which \( M_{BP}(t) \) is the band-passed MEG signal and \( H(M_{BP}(t)) \) is its Hilbert transform. The task-specific frequency band is determined by the contrast between two 1-s amplitude spectra calculated over about one hundred event-related EEG trials (Pfurtscheller and Lopes da Silva, 1999). One (serving as rest reference) is computed over the duration from 4s to 3s.
preceding the onset of movement, and the other (serving as reactive target) from 0.8s to 1.8s after the onset of movement (see Fig. 1a, b). All beta-frequency components with significant modulation in terms of post-movement amplitude increase (above 95% confidence level, i.e. $Z>3.09$, $P<0.01$) in the differential amplitude spectrum (see Fig. 1c) are taken as the task-specific frequency band for subsequent processing (Pfurtscheller G. and Berghold A., 1989).

Fig. 1. Determination of task-specific frequency band using two 1-s amplitude spectra. (a) “R” represents the reference period from -4s to -3s preceding onset of movement and “P” represents the post-movement duration from 0.8s to 1.8s after onset of movement. (b) Two spectra computed over the reference (R) and post-movement periods (P), respectively. (c) The task-specific frequency band for beta-band VAMW is defined as the one where the difference between two spectra exceeds the 95% confidence level.

The vector norm of AM waveforms (VAMW) at each sensor site is computed using the square root of the AM waveforms of each gradiometer pair, i.e.,

$$V(i,t) = \sqrt{m_x(i,t)^2 + m_y(i,t)^2},$$

in which $V(i,t)$ is the VAMW at the $i^{th}$ sensor location, and $m_x(i,t)$ and $m_y(i,t)$ are the AM waveforms in $\frac{\partial B_z}{\partial x}$ and $\frac{\partial B_z}{\partial y}$ directions of the $i^{th}$ sensor location. Event-related beta modulation is then computed as the difference in amplitude between the maximum amplitude of VAMW for each sensor site in the post-movement (0.8s to 1.8s) interval and mean activity between -2.5s and -2s (see Fig. 2a) (Leocani et al., 1997). Beta rebound (BR) is defined as the maximum amplitude of the computed event-related beta modulation from the subset of nine sensor sites in the vicinity of SMI (identified by SEF). The VAMWs of the BR calculation were averaged across the subjects (500 trials, 100 trials for each subject, 5 subjects pooled) to create the common temporal template, designated VAMW$_{template}$ (Fig. 2a).
2.3.3 Creation of a spatial template using topographical distribution of event-related beta modulation values

Individual spatial templates were first generated from the topographical distributions of event-related beta modulation values (see above). The five templates from the model generation group were then averaged to generate a common spatial template. In order to optimize conditions for spatial averaging, subjects’ heads were carefully positioned before actual measurements to keep head positioning and orientation as similar as possible. Distances between head centers of the five subjects and the reference point (the origin of the MEG sensor array) in the horizontal plane were less than 4mm, and angles between the vertical axis of the helmet and that of the head (the normal vector of the plane constituted by the three landmark points, i.e., nasion, and both pre-auricular points) remained within 5.5° (maximum deviation 1.5°) between subjects.

Only the left half of the spatial map (unshaded in Fig. 2b) was used as the spatial template because this study focused on beta event-related activities in the hemisphere contralateral to the side of finger lifting; however, the other half can be generated analogously to extract activities in the ipsilateral hemisphere. Correlations among individual spatial templates ranged from 0.92 to 0.68. Respective correlations between the common spatial template and the individual spatial templates were 0.973, 0.811, 0.881, 0.904, and 0.915. These high correlation values support the use of the spatial template in component selection for each individual’s magnetic signals.

2.3.4 Selection of pertinent independent components for the reconstruction of reactive beta activities

A spatial map (Eq. (4)) and corresponding VAMWs of each IC were generated for the selection of task-related ICs. Since the original signals may be decomposed into multiple ICs, the spectrum of each IC may vary from the one in the original signal due to the decomposition process. When settings for band-pass filtering for VAMW computation cannot be optimally determined using two-spectrum comparison for the generation of a VAMW template (Pfurtscheller & Lopes da Silva, 1999), three standard beta bands, 12-16, 16-20 and 20-24 Hz (Pfurtscheller G., 1981), enclosing the event-related beta activities in motor task, were used to band-pass filter (zero-phase, tenth-order, IIR Butterworth filter) for each single-trial IC such that the three frequency-laden resultant VAMW_{IC} (the VAMWs band-pass filtered in three frequency bands of each IC) retained all task-related information. These VAMW_{IC} were subsequently used in the selection of task-related ICs, which must fulfill the following dual criteria: 1) at least one of three corresponding VAMW_{IC} has a correlation with the VAMW_{template} higher than 95% (Z>1.63, P<0.05) among VAMW_{IC} of all the ICs for that single epoch, and 2) correlation between the spatial map and spatial template is above 95% (Z>1.63, P<0.05) for the spatial maps of all ICs. Data processed via 3-standard band filtering are not used in subsequent data reconstruction, but rather are used in conjunction with the dual-criteria only in the procedure “selecting” the pertinent ICs. Unselected columns, i.e., task-unrelated components, of mixing matrix U (Eq. (3)) are zeroed to produce a matrix \( \hat{U} \) such that task-related rhythmic signals are reconstructed by multiplying \( \hat{U} \) and \( S \) (Fig. 3). The reconstructed data in each trial are then filtered within a trial-specific frequency band to extract reactive beta activities.
Fig. 2. Creation of common temporal and spatial templates. (a) The common temporal template, $VAMW_{\text{template}}$, is created by averaging VAMWs (500 trials, 100 trials for each subject, 5 subjects pooled). Event-related beta modulation is defined as the amplitude difference between the mean amplitude of baseline activity (-2.5 to -2 s) and maximum amplitude in the post-movement interval (0.8 to 1.8 s). (b) The common spatial template is the average of the topographical distributions of event-related beta modulations of five subjects from model generation group. Only the half the spatial map (unshaded) contralateral to the side of finger lifting is used as the spatial template.
Fig. 3. Examples of IC-selection and signal reconstruction procedure. (a) Spatial maps, IC waveforms, Fourier spectra of IC waveforms and VAMW ICs of five ICs obtained from one
single epoch by ICA. Only ICs fulfilling the dual criteria are selected for signal reconstruction. For example, IC 3 meets the dual criteria (underscored in red): i) correlation value between spatial map and spatial template is 0.84 (rank= 97%, \(Z = 1.89\), \(P=0.03\)); ii) correlation value between 16-20 Hz VAMW_{IC} and 20-24 Hz VAMW_{IC} with VAMW_{template} is 0.8 (rank= 99%, \(Z=3.08\), \(P=0.01\)) and 0.78 (rank= 97.8%, \(Z = 2.85\), \(P=0.022\)), respectively. (b) Noise identification and removal. The deselected IC 2 in Fig. 2a may emanate from background noise since it resembles the IC 1 extracted from empty room measurement. (c) The impact of including task-unrelated IC into signal reconstruction. This figure illustrates (a different trial from Fig. 3a) that inclusion of task-unrelated IC (IC 9) with a high spatial correlation (correlation value=0.61, rank= 95.2%, \(Z=1.67\), \(P=0.048\)) but poor temporal correlation (correlation value=0.28, rank=13%, \(Z=0.34\), \(P=0.87\)) causes deterioration in the beta BR from 28.9 fT/cm (arrows and trace in red; IC9 eliminated from reconstruction) to 18.6 fT/cm (arrows and trace in blue; IC9 included for reconstruction).

2.3.5 Detection of task-laden trial-specific frequency band and extraction of reactive beta activities

The trial-specific frequency band detected in each trial is used to confine the reconstructed data within the most reactive beta band for further BR computation and source estimation. This frequency band is defined by the reactive beta band of the sensor site showing highest event-related beta modulation value (see creation of temporal template) over the nine SMI vicinal sensor sites (identified by SEF) and is identified using the aforementioned two-spectrum procedure which has been suggested as the best approach for the determination of reactive frequencies (Pfurtscheller & Lopes da Silva, 1999). Following data filtering with a trial-specific frequency band (zero-phase, tenth-order, IIR Butterworth filter), reactive beta activities in each single epoch can be extracted. The extracted reactive beta activities are then subjected to source estimation and beta rebound (BR) computations.

2.3.6 Calculation of VAMW_{recon} of reactive beta activities and single-trial epoch selection using a nonparametric sign test

Movement-related beta rebound (BR) can be quantified from single-epoch reactive beta activities and VAMW_{recon} (VAMW of reconstructed data) for reactive beta activity at each sensor site computed. The VAMW_{recon} of highest event-related beta modulation (see creation of temporal template) among the nine sensor sites vicinal to SMI is designated as VAMW_{recon_max} and is used in turn for single-trial epoch selection and BR computation, as the sensor site expressing VAMW_{recon_max} did not change throughout the experiment in our observations. A deterministic procedure, modified from Brovelli et al. (2002) approach, is used to select the significant trial. A nonparametric sign test is applied to the VAMW_{recon_max} designated for BR calculation in each single trial by computing the Z-score at each time point as

\[
Z(t) = \left( N^+(t) - \frac{1}{2} N \right) / \left( \frac{1}{2} \sqrt{N} \right),
\]

in which \(N^+(t)\) denotes the number of trials whose magnitudes are larger than the median value of their baseline activities at time point \(t\), and \(N\) the total number of trials. Time points with \(Z\) values greater than 3.09 (\(P<0.01\)) are defined as the time interval-of-interest (IOI). After the determination of IOI for each subject, another sign test is then applied to find epochs showing significant increases in amplitude (\(Z>1.63, P<0.05\)) using

\[
Z_{IOI}(i) = (N_{IOI}^+(i) - 1)(N_{IOI}^-) / \left( \frac{1}{2} \sqrt{N_{IOI}^-} \right),
\]

in which \(Z_{IOI}(i)\) is the Z value of the
\(i^{th}\) trial, \(N_{IOI}^{-}(i)\) is the number of data points in post-movement IOI with values larger than the median of baseline activities of the \(i^{th}\) trial, and \(N_{IOI}^{+}\) is the total number of time points in post-movement IOI (Brovelli et al., 2002). An example of single-trial epoch selection is given in Fig. 4 (Subject I). The first trial in Fig. 4 with a \(Z_{IOI}\) score equal to -4.53 is marked as an insignificant epoch and eliminated from further analysis.

### 2.3.7 Source estimation of the reactive beta activities

Source estimation of the MEG reactive beta activities was done using equivalent current dipole (ECD) analysis and minimum current estimation (MCE, Uutela et al., 1999; toolbox provided by Neuromag Ltd, Helsinki, Finland). A single dipole model was applied to explain the field every 1ms, and only dipoles showing goodness-of-fit (Jensen and Vanni, 2002) values higher than 80% were used for data explanation. In MCE, the lattice constant of the triangular grid was 10mm and locations closer than 30mm to the center of the conductor were excluded from current estimates. Both analyses used a realistic head model for each subject. Template generation and single-trial data processing procedure are schematized in Figs. 5a and 5b respectively. Epochs achieving significance in the increase of beta activities were chosen for subsequent BR calculation and dipole/source analysis.

\[\begin{align*}
N_{IOI}^{+} &= 336, \\
N_{IOI}^{-} &= 126, \\
Z_{IOI} &= -4.53
\end{align*}\]

\[\begin{align*}
N_{IOI}^{+} &= 336, \\
N_{IOI}^{-} &= 336, \\
Z_{IOI} &= 18.41
\end{align*}\]

\[\begin{align*}
N_{IOI}^{+} &= 331, \\
N_{IOI}^{-} &= 331, \\
Z_{IOI} &= 17.86
\end{align*}\]

Fig. 4. Example of single-trial epoch selection based on a nonparametric sign test. Single-trial VAMW\textsubscript{recon, max} of reconstructed data are examined through a nonparametric sign test. \(Z_{IOI}(i)\) is the Z value of the \(i^{th}\) trial, \(N_{IOI}^{+}(i)\) is the number of data points in post-movement IOI with values larger than the median of baseline activities of the \(i^{th}\) trial, and \(N_{IOI}^{-}\) is the total number of time points in post-movement IOI. Only epochs showing significant increase of beta activities are chosen for further analysis. The first trial with a \(Z_{IOI}\) score equal to -4.53 is marked as an insignificant epoch and eliminated from further analysis.

### 2.3.8 Validation of coupled common spatial and temporal templates for single-trial analysis

Since there are inevitably differences in head size and variations in head positions inside the MEG scanner among subjects, BR amplitude differences were compared using both individual spatial templates and the common spatial template. The use of a pair of common
spatial and temporal templates for the extraction of individuals’ neuromagnetic single-trial signals was further validated on one additional subject.

3. Results

Based on the known spatial location and temporal expression in terms of spatial and temporal templates, reactive beta activities were successfully extracted. Figure 3a shows that IC 3 meets the dual criteria: i) the correlation values between spatial map and spatial template is 0.84 (rank= 97%, Z= 1.89, P=0.03); ii) correlation values of 16-20Hz VAMW\textsubscript{IC} and 20-24 Hz VAMW\textsubscript{IC} vs. VAMW\textsubscript{template} are 0.8 (rank= 99%, Z=3.08, P=0.01) and 0.78 (rank= 97.8%, Z= 2.85, P=0.022), respectively. Fig. 3a illustrates that noise could also be identified and removed. IC2 in Fig. 3a correlates highly (=0.88) in spatial distribution with the IC1 extracted from empty room measurements (Fig. 3b), and is therefore rejected.

(a)

Measured MEG data

Determine the task-specific frequency band by comparing two short-time spectra. One spectrum is computed over reference time interval (-4s – -3s) and the other is computed over post-movement time interval (0.8s – 1.8s).

Filter the MEG epochs measured from each gradiometer with the task-specific frequency band and rectify the results using amplitude modulation method to obtain the AM waveforms: 

\[ m(t) = \sqrt{M_{\text{SP}}(t)^2 + H(M_{\text{SP}}(t))^2} \]

Compute the vector norm (VAMW) on each sensor site with a pair of gradiometers: 

\[ V(i,t) = \sqrt{m_x(i,t)^2 + m_y(i,t)^2} \]

Step 1: Average the VAMWs over many trials for each subject.
Step 2: Compute the event-related beta modulation of each sensor site, which is the difference between the maximum amplitude of VAMW in the post-movement (0.8s to 1.8s) interval and the mean amplitude of VAMW during -2.5s to -2s.

Find the sensor site displaying maximum event-related modulation in each of five subjects in model generation group, and average them to create the common temporal template, VAMW\textsubscript{template}.

For each subject in model generation group, construct an individual spatial template which is defined as the topographical distribution of event-related beta modulation of averaged VAMWs.

The common spatial template is created by averaging the individual spatial templates in model generation group.

Fig. 5a. Flow chart for creation of common spatial and temporal templates.
**Input MEG epoch**

\[ B_{mxn} \text{ with } m \text{ channels and } n \text{ time points} \]

**Preprocessing:** Bandpass filtering (6 Hz–50 Hz)

**Select IC component for data reconstruction:**

- **Step 1:** Compute the ICs using FastICA: \[ B = U \cdot S \]
- **Step 2:** Compute VAMWs of each IC with three standard beta bands, i.e. 12–16 Hz, 16–20 Hz, and 20–24 Hz, denoted as \( \text{VAMW}_{\text{betas}} \). The VAMW\(_{\text{betas}}\)s are subsequently in conjunction with spatial map for task-related IC selection.
- **Step 3:** Select ICs for data reconstruction. Dual criteria should be fulfilled: 1) the correlation between the temporal template and at least one of three VAMW\(_{\text{betas}}\)s should be higher than 95\% (\( Z > 1.63, P < 0.05 \)) among all ICs and 2) correlation between the spatial template and spatial map also should be higher than 95\% (\( Z > 1.63, P < 0.05 \)) among all ICs.

**Single-trial epoch selection:**

- Apply the nonparametric sign test for VAMW\(_{\text{recon max}}\) to detect the increase of beta activity emerging from baseline activity. Epochs with significant increase (\( Z > 1.63, P < 0.05 \)) in post-movement beta activity are selected for current source estimation and BR amplitude quantification.

---

**Reconstructed data**

**Determine the trial-specific frequency band:**

- **Step 1:** Determine the reactive frequency bands at the nine sensor sites in the vicinity of SMI using the two-spectrum comparison method.
- **Step 2:** Filter the reconstructed data of the nine SMI vicinal sensor sites using the reactive frequency bands and compute their VAMWs.
- **Step 3:** For each of nine sensor sites, compute the event-related beta modulation which is the difference between the maximum amplitude of VAMW in the post-movement (0.8s to 1.8s) interval and the mean amplitude of VAMW during -2.5s to -2s.
- **Step 4:** Find the maximum event-related beta modulation among nine sensor sites and designate its reactive frequency band as the trial-specific frequency band.

---

**Good epoch?**

- **Yes**
  1. Compute BR of the VAMW\(_{\text{recon max}}\).
  2. Estimate source locations of reactive beta activities.

**No**

---

Fig. 5b. Flow chart for ICA-based single-trial analysis method.
Figure 6a depicts the single-trial VAMW\textsubscript{rec} of subject I filtered within the trial-specific frequency band (Fig. 6b). The conventional AM method on the average of 100 epochs reveals a bilateral post-movement rebound pattern with contralateral (left hemisphere) dominance, whereas the current ICA-based single-trial analysis (one hemisphere template) yields only activation (one trial) in the left hemisphere (Fig. 6a and 6c).

Epoch acceptance rates were 84% (65/78), 89% (83/91), 71% (60/85), 73% (68/93), and 87% (76/87), respectively for the model generation group and 81% (71/88) for the validation subject; the average for all six was 80.8%. The IOIs of significance were 0.76s - 2.1s, 0.66s - 1.5s, 0.8s - 1.75s, 0.46s - 1.49s, and 0.71s - 1.28s for the five subjects in the model generation group, and 0.88s - 1.67s for the validation subject. Averaged magnitude of BR was calculated from the reconstructed data on trials that survived the epoch-selection procedure.
Fig. 6. Sensor-array display of VAMW$_{\text{recon}}$s and VAMWs. (a) One example of ICA single-trial VAMW$_{\text{recon}}$s of all sensor sites in subject I. The single-trial result shows only left sensorimotor area dominance of event-related activities, as the present study focuses on the area contralateral to movement side and only the left spatial template is used. The dashed trapezoid marks the nine SMI vicinal sensor sites and the VAMW$_{\text{recon max}}$ is marked with the red circle. (b) Trial-specific frequency band used for VAMW$_{\text{recon}}$s calculation in Fig. 6a. (c) VAMWs obtained from the conventional averaging method over 100 trials in subject I. This figure shows a bilateral beta rebound pattern with contralateral (left hemisphere) dominance.

The BR amplitudes computed from individual spatial templates were 20.9±7.1 (mean±sd), 18.1±10.3, 16.2±6.2, 23.2±10.89, and 6.2±2.7 for the first 5 subjects, respectively, and 27.6±11.1 fT/cm for the 6th subject (Table 1). Using the common spatial template, BR amplitudes were 21.1±7.97, 19.02±9.7, 15.5±5.3, 19.75±8.75, 5.91±3.2, and 27.1±10.2 fT/cm, respectively (Table 1). There was no significant difference between the results obtained with two approaches ($p=0.88$; unpaired two-tailed t test). BR amplitudes obtained with the conventional method of averaging on 100 trials were 18.2, 7.254, 12.92, 16.4, 2.9, and 23.12 fT/cm, respectively. Means for single-trial ICA-derived BRs, using either individual or common spatial templates, were significantly higher than those obtained using the conventional method of averaging ($p<0.005$; Matched-pair Wilcoxon test; Table 1). The comparisons of BR amplitude and task-specific frequency band between ICA-based single-trial and conventional methods are given in Table 1.
The ICA-based single-trial approach shows remarkable latency jittering and inter-trial variability throughout the whole measurement process. Both factors can result in attenuation and smearing of averaged movement-related MEG responses. Figure 7a shows the raster plot of sixty-five normalized single-trial VAMW\textsubscript{recon, max}s which survived the

(a)

(b)

Fig. 7. Smearing of MEG profile and decrease of BR magnitude due to latency jittering. (a) Raster plot of normalized VAMW\textsubscript{recon, max}s as sorted by the latency measured between the time of peak beta rebound and the movement onset. Black dashed line indicates movement onset time. (b) Latency jittering resulting in a smearing of the MEG profile and a decrease of BR magnitude when more VAMW\textsubscript{recon, max}s are averaged, as is common in the conventional averaging method.
selection procedure for subject I, sorted by VAMW$^*$ peak latency as indexed to movement onset. The mean latency of peak beta rebound for the 65 trials was 1.41±0.43 s (mean±sd). With more epochs (random selection) averaged as with the conventional method of averaging, the averaged BR was attenuated (25.3, 24.6, 22.3, 21.5, 20.3 and 21.1 fT/cm for 1, 15, 25, 35, 45 and 65 trials averaged, respectively; values taken from the averaged VAMW$^*$ using common spatial template) and the time-activity plots smeared (Fig. 7b).

<table>
<thead>
<tr>
<th>Subject index</th>
<th>Individual spatial template</th>
<th>Common spatial template</th>
<th>Individual spatial template</th>
<th>Common spatial template</th>
<th>Trial-specific frequency band (Hz)</th>
<th>BR amplitude (fT/cm)</th>
<th>Task-specific frequency band (Hz)</th>
<th>BR amplitude (fT/cm)</th>
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<tr>
<td>I</td>
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<td>18.04±2.62</td>
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Table 1. The comparison of BR amplitude and specific frequency bands for ICA-based single-trial and conventional methods.

Source estimation using ECD and MCE both showed a cluster of current sources centered (mean coordinates) in the anterior bank of the central sulcus (see Fig. 8e and 8f) on data points around the rebound peak of extracted reactive beta activities (see Fig. 8d, time interval between 1202ms – 1302ms of one single epoch of subject I). The ECD-located dipoles oscillate and span a sector. Furthermore, the center of MCE-estimated current sources (yellow dots) lies less than 2mm from the center of ECD-estimated dipoles (red dots) (see Fig. 8f). These results cross-verify the validity of the ICA-based single-trial method.

Figure 10 depicts the time-frequency plot of a normal subject and a Parkinson’s disease patient at an MEG channel in the vicinity of left sensorimotor area. Clear suppression of post-movement ERS (red circle) and an attenuated ERD (yellow circle) are observed in the Parkinson’s disease patient. The VAMWs was significantly larger both in alpha band and beta band in the normal subject than in the Parkinson’s patient. These imply the slowed and reduced recovery after motor act may impede cortical preparation of the next movement.
Fig. 8. Overlay of extracted reactive beta activities on MR image. (a) Spatial map reconstructed using $\tilde{x}_{\text{recon}} = \sum_{j=1}^{k} \tilde{x}_{j}$ (see Method Section). (b) and (c): Different views of superposition of the isocontour spatial map on the segmented MRI brain. (d) Representative trace of reconstructed reactive beta activities in the vicinity of SMI. (e) Upper panels are isocontour maps of reconstructed neuromagnetic signals at 1202 ms post movement. Lower panels show that all dipoles (from 1202 to 1302 ms after movement onset as box-framed in
(d)) are located in the primary motor area and oscillate accordingly. (f) The center of the MCE-estimated current sources (yellow dot) overlays the source location determined using the equivalent current dipole method (ECD) (red dot). Upper-left panel: coronal view. Upper-right panel: sagittal view. Lower-left panel: axial view. Lower-right panel: distribution of MCE estimated current sources.

Examining the single-trial variability using the proposed ICA-based method (Fig. 11), subtle dynamics of the beta rhythmic activities can be further studied. Figure 11 shows the ongoing trial-by-trial variabilities in amplitudes and latencies over 60 ICA de-noised post-movement ERS trials. With the utilization of ICA-based single-trial analysis, it is possible to investigate the reasoning of ERS vanishment is due to suppression of post-movement beta rebound in each single-trial, rather than the cause of temporal jittering and/or loss of synchronization. Even though the patient could perform lifting behavior well, his neuron activities show distinct sensorimotor patterns from normal subject, regardless of movement performance.

3. Discussion

The movement-related oscillatory modulations (ERD/ERS of alpha, beta and gamma) have been reported to be spatially extended (Babiloni et al., 1999; Crone et al., 1998a; Crone et al., 1998b; Leocani et al., 1997; Neuper and Pfurtscheller, 2001; Salmelin and Hari, 1994a; Taniguchi et al., 2000; van Burik et al., 1998). Source localizations using conventional filtering have also been reported to disperse among several regions (Salmelin & Hari, 1994a). However, our results strongly indicate that proper treatment when trial-by-trial dynamics can be accounted for yields clustered localizations congruent to neuroanatomical representations.

The present ICA-based spatiotemporal approach for single-trial analysis study is dedicated to the extraction of neuromagnetic measurements of event-related beta oscillatory activities. One distinct feature of the current ICA-based method as compared with other single-trial approaches (Guger et al., 2000; Ioannides et al., 1993; Jung et al., 2001) is the simultaneous use of a spatial template and a temporal template for component selection. The spatial template provides a priori spatial information for brain signals, while the temporal template contains temporal characteristics of event-related responses. Using the paired criteria for component selection, identification specificity of task-related components for signal reconstruction is significantly improved. As shown in Fig. 3c, the inclusion of IC 9 with high spatial correlation (correlation value=0.61, rank= 95.2%, Z=1.67, P=0.048) but devoid of temporal congruence (correlation value =0.28, rank=13%, Z=0.34, P=0.87) causes beta BR to deteriorate from 28.9 fT/cm (red curve) to 18.6 fT/cm (blue curve). The ICA-preprocessed dataset yields cleaner field maps (Fig. 9a), which result in circumscribed localizations (Figs. 9b-9c and 9e-9f., Salmelin and Hari, 1994a).

Significantly, the current method also makes possible the analysis of the reactive frequency band for every single trial once task-related rhythmic activities are extracted. The conventional method discounts this subtle but potentially important information. Notwithstanding, the idea of using a fixed window for signal filtering is neurophysiologically not optimal. We emphasize the precise identification of reactive trial-specific frequencies for BR calculation, since task-related frequency modulation might exist in one or multiple bands (Pfurtscheller & Lopes da Silva, 1999). The three-standard frequency band procedure is used for generation of VAMW to recover all possible task-related information and is followed by a two short-time spectra
Fig. 9. Comparisons of magnetic fields and source locations preprocessed with ICA-bandpass trial specific (upper panel) and task-specific bandpass filtering (lower panel). (a) Neuromagnetic field maps. Data preprocessed with ICA-trial specific bandpass filter (15.57±3.21–22.17±3.3 Hz) gives a much less noisy neuromagnetic field pattern than that processed with the task-specific bandpass filtering method (15–21 Hz) (Pfurtscheller et al. 1999). Black vertical lines in the tracings of the left column denote time points of the corresponding field maps in the right column. (b) Source localizations by ECD model. Only dipoles in post-movement IOI (interval-of-interest) with goodness-of-fit higher than 80% are accepted. The one with highest goodness-of-fit value out of each trial is rendered onto the subjects’ 3D MRI surfaces. The estimated source positions preprocessed by ICA-bandpass filtering (upper panel) are (x, y, z)=(-45±4.45, -3.9±6.33, 80.7±3.63mm; goodness-of-fit=97.5±3.7%) in subject I (65 trials) and (x, y, z)=(-35.3±3.5, 5.7±6.02, 88.7±5.61mm; goodness-of-fit=96.9±3.7%) in subject VI (71 trials), whereas task specific bandpass filtering (lower panel) yields (x, y, z)=(-46.3±11.6, -9.99±10.3, 84.51±6.7; goodness-of-fit=89.7±3.4%) in subject I (65 trials) and (x, y, z)=(-31.8±8.13, 0.5±14.01, 87.9±12.54mm; goodness-of-fit=87.2±4.5%) in subject VI (71 trials), respectively. The ICA-trial specific bandpass procedure yields better
results in terms of much focused source locations and higher goodness-of-fit. \( x, y, \) and \( z \) denote the dipole location in the head coordinate system as anchored by the HPI (head position indicator) coils. The x-axis passes through the preauricular points, pointing to the right; the positive y-axis traverses the nasion and is normal to the x-axis; the positive z-axis points upward and normal to the xy-plane.

![Fig. 10. The comparison of neural activity via time-frequency, VAMW and spectrum analysis obtained from one normal subject and a parkinson’s disease patient. The ensemble averaging results reflect the ERD attenuation (yellow circle) and the ERS disappearance (red circle) in the parkinson’s patient.](image)

![Fig. 11. Trial-by-trial comparison of VAMWs could be performed via the proposed ICA-based approach between the normal subject and the Parkinson’s disease patient.](image)
Extraction of Single-Trial Post-Movement MEG Beta Synchronization in Normal and Parkinson’s Patient Using ICA-Based Spatiotemporal Approach

comparison procedure (Pfurtscheller & Lopes da Silva, 1999) for the identification of the optimal reactive trial-specific frequency band in the reconstructed epochs. The present approach not only extracts the specific reactive frequencies but also retains phase information on a trial-by-trial basis. The trial-specific frequency band of post-movement beta modulation anchors mainly (~85% of all trials) in the lower beta band (16Hz~20Hz) and less frequently (~15%) in the higher beta band (20~24Hz). Great variation of BR values is also seen, as reflected in large SD (Table 1). The revealed trial-by-trial dynamics provide a possibility for future profound study of subtle brain dynamics.

It is noteworthy that not all the data reconstructed from the selected ICs survives the statistical threshold. We have carefully monitored online and thoroughly checked offline the EMG measurements in terms of EMG onset ($p=0.61$, unpaired two-tailed t-test), termination ($p=0.53$, unpaired two-tailed t-test) and the EMG duration ($p=0.573$, unpaired two-tailed t-test) during finger lifting between significant and insignificant trials as indexed to the movement registration by the optic pad (Abbink et al., 1998). The data indicate an absence of prominent behavioral difference commensurate to the differential neuromagnetic responses. Some epochs with a fluctuating baseline, e.g., non-task-related spontaneous bursts of beta oscillatory activities, may manifest high baseline activity, which in turn results in a decrease in BR readout leading to exclusion after statistical manipulation (Fig 4). It has been suggested that baseline spontaneous activities may carry important information relevant to attention level, wakefulness, task difficulty, etc. (Buser & Rougeul-Buser, 1999; Sterman, 1999). The jittering of the neuromagnetic beta ERS is likewise interesting and may be also physiological. A zero-phase Butterworth filter was used to bandpass filter the raw data. The symmetric property of the zero-phase filter means that processed signals have precisely zero phase distortion and therefore no time shift of peak beta rebound was introduced. Hence, fluctuations of significance level and the jittering of central processing despite similar behavioral performance may be ascribed to the subject’s variant cognitive states or the degree of training (Buser & Rougeul-Buser, 1999; Sterman, 1999; Flotzinger et al., 1992; Wolpaw et al., 1994; Bastiaansen et al., 2001; Bastiaansen et al., 1999; Earle, 1988; Haig et al., 1995; Hoffman et al., 1991; Yabe et al., 1993). The exploration of underlying mechanisms mandates more meticulous designs in the future. Using the conventional method of averaging, certain diseases, such as Parkinson’s and Unverricht-Lundborg myoclonic epilepsy, have been observed to show either attenuated, prolonged or abolished ERS responses (Silent et al., 2000; Tamas et al., 2003). Such cases can be further examined using the current ICA-based single-trial method for the time course and trial-by-trial dynamics to disclose hitherto unexplored mechanisms underlying these phenomena.

A concern with any data driven method is that prominent artifacts or noise can be intermingled with task-specific information (Ermer et al., 2000; Lins et al., 1993a; Lins et al., 1993b). However, previous ICA reports (Makeig et al., 2002; McKeown & Radtke, 2001) indicate that brain rhythmic signals generated from different sources usually have their own oscillatory frequencies with distinct phases and are located in specific brain regions with patterns that are distinct from artifacts or noise (see also Fig. 3). This endorses the feasibility of using ICA to separate targeted rhythmic signals from irrelevant ones. The high epoch-acceptance rate (~80%) can be attributed to an improved SNR as compared to other studies on single-trial approaches to sensorimotor oscillatory activities (Brovelli et al., 2002; Wolpaw and McFarland, 1994). For instance, the spatial map of IC2 in Fig. 2a correlates highly (0.88) with the spatial map of IC1 from empty room measurement as shown in Fig. 3b; this
suggests that the neuromagnetic signal IC2, deselected for subsequent processing, can be accounted for by background noise in the shielding room. IC11 in Fig. 3a has a stationary cycle around 1.2 Hz, and its spatial map has higher weights at the outer rim of the MEG sensor array, which suggests a plausible connection with cardiac cycles. It was also observed (Fig. 3) that rhythmic activities in left and right SMIs as well as the occipital areas could be extracted into separate ICs that can be reminiscent of various mechanisms and time courses of different brain oscillatory activities (Pfurtscheller & Lopes da Silva, 1999; Pfurtscheller et al., 1997; Pfurtscheller et al., 1998b; Stancak and Pfurtscheller, 1996a; Stancak & Pfurtscheller, 1996b; Andrew & Pfurtscheller, 1999).

Since most task irrelevant signals, e.g., internal and external noises, can be removed by proper de-selection of ICs, it is possible to reconstitute the representative spatial map of all contributing ICs using $\tilde{x}_{\text{recon}} = \sum_{j=1}^{k} \tilde{x}_j$, in which $\tilde{x}_{\text{recon}}$ is the reconstructed spatial map, $k$ is the number of selected ICs and $\tilde{x}_i$ is the spatial map of the $i$th selected IC in Fig. (4) (Fig. 8a). This spatial map of reconstructed signals, which is a topographical distribution of weighting factors on the sensor array, can be overlaid with the segmented MRI brain (Fig. 8b & 8c; ASA program, ANT Software, Dutch). The highest weight is shown to project over the SMI area, which demonstrates that the high SNR of the ICA-extracted rhythmic activities of each trial has made possible the use on single-trial data of source estimation methods that require high SNR on input data for processing, e.g., the equivalent current dipole technique (ECD), minimum current estimation (MCE), and minimum norm estimation (MNE) (Delorme et al., 2001; Delorme et al., 2002; Jung et al., 2001; Makeig et al., 1997; McKeown et al., 2001). Conventionally, these estimation methods exploit averaged data out of a large amount of trials.

Another reason why the intricate phase-unlocked signal can be preserved is the fact that no averaging procedure is needed; such a procedure would otherwise inherently distort the embedded information. Accordingly, as shown in Fig. 8d, source modeling with a moving dipole on a msec by msec basis on the reconstructed oscillatory beta signals during the rebound period (Brovelli et al., 2002) of a single-trial epoch results in a focused clustering of dipole foci at the pre-central area, i.e., the primary motor cortex (Fig. 8e). Figure 8 shows the result of MCE modeling (Uutela et al, 1999), where the center of MCE-estimated current sources (yellow dot) is very close (< 2mm distance) to the dipole location as estimated using the ECD approach (red dot).

It can be argued that one can first localize the generator area and then build a spatial filter for extracting single-trial data so that the subsequent analysis can be conducted on the source level instead of the sensor level. One premise and justification of using a source-area-generated spatial filter is that the source area can be precisely localized for the generation of a spatial filter (Tesche et al., 1995). The very first step is to filter the signals to obtain a presupposed reactive frequency band. However, using conventional simple filtering techniques, ambient noise with ~20Hz components cannot be optimally removed, and this will cause localization uncertainty for the probed sources (Fig 9). However, ICA pre-processing decomposes the compound neuromagnetic signals into various independent task-related and task-unrelated/noise components so that ~20Hz activities not related to the a priori spatiotemporal profile will not confound the selected ones. Furthermore, our ICA-based method differs from other spatial filtering techniques, e.g., signal space projection (SSP) which is a fixed spatial filter for signal extraction (Tesche et al., 1995). The ICA-based
method blindly decomposes the MEG epochs ($B$) into a spatially distributed map ($U$) multiplied by temporal signals ($S$), i.e. $B = U \cdot S$, on the basis of independency among sources (Vigario & Oja, 2000), whereas SSP mandates a pre-defined spatial filter ($U_{sf}$) for recovering signals ($S$), i.e. $S = U_{sf}^{+} \cdot B$, where $+$ denotes pseudo inverse, based on orthogonal projection. When ambient noise and the spatial filter are not mutually orthogonal, the SSP has difficulty in resolving the two. Subsequent application of ICA following SSP does not ensure finer signal extraction or further noise removal since the data recovered from SSP are already linear mixtures of components out of a pre-defined spatial filter, which is a constraint drag on the optimal performance of ICA designed for blind decomposition.

Left and right sensorimotor rhythms can be decomposed into two distinct ICs (IC3 and IC5 in Fig. 3), implying possible independent modulatory mechanisms between the two hemispheres. This view is corroborated by an event-related coherence study (Andrew and Pfurtscheller, 1999) that reports a lack of interhemispheric coherence in human post-movement beta activities. Movement-related beta oscillatory activities of the right hemisphere can be extracted in the same way using spatial and temporal templates for right sensorimotor rhythm. The source locations for extracted right hemispheric beta activities were mainly in the right premotor area (data not shown), which agrees with previous studies (Brovelli et al., 2002; Ilmoniemi, R. J., 1991). Event-related beta activities in SMA and posterior parietal cortical areas (Brovelli et al., 2002; Joliot et al., 1999) are not observed in our data, possibly due to the fact that the contributing sources here are radial in orientation and thus could not be optimally detected by MEG (Salmelin and Hari, 1994b).

The agreement between the values of BR amplitude obtained with the common spatial/temporal templates and the individually generated ones (Table 1) promises a flexibility in both experimental design and analytical strategy. The proposed ICA-based spatiotemporal approach for single trial analysis can also be applied on fewer trials (Fig 7b), which is a great advantage over conventional methods. Given meticulous head positioning (see above the Method Section), common spatial and temporal templates can be used to extract pertinent movement-related neuromagnetic signals from subjects, which may shorten the overall time needed to run an experiment. We have no preference for the use of a grand averaged template over individual ones. On the contrary, the use of an individual template is suggested for any profound individual-based ERD/ERS study. However, the feasibility of using a grand averaged template provides an effective alternative in cases where lengthy procedures cannot be endured by the participants. This is particularly true for clinical settings where patients have attention problems or are incapable of sustaining long experiments so that individual templates cannot be optimally obtained. Nevertheless, caution should be exercised when applying the current ICA-based single-trial method for clinical studies. For patients whose heads cannot be properly positioned in the center of the MEG helmet, the use of a common spatial template may fail, making a customized individual spatial template mandatory for IC selection. For patients whose motor performance deviates significantly from normal, e.g., victims of motor stroke or severe movement disorders, the use of the common temporal template might not be justified since the time courses of event-related brain activities may be significantly altered due to primary deficit or secondary plasticity. Accordingly, in such situations, an individual spatial template can be applied without a temporal template as an aid to component selection. Our future investigations will combine the current dual-template approach with a source estimation method so that a spatial filter of better precision and higher dimensions can be
designed, which will make possible sophisticated analysis on the source level instead of the sensor level, eliminating the positioning problem.

Degeneration of the dopaminergic neurons in substantia nigra pars compacta (SNc) in Parkinson’s patients result in abnormal projection in thalamo-cortical pathway which causes an abnormal projection from thalamus to supplementary motor area (SMA). Pfurtscheller et al. (1998) also have demonstrated that Parkinson’s patients have delayed ERD and abolished post-movement ERS and speculated there is dysfunction in subcortico-cortical connections in Parkinson’s patients. In this study, we analyzed post-movement ERS in one Parkinson’s patient. The present ICA-based approach may be helpful for disclosing the mechanism of movement-related brain rhythms which could be used as a clinical index for diagnosing Parkinson’s patients.

4. Conclusions

The present novel ICA-based spatiotemporal approach for single trial analysis features a paired-template matching for stringent component selection. The spatial template provides a priori spatial information for targeted brain signals while the temporal template contains temporal characteristics of event-related responses. The method promises not only a high extraction rate of post-movement beta synchronization but also better localization of the corresponding sources. Various source modeling methods commanding high SNR can now be applied to single trial data as extracted using the ICA-spatiotemporal procedure. Our method takes into account subtle trial-by-trial dynamics. The reconstructed MEG brain signals per trial unravel the temporal information and inter-trial variations of reactive oscillatory activities, which in turn may shed light on the subtle dynamics of brain processing. The embodied common template approach permits an effective alternative in cases where lengthy procedures cannot be endured by the participants or in clinical settings where patients have attention problems or are incapable of sustaining long experiments.

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6. References


Diagnostics and Rehabilitation of Parkinson's Disease presents the most current information pertaining to news-making topics relating to this disease, including etiology, early biomarkers for the diagnostics, novel methods to evaluate symptoms, research, multidisciplinary rehabilitation, new applications of brain imaging and invasive methods to the study of Parkinson's disease. Researchers have only recently begun to focus on the non-motor symptoms of Parkinson's disease, which are poorly recognized and inadequately treated by clinicians. The non-motor symptoms of Parkinson's disease have a significant impact on patient quality of life and mortality and include cognitive impairments, autonomic, gastrointestinal, and sensory symptoms. In-depth discussion of the use of imaging tools to study disease mechanisms is also provided, with emphasis on the abnormal network organization in parkinsonism. Deep brain stimulation management is a paradigm-shifting therapy for Parkinson's disease, essential tremor, and dystonia. In the recent years, new approaches of early diagnostics, training programmes and treatments have vastly improved the lives of people with Parkinson's disease, substantially reducing symptoms and significantly delaying disability. Written by leading scientists on movement and neurological disorders, this comprehensive book should appeal to a multidisciplinary audience and help people cope with medical, emotional, and practical challenges.

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