Time-Dependent Statistical and Correlation Properties of Neural Signals during Handwriting

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Abstract

To elucidate the cortical control of handwriting, we examined time-dependent statistical and correlational properties of simultaneously recorded 64-channel electroencephalograms (EEGs) and electromyograms (EMGs) of intrinsic hand muscles. We introduced a statistical method, which offered advantages compared to conventional coherence methods. In contrast to coherence methods, which operate in the frequency domain, our method enabled us to study the functional association between different neural regions in the time domain. In our experiments, subjects performed about 400 stereotyped trials during which they wrote a single character. These trials provided time-dependent EMG and EEG data capturing different handwriting epochs. The set of trials was treated as a statistical ensemble, and time-dependent correlation functions between neural signals were computed by averaging over that ensemble. We found that trial-to-trial variability of both the EMGs and EEGs was well described by a log-normal distribution with time-dependent parameters, which was clearly distinguished from the normal (Gaussian) distribution. We found strong and long-lasting EMG/EMG correlations, whereas EEG/EEG correlations, which were also quite strong, were short-lived with a characteristic correlation durations on the order of 100 ms or less. Our computations of correlation functions were restricted to the β spectral range (13–30 Hz) of EEG signals where we found the strongest effects related to handwriting. Although, all subjects involved in our experiments were right-hand writers, we observed a clear symmetry between left and right motor areas: inter-channel correlations were strong if both channels were located over the left or right hemispheres, and 2–3 times weaker if the EEG channels were located over different hemispheres. Although we observed synchronized changes in the mean energies of EEG and EMG signals, we found that EEG/EMG correlations were much weaker than EEG/EEG and EMG/EEG correlations. The absence of strong correlations between EMG and EEG signals indicates that (i) a large fraction of the EEG signal includes electrical activity unrelated to low-level motor variability; (ii) neural processing of cortically-derived signals by spinal circuitry may reduce the correlation between EEG and EMG signals.

Introduction

Since the first publication by D. Walter [1], the coherence method, developed for the analysis of stationary random data in linear systems (see, e.g., [2]), has been employed in hundreds of papers dealing with the analysis of neural signals such as EEGs and EMGs. In these publications, the level of coherence was used as a measure of coupling between the processes generating neural signals and of the functional association between neuronal structures [3–6].

This analysis of relationships between neural signals is based on computations of the coherence and phase of the two signals. For Fourier harmonics, \(X(\omega)\) and \(Y(\omega)\), of two time-dependent signals \(X(t)\) and \(Y(t)\), the coherence is defined as the square of the modulus, \(C(\omega) = |P(\omega)|^2\), and the phase is defined as \(\Phi(\omega) = \arctan(Im P(\omega)/Re P(\omega))\), of the complex coherence function

\[
P(\omega) = \frac{X(\omega)Y^*(\omega)}{|X(\omega)X^*(\omega)|^{1/2}|Y(\omega)Y^*(\omega)|^{1/2}}
\]

Citation: Rupasov VI, Lebedev MA, Erlichman JS, Lee SL, Leiter JC, et al. (2012) Time-Dependent Statistical and Correlation Properties of Neural Signals during Handwriting. PLoS ONE 7(9): e43945. doi:10.1371/journal.pone.0043945

Editor: Natasha M. Maurits, University Medical Center Groningen UMC, The Netherlands

Received April 19, 2012; Accepted July 27, 2012; Published September 11, 2012

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Funding: This work was supported by National Science Foundation (www.nsf.gov) grants 0848523 (IIP), 1048430 (RAHSS), 1048428 (RET) (Principal Investigator: ML), and by a National Instruments (www.ni.com) grant. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing Interests: ML is paid employee and the owner of Norconnect Inc., and VIR is an employee of Norconnect Inc. ML is going to develop products from this and future research. He is the author of pending patents with United States Patent Office. Recordation of handwriting and hand movement using electromyography 11640954, Handwriting EMG for Medical Diagnosis 61108603, Method and Apparatus for Biomedical Analysis Using EEG, and EMG signals 13341465. ML has 88% stock ownership in Norconnect Inc. and is the board chairman at Norconnect Inc. This research was supported by research grant #0848523 to Norconnect from National Science Foundation and a National Instruments grant. There are no further patents, products in development or marketed products to declare. This does not alter the authors’ adherence to all the PLoS ONE policies on sharing data and materials, as detailed online in the guide for authors.

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Here, Re \( P \) and Im \( P \) stand for the real and imaginary parts of the function \( P \).

Signals \( X(t) \) and \( Y(t) \) can also be sliced into \( M \) disjoint segments, \( \{X_j(t)\}, \{Y_j(t)\}, j = 1, \ldots, M \), and the function \( P(\omega) \) may be estimated as shown by Eq. (1) with auto- and cross-spectral density functions computed for each segment and averaged over \( M \) segments,

\[
P(\omega) = \frac{1}{M} \sum_j X_j(\omega) Y_j^*(\omega) \quad \text{subject to} \quad \frac{1}{M} \sum_j X_j(\omega) X_j^*(\omega) = 1, \quad \frac{1}{M} \sum_j Y_j(\omega) Y_j^*(\omega) = 1
\]  

(2)

Coherence is considered to be statistically significant if its magnitude exceeds some value \( L \). At any desired confidence level \( \alpha \% \), the confidence limit is estimated under a Gaussian assumption as:

\[
L = 1 - (1 - \frac{\alpha}{100})^{1/2}
\]  

(3)

A relatively small number of segments, \( M \), suffices to achieve any desired confidence limit due to the exponential dependence of the confidence limit on the number of segments.

Neural signals are obviously nonstationary. To address the nonstationarity of biological signals, the coherence method has been generalized for wavelet harmonics [9]. Wavelet-based methods allow a combined frequency- and time-domain representation of nonstationary signals.

In this paper, we propose and discuss an alternative approach to search for dynamical relationships between neural signals [10]. In this method, which is broadly employed in statistics and, in particular, in statistical physics, a relationship between two random time-dependent signals \( x(t) \) and \( y(t) \) is determined by the correlation function

\[
C(t_1, t_2) \sim \int dx dy [x(t_1) - \mu_x(t_1)][y(t_2) - \mu_y(t_2)]p(x, y).
\]  

(4)

Here \( p(x, y) \) is the joint probability density function of two random variables, \( \mu_x \) and \( \mu_y \) are the corresponding mean values \( \mu_x(t) = \int x(t)p(x)dx \) and \( \mu_y(t) = \int y(t)p(y)dy \), where \( p \) is the probability density function. It should be emphasized that for nonstationary systems, the time dependence of correlation functions is determined not only by the time dependencies of the signals themselves, but also by the time dependencies of the probability density functions.

For two independent random variables, the joint probability density function can be factorized, i.e., \( p(x, y) \sim p(x)p(y) \), and the correlation function \( C \) vanishes.

The probability density functions of neural signals are not known a priori. Therefore, one needs to have a sufficiently large set (statistical ensemble) of neural signals \( \{x_j(t)\} \) and \( \{y_j(t)\}, j = 1, \ldots, N \) recorded during \( N \) epochs - in our case, trials during which a subject repeatedly performs an identical task - in order to apply the statistical method. In this approach, the integration of probability density functions in Eq. (4) is replaced by the ensemble average over many trials,

\[
C(t_1, t_2) \sim \frac{1}{N} \sum_{j=1}^{N} [x_j(t_1) - \mu_x(t_1)][y_j(t_2) - \mu_y(t_2)].
\]  

(5)

where \( \mu_x(t) = \frac{1}{N} \sum_j x_j(t) \) and \( \mu_y(t) = \frac{1}{N} \sum_j y_j(t) \) are the mean values.

This expression becomes exact only in the limit when the number of trials goes to infinity, \( N \to \infty \). In our experiments the number of trials was about 400, and we used Fisher’s theorem [11] to compute the confidence interval for the correlation functions.

Here we apply this statistical method to an analysis of EEG and EMG signals recorded during handwriting. Handwriting [12] provides an excellent neuromuscular task for quantitative studies of statistical and correlational properties of biological signals in the time domain. Handwriting consists of relatively simple, stereotyped hand movements that involve two basic motor components: firmly holding a pen by the fingers and moving the hand and the fingers to produce written text. We recorded the mechanical events of handwriting using a digitizing tablet that electronically recorded each epoch when the pen touched the paper (pen-on-paper period), and we simultaneously recorded the cortical EEG and EMG of intrinsic muscles of the hand.

Our dataset allowed us (i) to align the mechanical events, EEG signals and EMG signals associated with handwriting and (ii) to divide all of these signals into trial segments that correspond to a sequential set of epochs as each subject performed a pen-on-paper activity.

The recorded data consisted of \( N \) trials, each trial captured the activity associated with a single handwritten letter. Each trial contained EEGs recorded in \( N_{\text{elec}} \) channels, and EMGs recorded in \( N_{\text{emg}} \) channels, i.e., in total, we obtained \( N \times N_{\text{elec}} \times N_{\text{emg}} \) data files each of 2000 ms duration.

In contrast to the coherence methods used to study the relationship between neural signals in the frequency domain, our statistical method enabled us to study the statistical and correlational properties of neural signals directly in the time domain. That allows us to elucidate both dynamical patterns of activity in different cortical areas and the functional relationships between different cortical areas.

As in the case of EMG signals recorded from muscle groups involved in handwriting [13], we found that trial-to-trial variability of the “energy” of the EEGs recorded from the motor cortex area had a log-normal distribution, which was clearly distinguishable from the normal (Gaussian) distribution. The log-normal distribution fitted the experimental data in all time intervals during a trial, but its parameters - the mean value and dispersion - depended on time. These two variables suffice to completely describe both qualitatively and quantitatively the trial-to-trial variability of neural signals during handwriting.

We also studied time-dependent EMG/EMG and EEG/EEG correlations. We found strong and long-time EMG/EMG correlations. Correlations between EEG signals over the motor cortex area were also quite strong, but the correlations existed over relatively short durations with characteristic correlation times on the order of 100 ms or less. All subjects involved in our experiments were right-handed and wrote with their dominant hand. However, we observed equally strong inter-channel correlations when both channels were located on either the left or right hemispheres, while inter-hemispheric correlations were 2–3 times weaker.

Although we observed task-related changes in the mean energies of both EEG and EMG signals, we found that EEG/EMG correlations were much weaker than EEG/EEG and EMG/EMG correlations. In other words, trial-to-trial variations
of the magnitudes of the EEG and EMG changes were related only weakly.

**Methods**

This study was approved by the Institutional Review Board of Human Participants Research of Dartmouth College, Hanover, NH. No personal information was recorded during the sessions, and all data were analyzed anonymously. Written informed consent was obtained from the subjects prior to the recording sessions.

A schematic of the experimental setup for simultaneous recording EMG and EEG signals during handwriting is shown in Figure 1. We used LogiPen to record handwriting characters. EMG signals were sampled using bipolar surface EMG electrodes (Kendall Arbo). We used EMG analog amplifiers with gain 1000, low pass filter of 450 Hz, and high pass filter of 12 Hz. The EMG signals were digitized at a sample rate of 1000 Hz. The surface EMGs of the intrinsic hand muscles were recorded by two electrode pairs. One pair recorded EMG activity from flexor pollicis brevis and abductor pollicis brevis. The second pair recorded from the first dorsal interosseus muscle (Figure 2). EEG activity was sampled with a standard 64-channel device. The data from the 64-channel EEG recording system were high-pass filtered with 8 Hz cutoff, digitized at 1000 Hz and saved on a computer. Handwriting and EMG inputs were synchronized by a LabView data acquisition program based on the pen-down event associated with each handwriting trial. We used LogiManage as the digitizer of handwriting traces. It recorded X and Y coordinates, as well as the pen-on-paper signal. EMG signals were recorded on a dedicated computer that was also connected to the EEG recording system via a parallel port. The pen-on-paper signals were used to align EEG and EMG recordings.

To carry out quantitative studies of time-dependent statistical and correlation properties of neural signals, we instructed seven subjects to write the digit “3.” This task produced consistent motor patterns with some trial-to-trial variability. The single character was written about 400 times in 10 blocks of 40 trials. The blocks were separated by 5 minute rest intervals.

Handwriting trials were defined as the epochs starting 1000 ms before the time moment when the pen touches the tablet at the onset of writing a particular character and ending 1000 ms later. In all cases, the character was completely written before the end of the trial. After filtering the signals, their amplitudes, \( A_{emg} \) and \( A_{eeg} \), containing frequencies from 8 to 450 Hz, were squared to get the signal “intensities” \( I_{emg} = A_{emg}^2 \) and \( I_{eeg} = A_{eeg}^2 \).

To study time-dependent statistical and correlation properties of the signals, 2000-ms trials were subdivided into 20 time intervals, each with a duration of 100 ms, and the signal “energy” was calculated for each of 20 intervals as the sum,

\[
E_{emg}(n,z,j) = \sum_{\text{interval}} I_{emg}(n,z,j),
\]

(6a)

\[
E_{eeg}(n,z,j) = \sum_{\text{interval}} I_{eeg}(n,z,j),
\]

(6b)

where \( n = 1 \ldots 20, \; z = 1 \ldots 2 \) for EMG signals and \( z = 1 \ldots 64 \) for EEG signals, and \( j = 1 \ldots N \) (where \( N \) is the total number of trials), enumerate the time intervals, recording channels, and trials, respectively.

The minimal length of the time intervals was dictated by the accuracy with which trials could be aligned with respect to each other. In our experiments, the accuracy of alignment did not exceed a few tens of milliseconds, which was determined by the accuracy of the digitizing tablet used to detect the first moment of time when the pen touches the paper. Therefore, we chose a 100-ms time interval. This interval substantially exceeds the temporal resolution of the digitizing tablet and ensures an accurate trial alignment.

To obtain dimensionless variables for each interval and trial, the energies \( E_{emg}(n,z,j) \) and \( E_{eeg}(n,z,j) \) were normalized by dividing by their mean values \( \bar{E}_{emg}(n,z) = \langle E_{emg}(n,z,j) \rangle \) and \( \bar{E}_{eeg}(n,z) = \langle E_{eeg}(n,z,j) \rangle \). Where the symbol \( \langle \ldots \rangle \) stands for averaging over trials,

\[
\langle \ldots \rangle = \frac{1}{N} \sum_{j=1}^{N} \ldots
\]

(7)

Thus, the EMG and EEG signals for each recording channel \( z \) and each trial \( j \) were characterized by dimensionless energies

\[
\bar{E}_{emg}(n,z,j) = \frac{E_{emg}(n,z,j)}{\bar{E}_{emg}(n,z)},
\]

(8a)

\[
\bar{E}_{eeg}(n,z,j) = \frac{E_{eeg}(n,z,j)}{\bar{E}_{eeg}(n,z)},
\]

(8b)
and their time dependence was described by the discrete variable \( n \) (time interval number).

**Results**

1. Statistics of EMG signals

We previously studied the statistical properties of EMG signals during handwriting [13]. The results of this previous study are summarized in this Section for the convenience of readers and for further comparison with the properties of the EEG signals. Correlations properties of EMG signals, which have not studied earlier, are discussed in the next Section.

**1.1. Mean values and variation coefficients.** Trial-averaged intensities of raw EMG signals, or EMG templates, \( I_{emg}^{(temp)} = \langle I_{emg} \rangle \) described the characteristic pattern of EMG activity for the handwritten character for each of two muscle groups (Figure 3a). The time dependency of the mean energies \( E_{emg}(n,x) \) is shown in Figure 3b. The first 9 time intervals represent the epoch preceding the first pen touch. This subject wrote the digit “3” with a mean duration of the pen-on-paper period (i.e., time during which the pen touched the paper) of about 600–700 ms, which corresponds to time intervals 10 through 15 or 16, while the intervals 17 through 20 correspond to the time period following the pen liftoff from the paper.

The variation coefficients, \( V_{emg}(n,x) \), which characterize the dispersion of the data distribution, were computed as the ratio of the standard deviation

\[
S_{emg}(n,x) = \sqrt{\langle (E_{emg}(n,x,j) - E_{emg}(n,x))^2 \rangle} \quad (9a)
\]

to the mean energy,

\[
V_{emg}(n,x) = \frac{S_{emg}(n,x)}{E_{emg}(n,x)} \quad (9b)
\]

The variation coefficients were large for all time intervals and for both EMG channels (Figure 3c) and ranged from about 0.3 to 1. Thus, there was a substantial dispersion of EMG signals from trial to trial, and the EMG activity during handwriting could not be adequately described in terms of mean values only. More detailed analysis of the statistical properties of the EMG signals was required.

It should be noted that while the mean energies increased during the pen-on-paper period, the dispersion decreased. EMG patterns were more consistent when the subject wrote characters than when he/she lifted the pen off the paper. This pattern of the mean-energy and dispersion was observed for all subject, characters, and muscles.

**1.2. Distribution functions.** We recorded a sufficiently large number of trials to approximate the theoretical probability distribution from our data. In Figure 4, the probability plots for experimental data (with no trial selection) are shown together with probability plots for the theoretical normal and log-normal distributions. It is clear that the experimental data are fitted well with the log-normal probability density function

\[
p(x) = \frac{1}{\sqrt{2\pi}\sigma} \exp \left\{ - \frac{(\log x - \mu)^2}{2\sigma^2} \right\}, \quad (10)
\]

where \( \log \) stands for natural logarithm.

Moreover, it is clear from Figure 4 that the fitted curves for log-normal and normal distributions are easily distinguished, and that the log-normal distribution fits the data much better than the normal one.

2. Correlation functions of EMG signals

We observed that activation patterns of different muscles were coordinated during handwriting and the EMG bursts occurred at consistent interburst intervals. Mathematically, this relationship can be described by statistical correlation functions as described in the Introduction.

Since the logarithms of the random dimensionless energies \((8a)\),

\[
\epsilon(n,x,j) = \log(E_{emg}(n,x,j)), \quad (11)
\]

are normally distributed across trials for each time interval, we may assume that the joint probability density function \( P(\epsilon(n,x,j), \epsilon(m,y,j)) \) for pairs of the logarithms of dimensionless energies is the joint (second order) normal probability density function [2].
This allows one to compute both the correlation functions
\[ C(n, m, a, b) \sim S_{\text{emg}}(n, a) S_{\text{emg}}(m, b) \times S_{\frac{1}{2}} e^{(n, a, j)} C_{22} / C_{13} S_{\frac{1}{2}} e^{(m, b, j)} C_{22} / C_{13} T, \]
where \( S_{\text{emg}}(n, a) = \langle s(n, x, j) \rangle \), and, using the Fisher transform [11], to estimate the confidence intervals for the correlations. The Pearson correlations functions, defined in Eq. (12), are normalized to the standard deviations; therefore, their moduli do not exceed 1. We have described those correlation coefficients that approached 1 and were statistically significant as “strong” and those that approached 0 as “weak” correlations.

Figure 5 shows the cross-correlation function \( C(n, m, x, \beta = 1, \beta = 2) \) for two groups of muscles (EMG channels 1 and 2) involved in handwriting as a function of two times \( n \) and \( m \) (Figure 5a) and the diagonal elements of this function at coincident time intervals \( n = m \) (Figure 5b). It is easy to see strong and long-time correlations between activities of muscle groups (Figure 5a) and particularly strong correlations at coincident time intervals. Correlations were higher outside the pen-on-paper period, when the subjects held the pen in the air, and decreased shortly before and during the pen-on-paper period.

Finally, autocorrelation functions, \( C(n, m, x = 1, \beta = 1) \) and \( C(n, m, x = 2, \beta = 2) \) for both EMG signals are shown in Figure 6. Here again, strong correlations are clear over long time for each muscle group.

3. Statistical properties of EEG signals
As can be seen from the EMG templates, which indicate distinct bursts of activity (Figure 3a), and from the time-dependency of the EMG energy (Figure 3b), muscle activity was clearly patterned during the handwriting of the character. EEG activity even in the motor cortex area should not be expected to be limited to the handwriting itself and can contain other components that are not necessarily related to handwriting. Therefore, the total amplitude of EEG signal reads as

\[ A_{\text{eeg}} = A_{\text{eeg}}^{(\text{handwriting})} + A_{\text{eeg}}^{(\text{other})} \]
and the non-handwriting contribution to the EEG obviously obscures the handwriting related component.

If a signal with amplitude \( A_{\text{eeg}}^{(\text{other})} \) is unrelated to handwriting trials, averaging over many trials should reduce its contribution to the mean values and correlation functions. Nevertheless, since we square the total amplitudes of EEG signals, the contribution of the non-handwriting activity does not vanish completely.

Figure 4. Probability plots for EMG signals. (a) Plot of empirical cumulative distribution function (time interval 11; EMG channel 1) together with theoretical cumulative distribution function plots for normal and log-normal distributions. The confidence bounds are shown for the confidence level of 95%. (b) Probability plot for experimental data for time interval 11, channel 1 together with probability plots for theoretical normal and log-normal distributions. Axis scales are chosen to have a straight line for the theoretical probability plot with the log-normal distribution.

doi:10.1371/journal.pone.0043945.g004

Figure 5. EMG/EMG correlation functions. (a) The cross-correlation function for EMG channels 1 and 2 (EMG1 and EMG2) as a function of two times: intervals for EMG1 and intervals for EMG2. (b) The cross-correlation function for EMG1 and EMG2 for coincident time intervals. Error bars indicate 95% confidence bounds.

doi:10.1371/journal.pone.0043945.g005
To indicate each EEG channel, we used conventional electrode names shown in Figure 7.

3.1. Mean values and variation coefficients. In contrast to EMG signals, we did not find prominent bursts in the EEG templates $I_{\text{eeg}}^{(\text{temp})} = \langle I_{\text{eeg}} \rangle$ for any of 64-channels that could be attributed to handwriting activity. Figure 8a represents the time behavior of the mean energy of EEG signal, $E(n,z)$, of channel C1, which was located over the left hemisphere.

The mean energy and variation of the EEG signal decreased about 300–400 ms before the pen-on-paper period and remained low during the actual period of writing. A similar time dependency of the mean energy was observed for channels FC5, FC3, FC1, C5, C3, CP5, CP3, and CP1 over the left hemisphere. It should be noted that during the 300-ms time period preceding the pen-on-paper period, EMG activity was increased (see Figure 3a and Figure 3b).

Thus, we found good correspondence between changes in EMG activity and EEG activity (Figure 8a) over the motor cortex. Even
though all subjects involved in our studies were right handed, similar EEG changes were observed for the midline channels Cz and CPz, and the channels FC2, FC4, FC6, C2, C4, C6, CP2, CP4, and CP6 corresponding to the right hemisphere, locations that were mirror images of the channels with synchronized EMG/EEG activity over the left hemisphere.

As in the case of EMG signals, we defined the variation coefficient as a ratio of the standard deviation,

\[ S_{\text{EEG}}(n, z) = \frac{\sqrt{\langle [E_{\text{EEG}}(n, z)]^2 - E_{\text{EEG}}(n, z)^2 \rangle}}{E_{\text{EEG}}(n, z)} \]  

(14a)

to the mean energy,

\[ V_{\text{EEG}}(n, z) = \frac{S_{\text{EEG}}(n, z)}{E_{\text{EEG}}(n, z)} \]  

(14b)

As in the case of EMG signals, the variation coefficients decreased shortly before and during the pen-on-paper period. Since the variation coefficients were quite large (Figure 8b), EEGs during handwriting could not be adequately described in terms of mean values only, and more detailed studies of their statistical properties were required.

The relative magnitude of the EEG energy decrease during the pen-on-paper period was variable from subject to subject. It was maximal in the \( \alpha \) (8–13 Hz) and, especially, \( \beta \) (13–30 Hz) spectral ranges and less apparent in the \( \gamma \) (30–100 Hz) spectral range (Figure 9).

It should be emphasized that the variation coefficients in the \( \beta \) spectral range, where we observed the largest relative decrease in the mean EEG energies, increased quite significantly (up to two times) in comparison to other spectral ranges.

3.2. Distribution functions. Figure 10 shows that, as in the case of EMG signals, the theoretical log-normal function fits our EEG data better than a normal distribution. The theoretical log-normal cumulative distribution function lies inside the confidence bounds computed with the confidence level of 95%. Similar results were found in all time intervals for the EEG channels corresponding to the motor cortex in both left and right hemispheres. Some small deviations observed in some of the time intervals could be attributed to a contribution of the component of EEG activity not associated with handwriting activity, \( A_{E_{\text{EEG}}}^{(\text{other})} \).

Thus, the log-normal distribution effectively approximated the trial-to-trial variability of both EMG and EEG energy.

Since the relative magnitude of the EEG energy decrease during the pen-on-paper period was maximal in the \( \beta \) spectral range, we restricted our computations of correlation functions for EEG signals to this spectral range.

4. Correlation functions of EEG signals

Since logarithms of dimensionless energies of EEG signals, including energies of EEG signals in the \( \beta \) spectral range,
Standardize the presentation of EEG/EEG correlations for the motor cortex area, it was convenient to average the function over the time intervals, i.e. the diagonal elements of the correlation matrices for each time interval to get a dynamical picture of the functional connectivity between different neural regions of the cortex during handwriting (see Figure 11, Figure 12, and Figure 13).

The statistical method enabled us to easily derive the correlation coefficients for each time interval to get a dynamical picture of functional connectivity between different neural regions of the cortex during handwriting (see Figure 12 and Figure 13).
...intervals before and during handwriting. Thus, volume conduction appears to have a small impact on the correlations that we computed based on the ensembles of identical trials.

Moreover, in Figure 15 we present the correlation function between EEG signals recorded in channels FC5 and CP6. We observed time-dependent and statistically significant correlations on the order of 0.2–0.3 despite the maximally large (see Figure 7) spatial separation between electrodes located in the motor cortex area with the inter-electrode spacing, \( L \), about of \( \sqrt{160} \) cm. The contribution of the dipole electric field generated by the cortex located under the electrode FC5 to the electrical potential recorded by the electrode CP6 was estimated in the simplest way as \( (h_L) \sim 1/160 \), where \( h \) (\( \sim 1 \) cm) is the distance between the electrode and cortex. Thus, the contribution of the electrical field generated at FC5 to activity at CP6 was very small, and the correlation between these EEG signals cannot be attributed to the volume conduction effect.

Finally, we studied the correlation function \( D(n,m,x,\beta) \) dependent on two times, or two sets of time intervals (Figure 16). In sharp contrast to the case of muscle activity (Figure 7 and Figure 8), we did not find any long-time correlations between EEG signals recorded over the motor cortex. The characteristic correlation time for a single channel was of the order of 100 ms or less, as seen from Figure 16a, where the correlation function \( D_C(n,m) = D(n,m) \) for channel C3 is plotted.
Moreover, we did not find long-time correlations between channels lying within the left, right, or central motor cortex sites. The characteristic correlation time in this case, which corresponded to the characteristic time of cross-talk between channels, was also on the order of 100 ms or less. More accurate computations of the characteristic correlation times would require a more precise trial alignment, which in our experiments was within several tens of milliseconds.

Finally, we found that correlations between channels that overlapped different parts of the motor cortex, e.g., channel C3 (the left part) and channel C4 (the right part) [Figure 16b], were strongest at coincident time intervals and quickly became insignificant as the time interval between recorded events increased. Thus, there were no significant long-time correlations between channels among different parts of the motor cortex.

5. Correlation functions of EMG and EEG signals

Since the logarithms of both EMG energy and EEG energy in motor cortex channels were normally distributed over trials, we computed the correlation functions between EMG and EEG signals as

\[
K(n,m,x,y) = \frac{1}{S_{\text{emg}}(n,x)S_{\text{emg}}(m,y)} \times \frac{\langle \bar{e}(n,x) \bar{e}(m,y) \rangle}{\langle \bar{e}(n,x) \rangle \langle \bar{e}(m,y) \rangle}
\]

which detects simultaneous trial-to-trial variations of EEG activity and EMG activity [Figure 17]. As in the case of EEG/EEG correlation functions, we restricted our computations to the \( \beta \) spectral range of EEG signals.

The energy of EMG signals grew in the time period shortly before and during the pen-on-paper period [Figure 3a and 3b], while the energy of EEG signals in the motor cortex channels decreased during the same time period [Figure 9b]. Therefore, we expected to find an anticorrelation between EMG and EEG signals, i.e., the correlation function \( K \) would be negative in the time intervals shortly before and during the pen-on-paper period.

Although correlation coefficients at coincident time intervals, shown in Figure 18, were statistically significant at some time intervals, they were smaller than EEG/EEG and EMG/EMG correlation coefficients. When statistically significant EMG/EEG correlations occurred, they were inconsistent among subjects.

The absence of strong correlations between EMG and EEG signals may indicate that strong trial-to-trial variability may be in part introduced by spinal motoneuron activity, which results in wide dispersion of EMG signals with respect to EEG signals. Additionally, EEG components unrelated to motor output may decrease correlations between EEG and EMG signals.

Discussion

To derive a dynamical picture of neural activity and of functional relationships between different neural regions, we examined time-dependent statistical and correlation properties of EMG and EEG signals recorded simultaneously during handwriting of digit “3” by 7 subjects. We recorded signals in approximately 400 2000-ms trials during which each subject performed an identical handwriting task. The trials started 1000 ms before the moment of time when the pen touched paper the first time and finished 1000 ms after this moment of time. To study the time dependence of neural signals, the trials were divided into 20 100-ms time intervals.

We studied trial-to-trial variability of EMG signals and EEG signals derived from the motor cortex during the 2000-ms time interval during which subjects performed the same handwriting task. We found that the trial-to-trial distribution of the neural signal energy was described well by a log-normal distribution and not by a normal distribution. Whereas the distribution parameters - the mean value and the dispersion - depended on intra-trial time, the log-normal distribution was found for all 20 time intervals.

We computed the Pearson auto- and cross-correlation functions for EMG signals and EEG signals recorded from the motor cortex. We observed very strong correlations at coincident time intervals between EMG signals recorded from different muscle groups. Moreover, these correlations were long-time and remained quite strong during almost the entire 2000-ms time period.

We also found strong correlations between EEG signals in \( \beta \) spectral range, provided that channel pairs were located in the motor cortex of the same hemisphere. The correlations between signals recorded from different sides of the skull were much weaker. Moreover, in contrast to the EMG/EEG correlations,
Finally, we found cross-correlation functions between EEG and EMG signals. We suggest that low correlation coefficients between EEG and EMG activities may be explained by an abundance of EEG signals unrelated to low-level motor parameters and by non-cortical sources of a trial-to-trial variability of spinal motorneuron activity even though the subjects performed a stereotypical handwriting task.

Acknowledgments

We thank Newton de Faria of National Instruments for his assistance in hardware selection and experimental design, and Patrick Britz and the technical support teams of Brain Vision and Brain Products.

Author Contributions

Conceived and designed the experiments: ML. Performed the experiments: SLL ML. Analyzed the data: ML VIR MAL JSE JCL SLL. Contributed reagents/materials/analysis tools: ML. Wrote the paper: ML VIR MAL JCL JSE SLL. Designed the software used in analysis: MAL VIR ML.

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