Study of the MEG Background Activity in Alzheimer’s Disease Patients with Scaling Analysis Methods

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Abstract— Alzheimer’s disease (AD) is one of the most prominent neurodegenerative disorders. The aim of this research work is to study the magnetoencephalogram (MEG) background activity in AD patients using two scaling analysis methods: detrended fluctuation analysis (DFA) and backward detrended moving average (BDMA). Both measures have been designed to quantify correlations in noisy and non-stationary signals. Five minutes of recording were acquired with a 148-channel whole-head magnetometer in 15 patients with probable AD and 15 control subjects. Both DFA and BDMA exhibited two scaling regions with different slopes. Significant differences between both groups were found in the second region of DFA and in the first region of BDMA ($p < 0.01$, Student’s t-test). Using receiver operating characteristic curves, accuracies of 83.33% with DFA and of 80% with BDMA were reached. Our findings show the usefulness of these scaling analysis methods to increase our insight into AD.

I. INTRODUCTION

Magnetoencephalography (MEG) is a non-invasive technique that allows recording the magnetic fields produced by brain activity. It provides an excellent temporal resolution, orders of magnitude better than other methods for measuring cerebral activity, as magnetic resonance imaging, single-photon-emission computed tomography or positron-emission tomography [1]. A good spatial resolution can also be achieved due to the large number of sensors. Moreover, the activity in different parts of the brain can be monitored simultaneously with whole-head equipments, such as the magnetometer used in the present study [1]. On the other hand, the magnetic signals generated by the human brain are extremely weak. Thus, SQUID (Superconducting QUantum Interference Device) sensors are necessary to detect them. In addition, MEG signals must be recorded in a magnetically shielded room. Thus, MEG is characterized by limited availability and high equipment cost.

Alzheimer’s disease (AD) is a progressive and irreversible brain disorder of unknown aetiology. It is the main cause of dementia in western countries, accounting for 50-60% of all cases [2]. AD affects 1% of population aged 60-64 years, but the prevalence increases exponentially with age, so around 30% of people over 85 years suffer from this disease [3]. Additionally, due to the fact that life expectancy has significantly improved in western countries in the last decades, it is expected that the number of people with dementia increase up to 81 million in 2040 [3]. AD is characterized by neuronal loss and the appearance of neuritic plaques containing amyloid-β-peptide and neurofibrillary tangles [4]. Clinically, this disease manifests as a slowly progressive impairment of mental functions whose course lasts several years prior to death [4]. Usually, AD starts by destroying neurons in parts of the patient’s brain that are responsible for storing and retrieving information. Then, it affects the brain areas involved in language and reasoning. Eventually, many other brain regions are atrophied. Thus, AD patients may wander, be unable to engage in conversation, appear non-responsive, become helpless and need complete care and attention [5]. Although a definite AD diagnosis is only possible by necropsy, a differential diagnosis with other types of dementia and with major depression should be attempted. The differential diagnosis includes medical history studies, physical and neurological evaluation, mental status tests, and neuroimaging techniques.

The electromagnetic brain activity has been researched in the last decades by means of non-linear techniques. Correlation dimension has been widely used to study the brain activity in AD patients [6, 7]. Nevertheless, reliable estimation of this classical measure requires a large number of data points and stationary and noise-free time series [8]. As these problems cannot be solved for physiological signals, other non-linear methods are necessary to study brain recordings. In fact, Lempel-Ziv complexity [9-11], Higuchi’s fractal dimension [12], approximate entropy [11], sample entropy [10], synchronization likelihood [13], phase lag index [14], and auto-mutual information [15] have been already used to analyze the MEG activity in AD.

In this preliminary study, we have examined the MEG background activity in patients with probable AD and inagematched control subjects using two scaling analysis measures: detrended fluctuation analysis (DFA) and backward detrended moving average (BDMA). Our purpose is to test the hypothesis that the neuronal dysfunction in AD is associated with differences in the dynamical processes underlying the MEG recording.

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II. MATERIAL AND METHODS

A. Subjects and MEG recording

The signals were recorded using a 148-channel whole-head magnetometer (MAGNES 2500 WH) located in a magnetically shielded room. The subjects lay comfortably on a patient bed, in a relaxed state and with their eyes closed. They were asked to stay awake and to avoid eye and head movements. For each subject, five minutes of recording were acquired at a sampling frequency of 678.17 Hz. These recordings were down-sampled by a factor of four, obtaining a sampling rate of 169.55 Hz. Data were digitally filtered between 0.5 and 40 Hz. Finally, artifact-free epochs of 5 seconds (848 samples) were selected.

MEG data were acquired from 30 subjects. Cognitive status was screened in both groups with the Mini Mental State Examination (MMSE). The AD group consisted of fifteen patients (5 men and 10 women; age = 72.33 ± 9.04 years, mean ± standard deviation, SD) fulfilling the criteria of probable AD, according to the criteria of the National Institute of Neurological and Communicative Disorders and Stroke - Alzheimer’s and Related Disorders Association. The mean MMSE score for the patients was 17.67 ± 3.94 points. Patients were free of other significant medical, neurological and psychiatric diseases than AD. Moreover, any of the participants in the study used medication that could be expected to influence in the MEG recording.

The control group consisted of fifteen control subjects without past or present neurological disorders (7 men and 8 women; age = 72.53 ± 5.40 years, MMSE score = 29.00 ± 0.33 points). All control subjects and all caregivers of the patients gave their informed consent for the participation in the current study. The local Ethics Committee approved this study.

B. Detrended Fluctuation Analysis (DFA)

DFA is a measure widely used for the detection of long-range correlations and fluctuations in time series [16]. This method provides a simple quantitative parameter (the scaling exponent) to represent the correlation properties of a signal [17]. DFA permits the detection of long-range correlations embedded in seemingly non-stationary time series. Additionally, it avoids the spurious detection of apparent correlations and fluctuations in time series [16]. This approach to quantify correlation properties in non-stationary signals with underlying trends [19], Serletis and Rosenberg [20] suggested that DFA is an improvement over DMA. DMA does not require dividing the series into non-overlapping windows. Instead, the DMA method detrends the series by subtracting a continuous function, the moving average. Additionally, DMA is more accurate since the moving average is a better low-pass filter when compared to the polynomial filter used for DFA [20]. In this study, we have used the category of DMA called BDMA [21]. To perform the BDMA of a time series $X = x_1, x_2, ..., x_N$, we pursue the following steps [21]:

1) The signal $X$ is integrated:

$$y_k = \sum_{j=1}^{N} y_j - \bar{X}$$

where $\bar{X}$ represents the mean value of $X$.

2) The integrated time series $Y$ is divided into windows of equal size $k$. The size of the windows is ranged between 3 and 84, as one-tenth of the signal length can be considered as the maximum window size when using DFA [17].

3) Within each window, labeled $b (b = 1, 2, ..., B)$, perform a least-square fit of $Y$ by a straight line $F^b$. This is the semilocal trend for the $b^{th}$ window.

4) Define $F^2(k)$ to be the variance of the fluctuation $Y$ from $F^b$ in the $b^{th}$ window:

$$F^2(k) = \frac{1}{k} \sum_{j=1}^{k} (y_j - F^b)^2$$

5) The square root of the average of $F^2(k)$ over all windows is the rms fluctuation from the semilocal trends in $B$ windows, each having $k$ time points:

$$F(k) = \left( \frac{1}{B} \sum_{b=1}^{B} F^2(k) \right)^{1/2}$$

6) Finally, the study of the dependence of $F(k)$ on the window size $k$ is the essence of DFA. If it is a power-law behavior $F(k) \propto k^\alpha$, the scaling exponent is an indicator of the nature of the fluctuations in the MEG signals.

C. Backward Detrended Moving Average (BDMA)

The detrended moving average (DMA) method is a new approach to quantify correlation properties in non-stationary signals with underlying trends [19], Serletis and Rosenberg [20] suggested that DMA is an improvement over DFA. DMA does not require dividing the series into non-overlapping windows. Instead, the DMA method detrends the series by subtracting a continuous function, the moving average. Additionally, DMA is more accurate since the moving average is a better low-pass filter when compared to the polynomial filter used for DFA [20]. In this study, we have used the category of DMA called BDMA [21]. To perform the BDMA of a time series $X = x_1, x_2, ..., x_N$, we pursue the following steps [21]:

1) The signal $X$ is integrated:

$$y_k = \sum_{j=1}^{N} y_j - \bar{X}$$

where $\bar{X}$ represents the mean value of $X$.

2) The second step of the BDMA is to detect trends in data employing a moving average. For a window of size $k$, the backward moving average is defined as:

$$y^b_k = \frac{1}{k} \sum_{n=0}^{k-1} x_{n+b}$$

3) Afterwards, the signal is detrended by subtracting the trend $y^b$ from the integrated profile $Y$:

$$C_k = y_k - y^b_k$$

4) Then calculate the fluctuation for a window of size $k$ as:

$$F(k) = \frac{1}{N-k+1} \sum_{k=1}^{N-k+1} (C_k)^2$$

5) As in DFA method, a power law relation between the fluctuation function $F(k)$ and the scale $k$ indicates a self-similar behavior.
III. RESULTS

DFA method was applied for the 148 MEG channels using window sizes between 3 and 84 samples. To simplify the analyses, the results were averaged over all channels. \( F(k) \) as a function of \( k \) was plotted in a log-log scale, as Fig. 1 illustrates. If the plot displays a linear scaling region with a certain scaling exponent, then there is a power-law behavior in the time series [22]. Our results showed two scaling regions with different slopes: \( \alpha_1 \) for \( 3 \leq k \leq 7 \), and \( \alpha_2 \) for \( 14 \leq k \leq 73 \). No significant differences were found between the \( \alpha_1 \) values of AD patients and control subjects (\( p = 0.7107 > 0.01; \) Student’s \( t \)-test). On the other hand, the differences between both groups were statistically significant when the slopes of the second scaling region were analyzed (\( p = 0.0015 < 0.01; \) Student’s \( t \)-test).

We next apply the BDMA method to the MEG recordings with windows sizes between 2 and 84 samples. BDMA profiles were obtained representing the natural logarithm of \( F(k) \) versus the natural logarithm of \( k \) (see Fig. 2). This scaling analysis method also showed two scaling regions, \( \alpha_1 \) and \( \alpha_2 \). Significant differences between both groups were found with \( \alpha_1 \) (\( p = 0.0001 \)), whereas the \( p \)-value obtained with \( \alpha_2 \) was 0.1065.

Finally, receiver operating characteristic (ROC) curves were used to assess the ability of these measures to discriminate AD patients from controls. This method summarizes the performance of a two-class classifier across the range of possible thresholds. Fig. 3 and 4 represent the ROC curves obtained at both scaling regions with DFA and BDMA, respectively. The highest accuracy was achieved when the slope \( \alpha_2 \) of DFA was analyzed: 83.33%. In the first scale region of BDMA, we achieved the highest value of area under the ROC curve (AROC): 0.8667. Table I shows the sensitivity, specificity, accuracy and AROC values obtained with DFA and BDMA in each scaling region.

IV. DISCUSSION AND CONCLUSION

We analyzed the MEG background activity from 15 patients with probable AD and 15 age-matched control subjects by means of DFA and BDMA methods. Our purpose was to check the hypothesis that MEG background activity was different in AD patients than in control subjects. Both measures have proven to be effective in discriminating AD patients from controls. Our results revealed that \( \alpha_2 \) values of DFA were significantly lower for the controls than for the AD patients. On the other hand, significant differences between both groups appear in the first scaling region of BDMA. Because both groups were carefully matched for age, these changes in the fluctuations of MEG signals may well represent the cognitive dysfunction in AD. Additionally, accuracies of 83.33% and 80% were reached with DFA and BDMA by means of ROC curves.

Other non-linear methods have been already used to study the MEG activity in AD. In [9-11], Lempel-Ziv complexity values were significantly lower in the recordings from AD patients than in those obtained from control subjects. This complexity loss in AD was confirmed using the algorithm of fractal dimension proposed by Higuchi [12]. Nevertheless, van Cappellen van Walsum et al. [7] suggested that this decreased complexity in the MEG background activity of AD patients may appear only in the low frequency bands. Other MEG studies revealed that brains affected by AD show a more regular physiological behaviour [10, 11]. Stam et al. [13] found changes of long and short distances.
Therefore, future efforts will be addressed to characterize these measures can detect a gradation of the disease process. Secondly, our results do not show if the sample size is small. Thus, a larger database is needed to confirm our results. A neural complexity measure applied to MEG data in Alzheimer’s disease,” Clin. Neurophysiol., vol. 114, pp. 1034–1040, 2003.


REFERENCES


