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PERMUTATION DISALIGNMENT INDEX AS AN INDIRECT, EEG-BASED, MEASURE OF BRAIN CONNECTIVITY IN MCI AND AD PATIENTS

NADIA MAMMONE, LILLA BONANNO, SIMONA DE SALVO,
 SILVIA MARINO, PLACIDO BRAMANTI

*IRCCS Centro Neurolesi Bonino-Pulejo
 Via Palermo c/da Casazza, SS. 113, 98124 Messina, Italy
 nadiamammone@tiscali.it, nadia.mammone@irccsme.it*

ALESSIA BRAMANTI

*Institute of Applied Sciences and Intelligent Systems Eduardo Caianiello (ISASI)
 National Research Council (CNR), Messina (Italy)*

FRANCESCO C. MORABITO

*DICEAM Department of the Mediterranean University of Reggio Calabria
 Via Graziella Feo di Vito, 89060 Reggio Calabria, Italy morabito@unirc.it*

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OBJECTIVE: In this work, we introduce Permutation Disalignment Index (PDI) as a novel nonlinear, amplitude independent, robust to noise metric of coupling strength between time series, with the aim of applying it to electroencephalographic (EEG) signals recorded longitudinally from Alzheimer's Disease (AD) and Mild Cognitive Impaired (MCI) patients. The goal is to indirectly estimate the connectivity between the cortical areas, through the quantification of the coupling strength between the corresponding EEG signals, in order to find a possible matching with the disease's progression. **METHOD:** PDI is first defined and tested on simulated interacting dynamic systems. PDI is then applied to real EEG recorded from 8 amnesic MCI subjects and 7 AD patients, who were longitudinally evaluated at time T0 and 3 months later (time T1). At time T1, 5 out of 8 MCI patients were still diagnosed MCI (stable MCI), whereas the remaining 3 exhibited a conversion from MCI to AD (prodromal AD). PDI was compared to the Spectral Coherence and the Dissimilarity Index. **RESULTS:** Limited to the size of the analysed dataset, both Coherence and PDI resulted sensitive to the conversion from MCI to AD, even though only PDI resulted specific. In particular, the intrasubject variability study showed that the three patients who converted to AD exhibited a significantly ($p < 0.001$) increased PDI (reduced coupling strength) in delta and theta bands. As regards Coherence, even though it significantly decreased in the three converted patients, in delta and theta bands, such a behaviour was detectable also in one stable MCI patient, in delta band, thus making Coherence not specific. From the Dissimilarity Index point of view, the converted MCI showed no peculiar behaviour. **CONCLUSIONS:** PDI significantly increased, in delta and theta bands, specifically in the MCI subjects who converted to AD. The increase of PDI reflects a reduced coupling strength among the brain areas, which is consistent with the expected connectivity reduction associated to AD progression.

Keywords: Electroencephalography, Alzheimer's Disease, Mild Cognitive Impairment, Permutation Disalignment Index.

1. Introduction

Dementia is a general term used to refer to a wide range of symptoms, associated with a decline in memory or other thinking skills, which reduce a person's ability to perform everyday life activities. Alzheimer's disease (AD) is the most common form of dementia and accounts for 60% of dementia cases¹. Current diagnosis of AD relies on quantifying the mental decline. However, by the time the patient is diagnosed, the disorder has already caused severe brain damage. Studies suggest that AD begins years before the clinical symptoms become visible². Furthermore, normal ageing is also characterised by a slow decline of cognitive functions, thus discriminating normal ageing from AD, at a very early stage, can be challenging when relying only on standard procedures based on clinical evaluation.

Researchers hope to develop an accurate way to early detect AD before its devastating symptoms appear. To this purpose, high-risk population should be periodically monitored through an early detection system. With regard to high-risk population, Mild Cognitive Impairment (MCI) is a condition in which the subject experiences minor problems with cognition (memory or thinking) and involves nearly 5-20% of people aged over 65. In MCI patients, cognitive issues are worse than we normally expect for a healthy person of the same age, but the symptoms are not severe enough to interfere significantly with daily life, thus, MCI is not labelled as dementia². In recent clinical studies, *amnestic* MCI subjects were shown to have an increased risk of developing AD. The incidence of AD, observed in 4 studies on amnestic MCI, ranged from 10.2% to 26.0% (median: 17.6%) over 1 year and from 17.6% to 36.3% (median: 24.2%) over 2 years³. Therefore, this population should be studied on the long term and longitudinally, in order to develop a system that can help to predict the transition from MCI to AD. Such a system should necessarily be non-invasive, well tolerated and low-cost; thus it could be based on Electroencephalography (EEG). The EEG consists in recording the electrical potentials generated by the brain through a set of electrodes located on the scalp. In AD patients, EEG abnormalities arise because of anatomical and functional deficits of the cerebral cortex. One of the effects of such deficits is the functional disconnection between some cortical areas: such a disconnection affects the behaviour of the electrical activity of the brain and, therefore, alter the EEG⁴.

Adeli et al. presented an overview of computational modelling for the analysis of EEGs recorded from AD patients^{5,6}. The hallmarks of EEG abnormality in AD patients, compared to healthy controls, are often a shift

of the power spectrum to lower frequencies and reduced coherences among cortical regions⁷, which are thought to be associated with functional disconnections among cortical areas resulting from the death of cortical neurons⁸. An EEG-based diagnostic tool should reveal the effects of such disconnection over the scalp; to this purpose, the mapping of descriptive features^{9,10,11,12}, might help. Stam et al. proposed to compute the synchronisation likelihood of multichannel EEG data in AD patients, subjects with MCI and subjects with subjective memory complaints (SC). The synchronisation likelihood significantly decreased in the 14-18 and 18-22 Hz bands in AD patients compared with both MCI subjects and healthy controls¹³.

Dunkin et al. hypothesised that decreased coherence would be associated with cognitive dysfunction as assessed by neuropsychological tests. They found that reduced coherence was associated with impairment on specific neuropsychological tests. The results endorsed the hypothesis that coherence reflects a functional breakdown in communication between brain areas¹⁴. Sankari et al.¹⁵ presented a comprehensive study of intrahemispheric, interhemispheric and distal EEG coherence in AD patients. Their study showed a pattern of decrease in AD coherence, by indicating a decline in cortical connectivity but they also report exceptions in specific bands where an increase in coherence can be attributed to compensatory mechanisms. Adeli et al. presented a wavelet coherence investigation of EEG readings acquired from patients with AD and healthy controls. Pairwise electrode coherence and wavelet coherence were calculated over each frequency band (delta, theta, alpha, and beta)^{16,17}.

High EEG upper/low alpha power ratio was associated with cortical thinning and lower perfusion in the temporo-parietal lobe. Moreover, atrophy and lower perfusion rate were both significantly correlated with memory impairment in MCI subjects¹⁸. In¹⁹, correlation between EEG markers and volumetric differences in mapped hippocampal regions was estimated in AD patients. Results showed that the increase of alpha3/alpha2 power ratio is correlated with atrophy of mapped hippocampal regions in AD. In²⁰, the most significant results of recent studies on correlation between scalp EEG, cognitive decline, and anatomical substrate have been reviewed, with particular attention to the relationships between EEG changes and hippocampal atrophy.

Morabito et al.⁴ proposed a technique, based on Complex Networks, to analyse the longitudinal modifications of brain connectivity and information transfer in AD patients. They found out that the progres-

sion of AD was characterised by a loss of connected areas, in terms of network parameters. Since the evolution of AD is characterised by the progressive loss of functional connectivity within neocortical association areas, Giannakopoulos et al. supposed that event-modulated EEG dynamic analysis could allow to investigate the functional activation of neocortical circuits. They reviewed and summarised clinically significant results of EEG activation studies in this field and discussed future perspectives of researches which aim at reaching an early and individual prediction of cognitive decline in healthy elderly controls²¹. Prichep et al. reported results from initial quantitative electroencephalography (qEEG) evaluations of normal elderly subjects. Source localisation algorithms were used to identify the most probable generators of abnormal features in the EEG. Abnormalities were found in the prodromal EEGs of those subjects who later converted to dementia²². Rossini et al. analysed the cortical connectivity (spectral coherence) and the low resolution brain electromagnetic tomography sources of EEG rhythms in MCI patients at baseline and follow-up²³. They found out that low midline coherence and weak temporal source were associated with a 10% annual rate of MCI to AD conversion. The National Institute on Aging and the Alzheimer's Association (NIA/AA) workgroup of experts postulated that what is commonly considered Alzheimer should rather be considered a mere stage of a long, complex degenerating process²⁴ and, therefore, they strongly encouraged the researchers to engage longitudinal studies. However, the literature lacks longitudinal studies on MCI/AD, because keeping such patients and their caregivers loyal to a periodical follow-up program can be challenging.

To our best knowledge, and also according to the review paper²⁵, the most promising work about MCI conversion to AD is²⁶, they reported that IFAST method was able to predict the conversion from amnesic MCI to AD with high accuracy (85.98%) in a 1-year follow-up study. This methodology was later improved but it has been validated only in a MCI vs AD vs Controls cross study so far²⁷. As already claimed above, MCI and AD are known as disconnection disorders, because it is well accepted that MCI/AD weakens the connectivity between the areas of the brain. Changes in the connectivity between cortical areas are likely to induce changes in the coupling strength between the corresponding EEG signals, this is the reason why a measure of coupling strength between EEG signals might help to indirectly estimate the changes in the brain connectivity due the disease's progression. The present work resulted from a

translational research whose goal was to longitudinally evaluate the evolution of the brain connectivity in MCI and AD patients, indirectly, through the EEG. The specific aim was to compare the overall coupling strength between EEG signals at time T0 and at time T1, in MCI and AD patients, in order to assess if it could reflect the brain connectivity reduction that is expected to be induced by the disease's progression. Our first evaluation was carried out applying Spectral Coherence because it is the gold standard method for the indirect estimation of the brain connectivity from the EEGs of AD and MCI patients. The reason for that is Coherence-based analysis comes with most of the EEG processing systems used in clinical applications on MCI/AD²⁷. As Coherence showed a trend that was peculiar to converted MCI but also exhibited a false positive, even though our dataset was small, we decided to develop our own metric for the estimation of coupling strength between time series. We introduced the Permutation Disalignment Index (PDI), which is a novel measure of coupling between time series, that can help whenever a multivariate, amplitude invariant, robust to noise, nonlinear coupling strength analysis is necessary. With application to EEG signals, PDI can be interpreted as an indirect measure of the coupling strength between two or more cortical areas, estimated through the quantification of the alignment randomness between the corresponding EEG projected time series. From the theoretical point of view, we can address the advantages of analysing the EEG through PDI, as compared to Coherence, to endorse our choice to develop a novel, nonlinear, multivariate, symbolic coupling strength descriptor: 1) Coherence is not a multivariate but only a pairwise descriptor, thus it cannot be exploited to estimate the joint coherence of a group of electrodes, on the contrary, PDI can be both multivariate and pairwise thus it can be exploited to estimate the coupling strength between a pair of electrode sites or among a group of electrodes covering a more extended area; 2) Coherence is linear and can fail to capture the nonlinear dynamics of the EEG; 3) Coherence is not a symbolic methodology and can be more sensitive to noise, since our method relies on the projection onto a predetermined set of symbols (the motifs), the possible presence of amplitude burst in the EEG due to artifacts would not cause amplitude burst in the projected symbols, therefore in the descriptor. Considered such motivations, PDI was hereby defined, implemented and, first of all, tested on simulated unidirectionally coupled Henon maps, in order to validate PDI's ability to measure the coupling strength between interacting dynamic systems. Since PDI success-

fully reflected the coupling strength variations in the simulated coupled systems, it was tested on real EEG signals recorded during a follow-up study which involved 7 AD patients and 8 amnesic MCI patients. The patients were evaluated at time T0 and then, 3 months later, at time T1.

Besides Coherence, PDI was also compared to the Dissimilarity Index (Dm) proposed by Ouyang et al.²⁸, which is a nonlinear, symbolic dissimilarity measure between time series that, in our opinion, was the best-matching nonlinear method.

According to the achieved results, PDI seems to have a high potential in the analysis of MCI/AD EEG time series. We hope that the present work encourages longitudinal studies on MCI patients, which are of paramount importance in the prediction of the degeneration from MCI to AD.

The paper is organised as follows: Section 2 describes how the patients were selected and how the EEGs were recorded and preprocessed. Section 3 introduces the concept of PDI and describe how it was tested on simulated data and on the experimental EEG data. Section 5.3 introduces the standard comparative methodologies used in the quantitative EEG analysis (spectral coherence and dissimilarity index). Section 6 reports the achieved results, Section 7 discusses the results and Section 8 addresses the conclusions.

2. EEG data recording and preprocessing

2.1. Study population

Fifteen patients, 7 AD subjects and 8 MCI subjects, at various stages of clinical evolution, were recruited at the IRCCS Centro Neurolesi *Bonino-Pulejo* of Messina (Italy). All patients were enrolled within an ongoing cooperation agreement that also included a clinical protocol approved by the local Ethical Committee. All patients signed an informed consent form. The clinical diagnosis was performed, according to the Diagnostic and Statistical Manual of Mental Disorders (fifth edition, DSM-5)²⁹, by a multidisciplinary team including neurologists, psychologists, psychiatrists and EEG experts, through a complete medical assessment. The same examiners conducted all the cognitive and clinical evaluations. Every patient underwent a neuroradiological examination to exclude other conditions (tumors, strokes, damage from severe head trauma or buildup of fluid in the brain, etc) that may cause symptoms similar to AD but require different treatment. Current use of any medications (but particularly cholinesterase inhibitors (ChEis), Memantine, anti-depressants, anti-psychotics and anti-epileptic drugs) was also taken into account

in AD patients. MCI patients were not undergoing any medical treatment. All patients were assessed at baseline (time T0) and 3 months after the first evaluation (time T1). At time T1, 3 MCI patients exhibited a conversion from MCI to AD. Table I provides details about gender (Female/Male), age and diagnosis (MCI or AD) of every patient.

Table I: Dataset description

ID	Age	Gender	Diagnosis T0	Diagnosis T1
pt_03	68	M	MCI	AD
pt_23	84	F	MCI	MCI
pt_30	69	M	MCI	MCI
pt_41	78	M	MCI	MCI
pt_51	71	F	MCI	AD
pt_57	83	M	MCI	MCI
pt_71	79	F	MCI	AD
pt_72	65	F	MCI	MCI
pt_31	74	M	AD	AD
pt_54	83	F	AD	AD
pt_64	74	F	AD	AD
pt_65	76	M	AD	AD
pt_76	79	F	AD	AD
pt_86	83	F	AD	AD
pt_87	78	F	AD	AD

2.2. EEG recording

The EEG was recorded in a comfortable resting state, according to the 10-20 International System (19 channels: Fp1, Fp2, F3, F4, C3, C4, P3, P4, O1, O2, F7, F8, T3, T4, T5, T6, Fz, Cz and Pz), at a sampling rate of 1024 Hz, by using a notch filter at 50Hz, with linked earlobe (A1-A2) reference. Before the EEG recording, all patients and their caregivers underwent a semi-structured interview including questions about: (a) quality of the last night sleep; (b) duration of the last night sleep; (c) meal timing and content. The EEG recordings were performed in the morning. During the acquisition of the EEG, the patients kept their eyes closed but remained awake. The technician, keeping the subject alert by calling her/him name, prevented the drowsiness: the corresponding EEG segments were discarded to avoid the effects of auditory cortex activation. The patients did not sleep during the recording as confirmed by the EEG recordings which did not show any sleep pattern. The EEG was initially sampled at 1024Hz, according to the general EEG recording protocol adopted at the IRCCS Neurolesi Center. The sampling rate was set high in order to make the EEG dataset as general as possible, in order to be suitable also for possible future researches with different goals.

2.3. EEG preprocessing

EEG signals are commonly decomposed into the four major EEG sub-bands: delta (0-4 Hz), theta (4-8 Hz), alpha (8-12 Hz), and beta (12-32 Hz). Each of the four sub-bands relates to different functional and physiological parts of the brain. In this paper, the EEG was band-pass filtered at 0.1-32Hz, then it was split into the four sub-bands through a set of band-pass filters implemented in the toolbox EEGLab (<https://sccn.ucsd.edu/eeglab/>)³⁰. In particular, we used the function *eegfiltfft*, based on Fast Fourier Transform (FFT) and inverse FFT to reconstruct only the specific frequency range under consideration. Figure 1 shows an EEG signal (filtered between 0.1 and 32 Hz) and the corresponding sub-bands. Starting from one n -channels EEG recording, we eventually end up with 4 different n -channels EEG recordings, each one corresponding to a different frequency sub-band: EEG_{δ} , EEG_{θ} , EEG_{α} , EEG_{β} . Each sub-band-EEG was then downsampled to 256Hz. Considered the specific application, 256Hz is a sampling rate high enough to capture delta, theta, alpha and beta waves, which are typically analysed in AD/MCI patients.

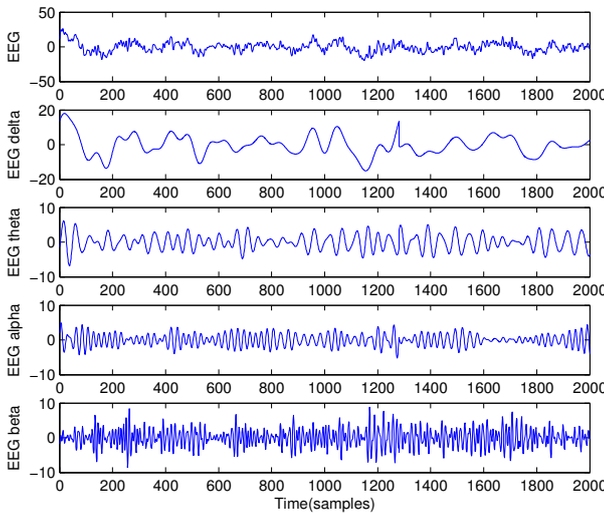


Fig. 1. An EEG signal (filtered between 0.1 and 32 Hz and expressed in μV), and the corresponding delta (0-4 Hz), theta (4-8 Hz), alpha (8-12 Hz), and beta (12-32 Hz) sub-bands.

Every sub-band-EEG recording was partitioned into 5 s non-overlapping windows (as the sampling rate after downsampling is 256Hz, each window included $N_s = 1280$ EEG samples); therefore, each sub-band-EEG was eventually partitioned into N_w non-overlapping windows, where N_w depends of the length of the recording. All the algorithms were implemented in Matlab

(The MathWorks, Inc., Natick, MA, USA).

Before processing the filtered EEG through the proposed algorithm, artifactual segments were manually labelled by EEG experts and then cancelled. However, it is worth to point out that, since the algorithm processes the EEG window by window and then it averages the estimated parameters over the windows, the effect of possible artifacts is mitigated. The average time length of the recordings, after artifact cancellation, is 5.44min.

3. A novel, EEG-based, indirect measure of brain connectivity: Permutation Disalignment Index

PE was introduced by Bandt and Pompe in 2002³¹ as a symbolic robust descriptor which detects dynamic complexity changes in time series. Thanks to the projection into symbols (motifs), PE allows to estimate the randomness of a time series regardless of its amplitude, which plays a key role when analysing EEG, whose amplitude depends on the location of the reference electrode. In fact, when processing the EEG recordings through amplitude dependent techniques, each EEG signal should be normalised to cancel the effect of closeness to the reference electrode. Normalisation is not necessary when a symbolic procedure like PE is used. However, PE is a univariate descriptor which can only describe the randomness of a single time series (i.e. a signal recorded at a specific electrode site) and cannot quantify the coupling between two or more time series (i.e. between two or more different cortical areas). We will now briefly recall the PE definition in order to properly introduce our PDI concept.

3.1. Definition of Permutation Entropy

Given a time series x with N samples, it can be mapped into a m -dimensional space, with m being the embedding dimension and L being the time lag. Starting from a given sample $x(t)$ in the EEG window under analysis, a m -dimensional X_t vector is constructed, picking up the remaining $m - 1$ samples with a L -samples shift forward:

$$X_t = [x(t), x(t + L), \dots, x(t + (m - 1)L)]^T \quad (1)$$

Then the algorithm moves to the next sample $x(t + 1)$ and reiterates the procedure, so a new vector $X_{t + 1}$ is constructed.

The values of X_t are ordered, in increasing order and a reshaped version Xr_t of the original X_t is defined:

$$Xr_t = [x(\tau_1), x(\tau_2), \dots, x(\tau_m)]^T \quad (2)$$

With $\tau_i = 1, \dots, m$, where τ_i tells us is what position the sample $x(\tau_i)$ was in the original vector X_t .

By reshaping the vectors (and then considering the original time points associated to the elements of the reshaped vectors) we are essentially discarding the amplitude of the components and only taking into account their relative levels. We now need to associate the observed pattern to a predetermined symbol (motif). The sequence of the time points will tell us what motif the observed pattern can be associated to. In other words, each vector X_t can be considered mapped onto a symbol vector $\pi_i = [\tau_1, \tau_2, \dots, \tau_m]$ (where $i=1, \dots, m!$ and $m!$ is the number of the possible permutations); π_i is a sequence of time points, therefore a sequence of integers. The occurrence rate of a given sequence π_i in the time series x is denoted as $p_X(\pi_i)$, and represents the probability of observing the specific symbol π_i in the time series under analysis.

The algorithm counts how many times each sequence π_i appeared in the time series, then this number is normalised by $N - (m - 1)L$ to estimate the probability of the motif: $p(\pi_i) = n(\pi_i)/(N - (m - 1)L)$. PE is finally computed as:

$$H(m) = - \sum_{i=1}^{m!} p(\pi_i) \log(p(\pi_i)) \quad (3)$$

Where \log is the natural logarithm. The authors have recently successfully applied a new version of PE, based on the concept of Renyi's Entropy, i.e. the Renyi's Permutation Entropy (PER)³², to the study the EEG of absence seizure patients³³. In particular, the randomness of the electrical activity of each cortical area was investigated through the estimation of the randomness of the signal recorded at the corresponding EEG channel (time series x). In that work, Renyi's theory³⁴ was introduced to enhance the standard definition of PE through the introduction of the parameter α ($\alpha \geq 0$):

$$H_R(m) = - \frac{1}{1 - \alpha} \log \sum_{i=1}^{m!} p(\pi_i)^\alpha \quad (4)$$

For $\alpha \rightarrow 1$, we obtain the standard PE definition. High α values emphasise the super-gaussian distributions whereas low α values emphasise the sub-gaussian ones. Therefore, α can tune the sensitivity to sub-gaussianity and super-gaussianity of pdf distribution. In particular, higher α values ($\alpha > 4$) emphasise the super-gaussianity, whereas average alpha values ($\alpha = 2 \sim 3$) emphasise both of them.

3.2. Permutation Disalignment Index

The aim of the present work was to define a new measure of coupling strength between time series and then to test it an indirect, EEG-based, measure of brain connectivity in AD and MCI patients. As AD and MCI are known as disconnection disorders, we considered the coupling strength between the different cortical areas rather than the activity of the single cortical area. Furthermore, we were interested in symbolic descriptors, to ensure the independency from the reference-electrode. Summarising, we needed a symbolic, at least bivariate, descriptor of the coupling strength between EEG time series.

Hereby we propose a symbolic measure of coupling strength, referred to as *Permutation Disalignment Index* (PDI). In order to properly introduce PDI, we must refer to Section 3.1.

Our method is concisely illustrated in Figure 2. Let us, for example, consider $m=3$, for sake of clarity. When $m=3$, given two time series x and y , the time point t and the lag L , we can project the time series x into the vector X_t (Eq. 1) and the time series y into the vector Y_t (Eq. 5):

$$Y_t = [y(t), y(t + L), \dots, y(t + (m - 1)L)]^T \quad (5)$$

X_t and Y_t are both vectors with three elements because $m=3$. The values of X_t and of Y_t are reshaped in an increasing order, as described in the previous Section.

As explained in Section 3.1, by reshaping X_t and Y_t , and considering the original time points associated to the elements of the reshaped vectors, we discard the amplitude of the components and we only take into account their relative levels, which can be *low*, *medium* and *high* (when $m=3$). Six possible motifs can occur when $m=3$, in other words, six possible permutations of the *low*, *medium* and *high* levels (Figure 2). Given a time point t , we have to check if the same motif occurred both in x and y or not. For example, in Figure 2, at time t the same motif π_4 occurred both in x and y . The procedure is reiterated for every time point t , in this way, we eventually come up with a final simultaneous occurrence rate $p_{X,Y}(\pi_i)$ of every motif π_i in x and y . Once the occurrence rate $p_{X,Y}(\pi_i)$ of every motif has been estimated, we can define the PDI between x and y as:

$$PDI(X, Y) = \frac{1}{1 - \alpha} \log \left[\sum_{i=1}^{m!} p_{X,Y}(\pi_i)^\alpha \right] \quad (6)$$

The more coupled x and y are, the lower the PDI is expected to be, as we expect the time series to show the same motifs with a high probability. In this work, PDI

has been defined according to the Renyi's theory and the parameter α is introduced alongside m and L , because of the reasons discussed in Section 3.1.

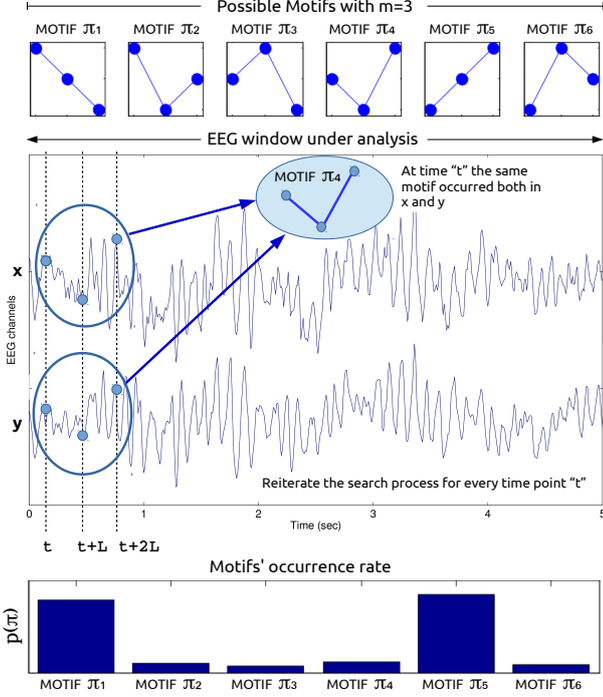


Fig. 2. Given two time series x and y (for example, two EEG signals within an EEG window under analysis), given an embedding dimension m , a time point t and a lag L , we can project the time series x into the vector X_t and the time series y into the vector Y_t , which are both vectors with three elements. Six possible motifs (permutations of the *low*, *medium* and *high* levels) can occur when $m=3$. Given a time point t , the algorithm checks if the same motif occurred both in x and y or not. In the present Figure, for example, at time t the same motif π_4 occurred both in x and y . The procedure is reiterated for every time point t so that, in the end of the analysis, we come up with a final occurrence rate $p_{X,Y}(\pi_i)$ of every motif π_i in x and y .

PDI can be extended to the analysis of a n -dimensional random variable X to estimate the overall coupling among all of its n components:

$$PDI(X_1, \dots, X_n) = \frac{1}{1 - \alpha} \log \left[\sum_{i=1}^{m!} p_{X_1, \dots, X_n}(\pi_i)^\alpha \right] \quad (7)$$

where the simultaneous occurrence of a given motif π_i in all of the time series X_1, \dots, X_n will be investigated, where $i = 1, \dots, m!$.

3.3. Henon Maps analysis with PDI

In this work we aimed at introducing PDI as a novel measure of coupling strength between time series and

testing its ability in estimating the coupling strength between real EEG signals from MCI and AD patients. First of all, we tested PDI on simulated data in order to assess its sensitivity to the changes in the coupling strength between interacting dynamic systems. The analysis of the simulated time-series yielded suggestions on the selection of the relevant configuration parameters for the computation of PDI. In particular, we applied PDI to the detection of nonlinear interdependency of two unidirectionally coupled Henon Maps X and Y , which have been extensively used in literature to validate measures of coupling strength^{35 36}, they are defined as:

$$\begin{cases} X : x_{n+1} = 1.4 - x_n^2 + b_x x_{n-1} \\ Y : y_{n+1} = 1.4 - [c x_n + (1 - c) y_n] y_n + b_y y_{n-1} \end{cases} \quad (8)$$

System X drives system Y with a nonlinear coupling strength c , which ranges from 0 to 1 with, 0 representing no coupling and 1 representing complete coupling. In the analysis of Identical Systems, b_x is set to 0.3 and b_y is set to 0.3, whereas they are set to $b_x = 0.3$ and $b_y = 0.1$ for Non Identical Systems. A detailed discussion about the implications of identical and nonidentical systems can be found in³⁷.

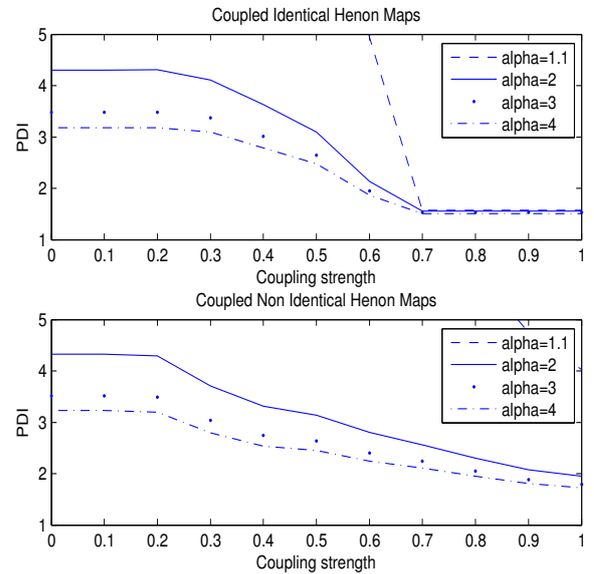


Fig. 3. PDI as a function of coupling strength c and α . The top sub-plot is related to the analysis of identical Henon Maps and the bottom sub-plot is related to non-identical Henon maps.

X and Y are initialised randomly and are iteratively computed making c increase of 0.01 every 1000 steps. A sample of PDI is computed every 1000 simulated samples of X and Y . Therefore, in the range $c \div c + 0.1$, 10 PDI values are calculated because c increases with a

0.01 increasing rate, in order to make it grow smoothly. The ten PDI values estimated with a coupling strength ranging from a given value c to $c+0.1$ are then averaged and a single, average, PDI sample is associated to that c value.

In Figure 3, PDI is depicted as a function of both the coupling strength c and α , for identical (top sub-plot) and non-identical Henon maps (bottom sub-plot). PDI decreases as the coupling strength increases. This suggests that decreased projected alignment randomness reflects an increased coupling strength, as expected. The behaviour is similar for different α values, even though the absolute range decreases as α increases. $\alpha=1.1$ shows the widest range but a non-smooth behaviour, therefore, in this work $\alpha=2$ was selected. PDI for $c > 0.7$ reflects a strong synchronisation between the two coupled systems. The critical threshold $c = 0.7$ corresponds to the point when the maximum Lyapunov exponent of the response system becomes negative and identical synchronisation between the systems takes place. For $0 < c < 0.7$, PDI decreases monotonically as c increases, thus showing that PDI is sensitive even to weak coupling.

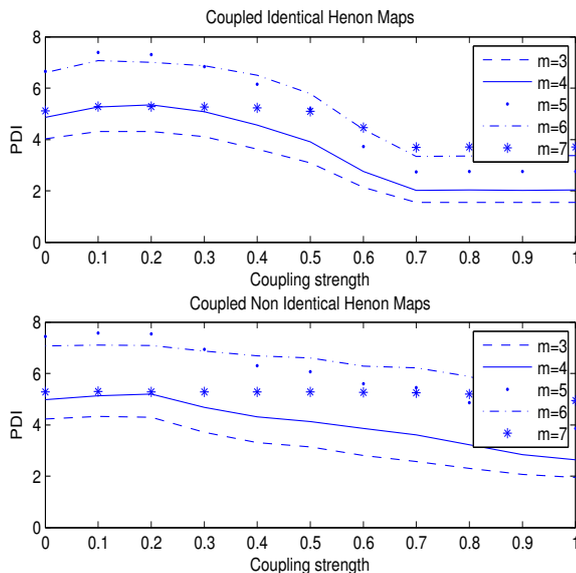


Fig. 4. PDI as a function of coupling strength c and embedding dimension m . The top sub-plot is related to the analysis of identical Henon Maps and the bottom sub-plot is related to non-identical Henon maps.

Figure 4 shows PDI as a function of the coupling strength c and the embedding dimension m , for both the identical (top sub-plot) and the non-identical Henon maps (bottom sub-plot). PDI decreases as the coupling strength increases.

With regard to the identical systems, for $c > 0.6$, PDI increases with m , which is intuitively plausible be-

cause the projected alignment randomness of two coupled systems becomes inherently lower, as m increases, because the probability of observing the same symbol in the two time series is lower (in fact, the number of possible symbols is equal to $m!$). An abrupt change can be observed in the transition from $m=6$ to $m=7$ as the trend appears flattened both in identical and non identical Systems. This suggests that, as m becomes too large, PDI becomes less sensitive to the coupling strength variation.

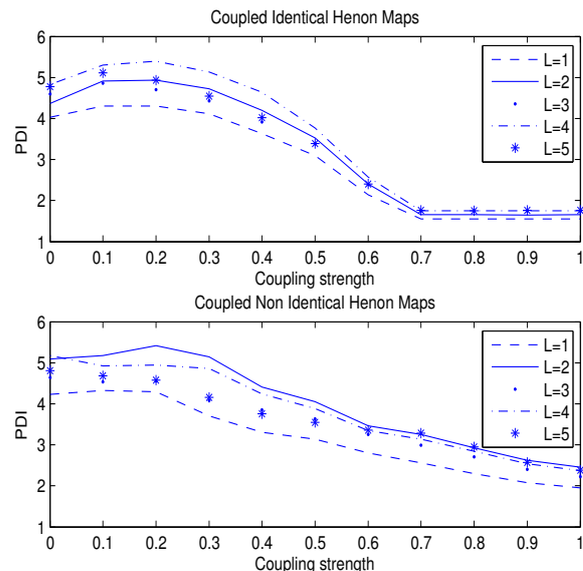


Fig. 5. PDI as a function of coupling strength c and lag L . The top sub-plot is related to the analysis of identical Henon Maps and the bottom sub-plot is related to non-identical Henon maps.

A relatively high m ($m > 3$), would inherently reduce the probability of a motif being detected both in signals x and y and, therefore, reduce the estimated synchronisation between them. In fact, as shown in 4 for the simulated Henon Maps, as m increases, the sensitivity to the coupling strength c is lost.

Furthermore, increasing m would increase the computational cost because the number of possible symbols, which we have to estimate the occurrence rate for, is equal to $m!$ According to the above mentioned observations, $m=3$ was selected in this paper.

Finally, Figure 5 shows PDI as a function of the coupling strength c and the lag L . Once again, PDI decreases as the coupling strength increases. The behaviour looks similar for different L values. The issue of optimal selection of m and L in AD/MCI EEG was discussed in detail in ³⁸. That paper showed that, by setting $m = 3$ and $L = 1$, the *slowing* effect typical of MCI/AD EEGs was better captured. Therefore, even though $L = 4$ looks associated to a wider range (Figure

5), since it showed the same trend as for $L = 1$, we selected $L = 1$. Our future work will be focused also on testing the method under different parameter settings, in order to assess how this affects the performance of the algorithm. This will be likely done on our upcoming, extended database, so that the parameter settings can be statistically optimised for MCI and AD patients. Until then, we would rely on the parameter settings $m = 3$ and $L = 1$, which was shown to work fine on AD/MCI EEG³⁸ and also worked fine on our theoretical assessment on Henon Maps.

Once proven the ability of PDI to follow the variation of the coupling strength between two simulated dynamical interacting systems, the next step was testing PDI in the quantification of the coupling strength between real systems like the electroencephalographic signals of AD and MCI patients. The goal is the indirect quantification of the connectivity between the cortical areas through the estimation of the coupling strength between the corresponding EEG signals.

4. Relative Power Analysis

As the Relative Power is a parameter commonly evaluated when analysing EEG from MCI and AD patients, even though it is a univariate measure and cannot directly be compared to PDI, we thought that showing a Relative Power analysis over our longitudinal database was appropriate. Given an EEG window under analysis and considering the generic x -th EEG channel, the corresponding time series is here denoted as x . The Power Spectral Density (PSD) of x is defined as the Fourier Transform of the autocorrelation function (ACF) of x :

$$PSD_{xx} = F\{ACF_{xx}(t)\} \quad (9)$$

Once each EEG channel was band-pass filtered as described in Section 2.3, and the four sub-bands-channels were generated ($x_\delta, x_\theta, x_\alpha, x_\beta$), we can compute the Relative Power (RP) of x in every sub-band (SB):

$$RP_x = \frac{PSD(SB)}{PSD(total)} \quad (10)$$

which is the ratio between the PSD of the signal in the specific sub-band ($f_1 - f_2$) and the overall PSD of in the range 0.1-32Hz. The power in the sub-band $f_1 - f_2$ was calculated as the integral of the PSD $P_{xx}(f)$, estimated between f_1 and f_2 . In this work, the popular Welch algorithm was used to estimate PSD³⁹.

Given an EEG_{SB} and given the generic window w under analysis, the relative power $RP_{x_1, \dots, x_k}^w(SB)$

among the electrodes belonging to a specific scalp area (frontal, temporal, central, parietal, occipital) was calculated by averaging the EEG of the electrodes x_1, \dots, x_k belonging to the sub-area under consideration and then calculating the RP over the averaged EEG. The $RP_{x_1, \dots, x_k}^w(SB)$ values were then averaged over the windows w and a single average RP value $\overline{RP}_{x_1, \dots, x_k}(SB)$ was computed for every sub-area, in every sub-band.

The average RP $\overline{RP}_{x_1, \dots, x_k}(SB)$ values were calculated both at time T0 and T1.

For every patient, the percent variation of $\overline{RP}_{x_1, \dots, x_k}(SB)$ (hereinafter simply denoted as \overline{RP}) was then estimated as:

$$\Delta(\overline{RP}(T1) - \overline{RP}(T0))\% = \frac{(\overline{RP}(T1) - \overline{RP}(T0))}{\overline{RP}(T0)} * 100 \quad (11)$$

5. Descriptors compared to PDI

5.1. Spectral Coherence

The magnitude squared coherence between two EEG channels x and y is defined as:

$$C_{xy}(f) = \frac{|P_{xy}(f)|^2}{P_{xx}(f)P_{yy}(f)} \quad (12)$$

where $P_{xy}(f)$ is the cross power spectral density of x and y , $P_{xx}(f)$ and $P_{yy}(f)$ are the power spectral densities of x and y , respectively. All of them are functions of the frequency f . $C_{xy}(f)$ is a function of frequency as well, and it is a measure of synchronisation between x and y . Coherence ranges from 0 to 1, which indicates how well x corresponds to y at a given frequency f . In this work, coherence was calculated through the Welch's averaged, modified periodogram method³⁹.

Given the sub-band EEG recording, EEG_{SB} , and given the window w under analysis ($w=1, \dots, Nw$, where Nw is the number of analysed windows), the average coherence $C_{x,y}^w(SB)$ between a given couple of electrodes x and y is calculated. The values $C_{x,y}^w(SB)$ are then averaged with respect to w , in order to come up with a single average $\overline{C}_{x,y}(SB)$ between channels x and y , for each EEG-sub-band recording.

5.2. Dissimilarity Index (Dm)

Given an EEG segment and given a pair of EEG channels (i. e. a pair of time series) x and y , they are projected into the phase space to reconstruct the ordinal patterns π_i ,

similarly to the procedure used in PE and PDI calculation (Sections 3.1 and 3.2).

The occurrence rates $p_X(\pi_i)$ and $p_Y(\pi_i)$ of each ordinal pattern π_i on each of the two time series x and y are calculated. After that, the distance between the rank-frequency distributions is estimated and represents the dissimilarity measure between the two time series:

$$Dm(X, Y) = \sqrt{\frac{m!}{m! - 1} \left(\sum_{i=1}^m (p_X(\pi_i) - p_Y(\pi_i))^2 \right)} \quad (13)$$

Further details can be found in Ouyang et al.²⁸. We used the algorithm shared by Ouyang et al. through the *File Exchange - MATLAB Central - MathWorks* website. The parameters m and L were set as for PDI.

5.3. Estimating PDI, Coherence and Dm from the EEG recordings

In order to carry out an overall scalp analysis, PDI, Coherence and Dm , between every possible pair of electrodes were computed. The aim is to evaluate the behaviour of the above mentioned descriptors, all over the scalp, in the follow-up of MCI and AD patients (at time T0 and at time T1), in every sub-band. Considering a sub-band EEG_{SB} , obtained as described in Section 2.3, and given a generic window w under analysis, the $PDI_{x,y}^w(SB)$, the coherence $C_{x,y}^w(SB)$ and the $Dm_{x,y}^w(SB)$ between every couple of electrodes x and y were calculated according to Sections 5.1 and 5.2. These values were then averaged over the windows (i. e. over the time) in order to come up with a single average PDI value $\overline{PDI}_{x,y}(SB)$, a single average coherence value $\overline{C}_{x,y}(SB)$ and a single average Dm value $\overline{Dm}_{x,y}(SB)$, for every couple, in every sub-band. For every patient, either EEG-T0 and EEG-T1, were analysed in this way.

6. Results

In this Section, we will first of all report the results of the Relative Power analysis (described in Section 4) and then we will report in detail the results of the analysis described in Section 5.

6.1. Relative Power analysis

The estimated percent variations of RP (Eq. 4) are represented as histograms in Figure 6. This analysis was carried out to detect possible shared trends in the changes of the EEG of MCI and AD patients from the point of view of RP variation. The top sub-plots, in Figure 6, are

associated to the MCI patients and the bottom ones are associated to the AD patients. Each sub-plot is associated to a specific sub-band (delta, theta, alpha and beta) and the x-axis represents the scalp sub-areas (Frontal, Temporal, Central, Parietal and Occipital). Within each group, every patient is depicted with a different colour so that his/her histograms can be identified. Within each group (AD or MCI), the patients showed some common behaviour, which can be summarised as follows:

- The temporal and parietal areas of MCI patients showed decreased RP in alpha band;
- The central and parietal areas of MCI patients showed decreased RP in beta band;
- The central area of AD patients showed increased RP in delta band;
- The frontal area of AD patients showed decreased RP in alpha band.

6.2. PDI vs Coherence vs Dm analysis

The main goal of the present paper was to develop a new indirect measure of connectivity and to apply it to the EEG of MCI and AD patients in order to investigate possible correlations with the disease's development. In other words, we aimed at finding possible EEG markers that are able to discriminate between stable MCI and MCI degenerating towards AD. We decided to examine in detail the overall scalp variation of PDI levels in the transition from T0 to T1, and to compare it with Coherence and Dm .

Figures 7, 8, 9 show the boxplots of PDI, Coherence and Dm , respectively. All of the three descriptors were estimated at time T0 and T1, according to the procedure described in Section 5. Boxplots are coloured according to the results provided by the statistical analysis discussed in Section 6.3.

Every boxplot in Figure 7, shows the $\overline{PDI}_{x,y}(SB)$ values calculated for every couple of electrodes x and y at time T0 or T1. Given a patient under consideration, the boxplot on the left is associated to time T0 and the one on the right is associated to time T1. Similarly, the single boxplot in Figure 8 and 9 shows the $\overline{C}_{x,y}(SB)$ values (Figure 8) and the $\overline{Dm}_{x,y}(SB)$ values (Figure 9), calculated for every couple of electrodes x and y at time T0 or T1. Inspecting the boxplots in the three Figures, we can infer that the patients who experienced the conversion from MCI to AD (pt 3, pt 51 and pt 71) showed a similar behaviour from the point of view of PDI's evolution and Coherence's evolution. In particular, we can point out that, in the above mentioned patients, the median of the Coherence decreased in the delta and theta bands, whereas the median of the

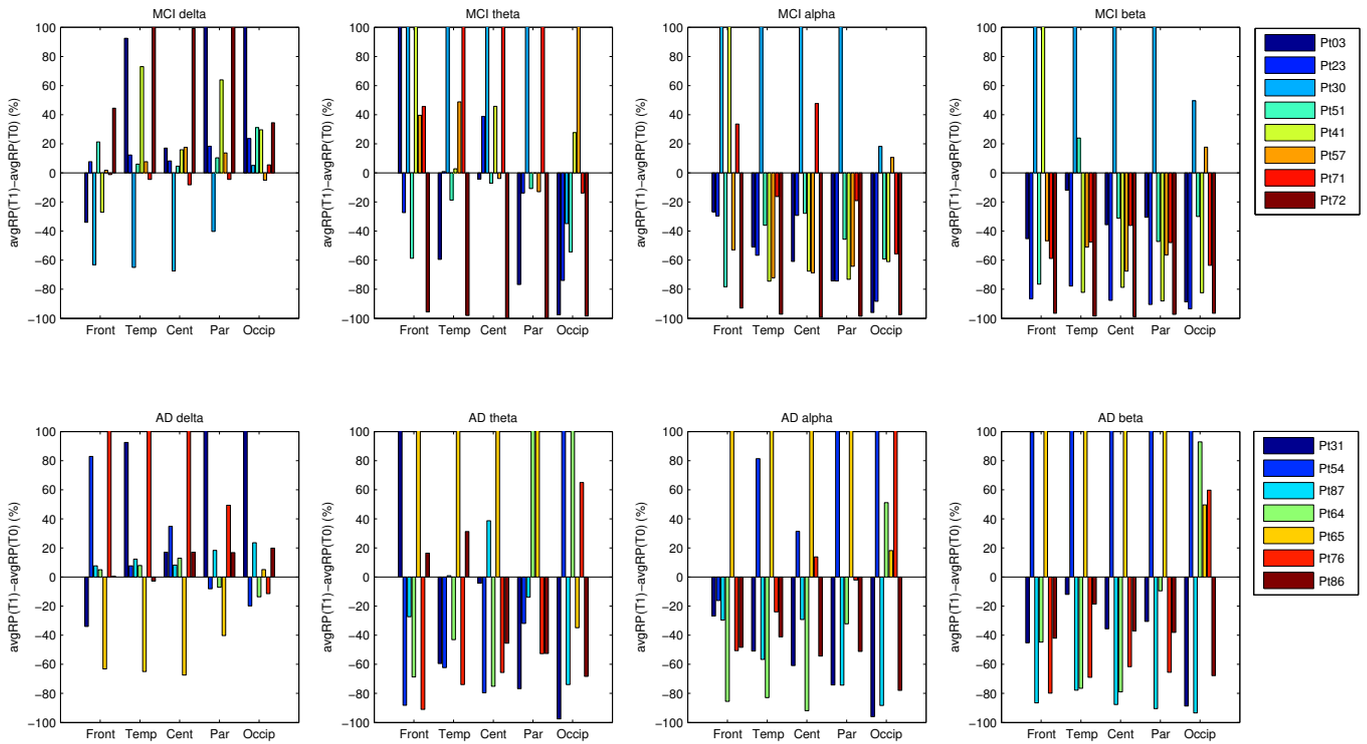


Fig. 6. Percent variation of the average RP (comparing time T0 and time T1). The top sub-plots are associated to the MCI patients and the bottom ones are associated to the AD patients. Within each group, the legend shows the colour associated to each patient. The x-axis represents the scalp sub-areas (Frontal, Temporal, Central, Parietal and Occipital) and the y-axis represents the percent variation of the average RP. Each sub-plot is associated to a specific sub-band (delta, theta, alpha and beta).

PDI increased in the delta and theta bands, thus, both descriptors reflected an overall reduced coupling strength in delta and theta band. Such a result is consistent with the commonly shared interpretation of AD and MCI as disconnection disorders. Boxplots in Figure 7 show that the increase of PDI is sharp in patient 3, 51 and 71 (who converted to AD), both in terms of increased median and absolute range. In particular, for patients 3, 51 and 71, the PDI boxplot range at T0 shows no overlap with the PDI boxplot range at T1 in delta band. The range of PDI in patients 51 and 71 showed no overlap also in theta and alpha sub-bands.

Four out of five stable MCI patients (pt 23, pt 41, pt 57, pt 72) showed stable or increased Coherence levels (i.e. stable or increased coupling strength), Figure 8, whereas pt 30 showed reduced Coherence levels, like the converted patients, and should be considered a false positive. On the contrary, from the PDI perspective, the totality of the stable MCI patients (pt 23, pt 30, pt 41, pt 57, pt 72) showed stable or reduced PDI (i.e. stable or increased coupling strength).

As regards Dm analysis, pt 23, pt 51 and pt 71 exhibited increased Dm levels (reduced coupling strength) in delta, alpha and beta band (Figure 9); pt 57 showed the same behaviour only in delta band and pt 72 only in theta band. Apparently, Dm seems to be not sensitive to MCI to AD conversion as there is no correlation between DM variation and disease progression towards AD. We can conclude that MCI patients who converted to AD exhibited:

- Decreased median Coherence in delta and theta bands (not specific because also one stable MCI patient, pt 30, exhibited a similar behaviour in delta band);
- Increased median PDI and increased PDI range in delta and theta bands (specific and sharp, particularly in delta band);

Summing all up, even though the analysed dataset is small, both Coherence and PDI resulted sensitive to the conversion from MCI to AD because both detected a reduced coupling strength in delta and theta bands. Nevertheless, Coherence looks less specific than PDI be-

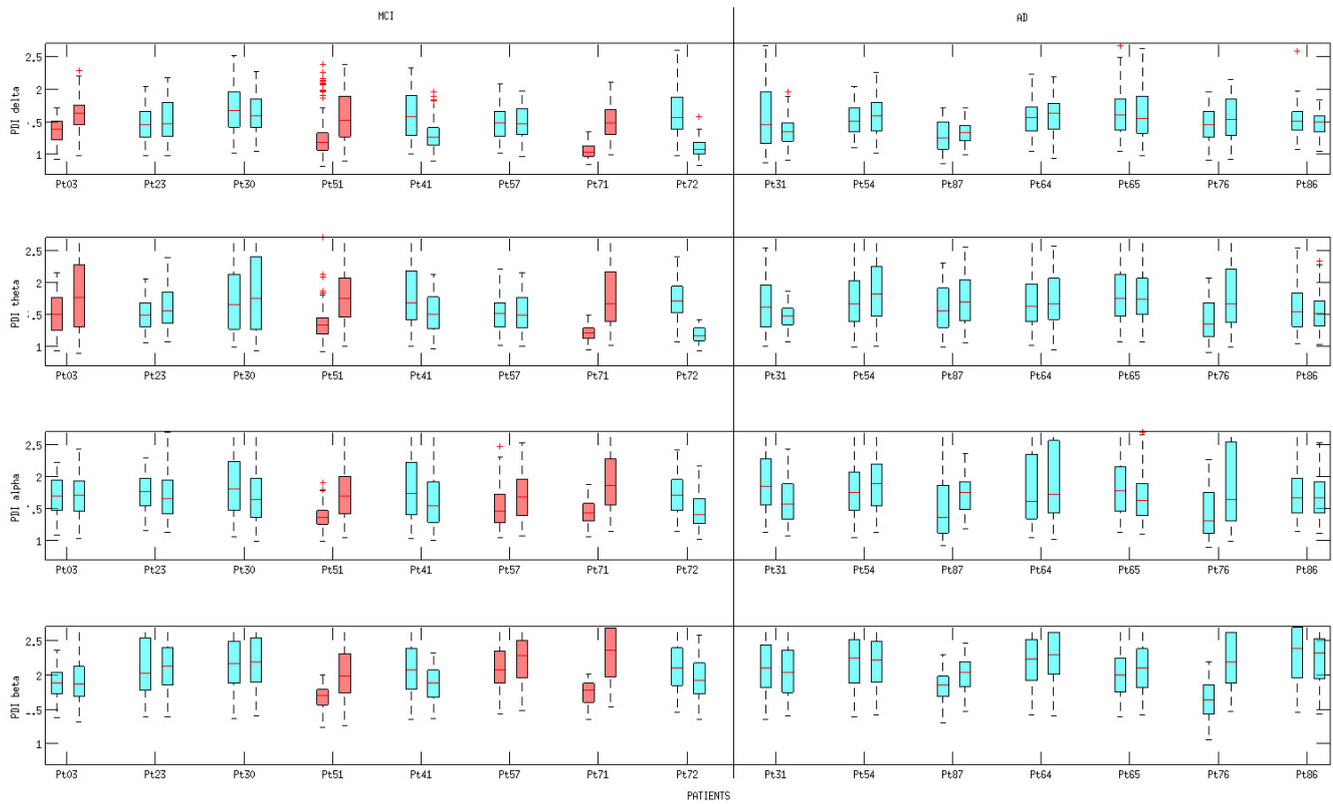


Fig. 7. Boxplot of PDI at time T0 and T1. Every row is associated to a specific EEG sub-band (delta, theta, alpha, beta). For each patient, the boxplot on the left represents the values assumed at time T0 whereas the boxplot on the right represents the values assumed at time T1. On each box, the central mark is the median, the edges of the box are the 25th and 75th percentiles, the whiskers extend to the most extreme data points not considered outliers. The red boxplots, in the MCI group, are associated to a statistically significant increase ($p < 0.001$) of PDI levels (see Section 6.3 Statistical Analysis.)

cause one stable MCI patient (patient 30) behaved like converted patients (patient 3, 51 and 71) as he/she exhibited a decreased coherence (Figure 8) in delta band, although at T1 he/she was still diagnosed MCI. From the PDI perspective, the reduced coupling strength in delta and theta bands was a behaviour that could be detected only on converted MCI patients.

6.3. Statistical analysis

In order to assess how significant was, for every patient, the overall variation of the PDI levels, between time T0 and T1, an intra-subject, descriptive analysis was performed. For every patient, given a sub-band, the two populations under analysis were the PDI values at T0 (given a patient, the boxplot on the left in Figure 7) and the PDI values at T1 (given a patient, the boxplot on the right in Figure 7). The same analysis was carried out with Coherence (Figure 8) and with Dm (Figure 9).

The Wilcoxon rank sum test⁴⁰ was used to test the null hypothesis that data in the two populations under consideration are independent samples from identical continuous distributions with equal medians. The statis-

tical analyses were performed in Matlab.

The achieved results are reported in Table II. The p values corresponding to a significant ($p < 0.001$) increase of PDI or Dm levels (reduced coupling strength) or with significant decrease of Coherence levels (reduced coupling strength), are highlighted in bold style. In Figures 7, 8 and 9, the boxplots associated with significant ($p < 0.001$) increase of PDI or Dm levels or with significant decrease of Coherence levels, were coloured in red in MCI patients, in order to unravel possible correlations with MCI progression towards AD.

As regards PDI analysis, the intrasubject variability study showed significantly ($p < 0.001$) increased PDI levels in: Patient 3 (delta, theta); Patient 51 (delta, theta, alpha, beta); Patient 57 (alpha, beta); Patient 71 (delta, theta, alpha, beta); Patient 54 (theta, alpha); Patient 87 (delta, theta, alpha, beta); Patient 64 (alpha); Patient 76 (theta, alpha, beta).

As regards Coherence analysis, the intrasubject variability study showed a significant ($p < 0.001$) decrease of Coherence, between T0 and T1, in: Patient 3 (delta, theta); Patient 30 (delta); Patient 51 (delta, theta,

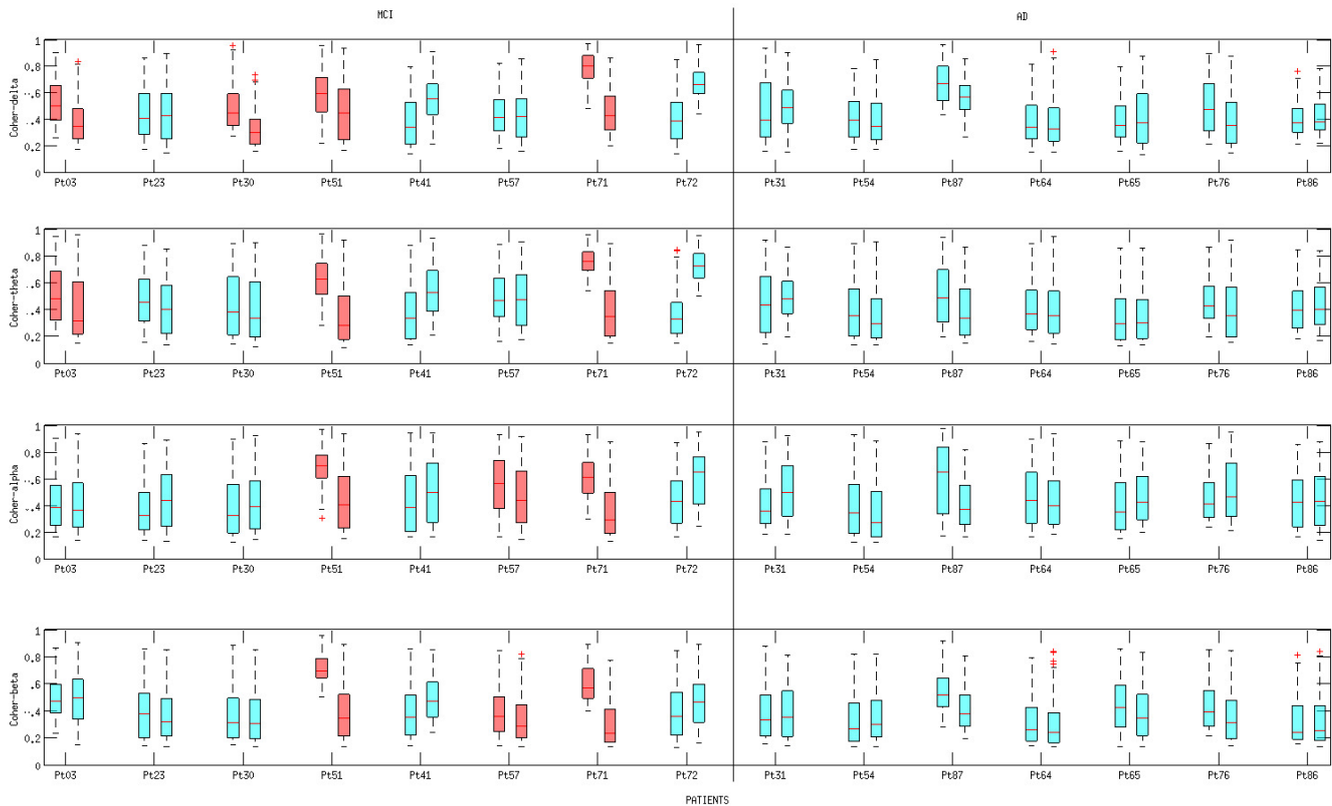


Fig. 8. Boxplot of Coherence at time T0 and T1. Every row is associated to a specific EEG sub-band (delta, theta, alpha, beta). For each patient, the boxplot on the left represents the values assumed at time T0 whereas the boxplot on the right represents the values assumed at time T1. On each box, the central mark is the median, the edges of the box are the 25th and 75th percentiles, the whiskers extend to the most extreme data points not considered outliers. The red boxplots, in the MCI group, are associated to a statistically significant decrease ($p < 0.001$) of Coherence levels (see Section 6.3 *Statistical Analysis*.)

alpha, beta); Patient 57 (alpha, beta); Patient 71 (delta, theta, alpha, beta); Patient 54 (theta, alpha); Patient 87 (delta, theta, alpha, beta); Patient 65 (beta); Patient 76 (delta, theta, beta).

As regards Dm analysis, the intrasubject variability study showed significantly ($p < 0.001$) increased Dm levels in: Patient 23 (delta, alpha and beta); Patient 51 (delta, alpha and beta); Patient 71 (delta, alpha and beta); Patient 57 (delta) and Patient 72 (theta).

Focusing on MCI patients, the intrasubject variability study showed that patients 3, 51 and 71 (who had converted to AD by the time T1) exhibited a significant ($p < 0.001$) increase of PDI in delta and theta bands and that patients 51 and 71 exhibited the same behaviour also in alpha and beta bands. As regards Coherence, patients 3, 51 and 71 exhibited a significant ($p < 0.001$) decrease of Coherence in delta and theta bands and patients 51 and 71 behaved similarly also in alpha and beta bands. However, also patient 30 (not converted to AD) exhibited a significant decrease of Coherence in delta band. As regards Dm analysis, it was not possible to detect a behaviour peculiar to the converted MCI.

Summing all up, a statistically significant increase of PDI medians was detected, in delta and theta bands, only in the MCI subjects who converted to AD. The average increase of PDI, in delta and theta bands, might be possibly correlated with the reduced connectivity, which is associated to the disease progression towards AD. The group analysis was not performed, due to the limited size of the dataset: this will be the goal of our future research. A large number of patients is, indeed, currently under recruitment at the *IRCCS Neurolesi Center Bonino-Pulejo* (Messina, Italy). In the near future, an extensive validation of the proposed methodology will be carried out on the extended dataset.

7. Discussion

In this paper, we introduced a novel measure of time series coupling, the *Permutation Disalignment Index* (PDI), and its possible application as an indirect estimation of the brain connectivity through the EEG, with application to the follow-up of AD and MCI patients. PDI is a multivariate measure which allows to estimate the

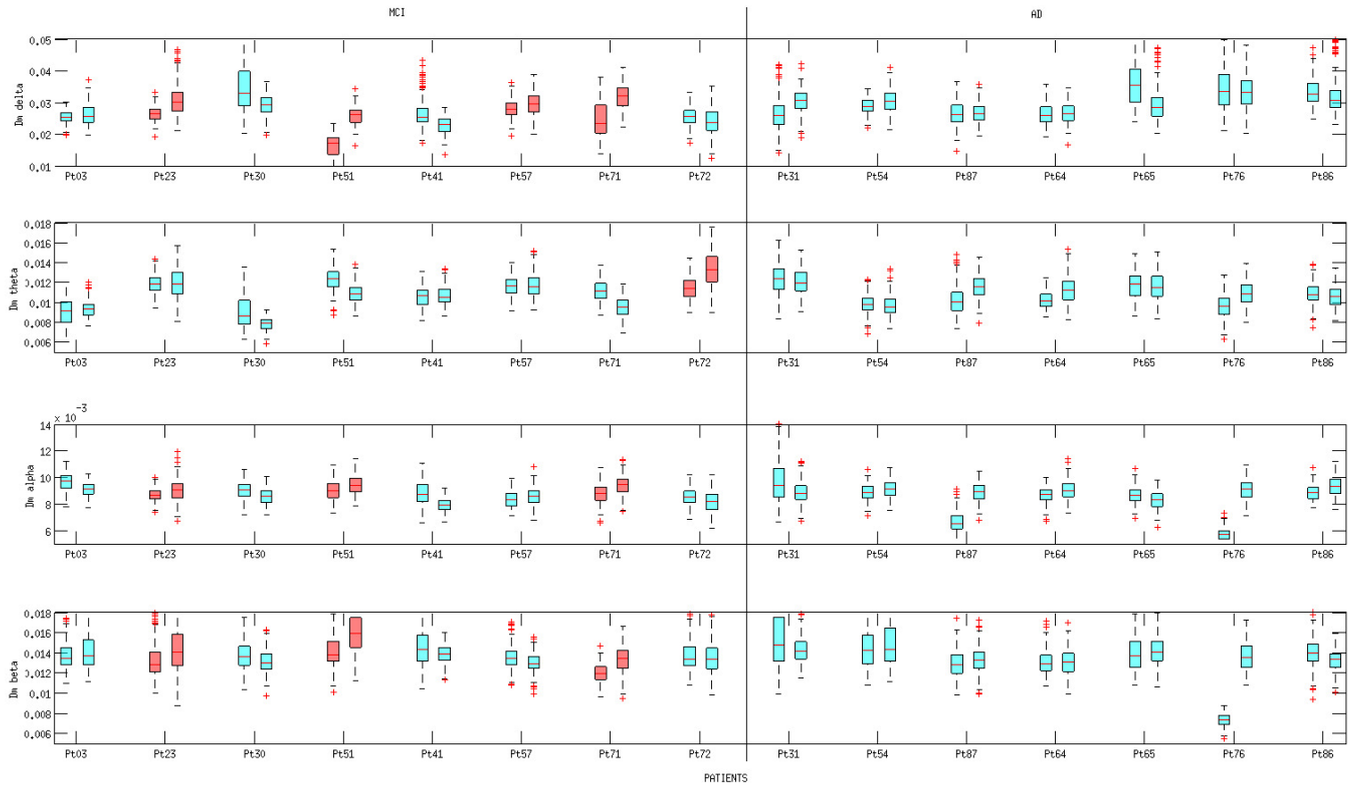


Fig. 9. Boxplot of Dm at time T0 and T1. Every row is associated to a specific EEG sub-band (delta, theta, alpha, beta). For each patient, the boxplot on the left represents the values assumed at time T0 whereas the boxplot on the right represents the values assumed at time T1. On each box, the central mark is the median, the edges of the box are the 25th and 75th percentiles, the whiskers extend to the most extreme data points not considered outliers. The red boxplots, in the MCI group, are associated to a statistically significant increase ($p < 0.001$) of Dm levels (see Section 6.3 *Statistical Analysis*.)

		PDI				Coherence				Dm			
		Delta	Theta	Alpha	Beta	Delta	Theta	Alpha	Beta	Delta	Theta	Alpha	Beta
MCI	Pt 03 (conv to AD)	2,86E-018	1,66E-005	0,81577343	0,44126899	1,90E-015	2,33E-006	0,44843899	0,711596705	0,112977469	0,021617034	5,71E-016	0,165451263
	Pt 23	0,145728533	0,00941136	0,03032865	0,34742543	0,95202795	0,00717482	0,00227497	0,164785168	5,46E-018	0,377389748	2,53E-005	6,653E-007
	Pt 30	0,133709567	0,24228364	0,00885711	0,60108012	0,00020795	0,24805842	0,09972153	0,714043459	2,94E-011	9,60E-012	1,19E-009	7,61E-005
	Pt 51 (conv to AD)	2,33E-015	1,12E-022	1,18E-019	5,10E-020	1,88E-007	4,61E-027	1,27E-022	2,75E-041	9,88E-056	1,86E-025	9,40E-008	2,99E-014
	Pt 41	1,33E-015	4,98E-007	4,65E-004	4,30E-008	2,15E-016	5,44E-013	6,92E-003	3,55E-008	2,03E-016	0,405800646	7,25E-023	0,000911609
	Pt 57	0,742800365	0,70508659	5,36E-006	0,00065882	0,49697015	0,40271885	9,05E-005	0,00674046	5,28E-005	0,937227236	0,001285828	1,55E-006
	Pt 71 (conv to AD)	5,04E-050	2,18E-034	3,06E-023	1,04E-034	9,60E-046	6,90E-042	2,38E-024	1,65E-035	1,37E-026	1,69E-032	1,71E-012	2,48E-025
	Pt 72	3,11E-047	4,42E-050	2,71E-013	5,53E-006	2,18E-036	3,03E-048	4,45E-013	2,15E-005	0,000160145	4,29E-023	0,000104076	0,223026449
AD	Pt 31	0,00423174	6,33E-005	5,47E-008	0,20410968	0,03748398	0,03448438	2,19E-006	0,897301284	2,00E-016	0,058887264	1,65E-006	0,006852411
	Pt 54	9,23E-002	0,00708134	0,02335987	6,89E-001	3,91E-001	6,22E-002	3,86E-002	0,009933283	2,18E-006	0,041789886	0,000318493	0,145728533
	Pt 87	0,006041909	0,0147187	4,17E-009	6,97E-012	2,43E-010	1,45E-007	2,27E-012	1,24E-014	0,562098917	9,95E-018	2,29E-051	0,007413375
	Pt 64	0,235306893	0,2344449	0,01981434	0,13976615	0,38512272	0,1664542	0,24316586	0,259895256	0,080855598	3,07E-013	2,41E-006	0,792079229
	Pt 65	0,688906413	0,75274547	0,00503846	0,08009788	0,89557052	0,88087989	0,00030665	0,00987057	5,40E-019	0,375618902	1,13E-007	0,016013818
	Pt 76	0,001583312	7,02E-012	4,49E-010	1,66E-031	1,29E-007	0,00026103	0,02691103	4,46E-006	0,137724263	2,67E-019	1,49E-057	1,46E-057
	Pt 86	4,53E-002	0,54162747	5,81E-001	2,60E-002	1,89E-001	5,98E-001	7,49E-001	0,653026532	0,000353685	0,016500072	1,58E-010	2,18E-007

Results of the statistical, intra-subject, descriptive analysis. We evaluated how significant was, for every patient, the overall variation of each descriptor (PDI, Coherence and Dm), between time T0 and T1. The Wilcoxon rank sum test was applied and the p values are reported. The p values corresponding to a significant ($p < 0.001$) increase of PDI or Dm levels (reduced coupling strength) or with significant decrease of Coherence levels (reduced coupling strength), are highlighted in bold style.

coupling strength between signals in terms of projected alignment randomness. PDI overcomes the limitations of the standard univariate Permutation Entropy (PE)³¹ whenever a pairwise or multivariate measure is necessary. Furthermore, it overcomes the issues that come with applying standard, amplitude dependent methodologies to EEG data. In fact, when processing the EEG,

the amplitude of the signals depends on the location of the reference electrode and, when using amplitude dependent methods, a preprocessing step to normalise EEG time series is required. The *permutation* concept allows us to bypass the normalisation as time series are projected into a symbolic space, where the effect of amplitude is lost and only the effect of variation is kept,

which also produces an improved robustness to noise and EEG artifacts in general. The ability of PDI in detecting nonlinear interdependency of two unidirectionally coupled Henon maps X and Y was validated theoretically. PDI was then tested in a follow-up study over 8 MCI and 7 AD patients. Every patient was evaluated at time T0 and then three months later (time T1): 5 MCI remained stable and 3 MCI exhibited a conversion to AD. The EEG recordings were processed, in each sub-band, in order to track the changes of the EEG's PDI in every range of frequency.

The study consisted in evaluating the evolution of PDI, all over the scalp, in comparison with Coherence and Dissimilarity Index. Coherence and PDI resulted both sensitive to the conversion from MCI to AD; in fact, MCI patients who converted to AD showed an increased median PDI and decreased median Coherence (reduced coupling strength) in delta and theta bands. However, only PDI resulted specific, because Coherence produced a false positive (a stable MCI subject exhibited a behaviour similar to the converted MCI). Analysing the EEG of MCI patients through PDI looks a promising way to quantify how the disease is affecting the EEG and to detect its possible degeneration towards AD. Developing EEG-based systems that are able to objectively quantify the progression of the disease is crucial and focusing the attention on periodic, long-term, follow-up studies is necessary. In this way, we could unravel if the variation of key EEG features may be the hallmark of the disease's progression and help to foresee its development. Furthermore, in the present paper, only resting state EEG were considered, future work will also be focused on the analysis of EEG during tasks, including evoked potentials. In this case, however, the traces will be more likely to be affected by artifacts and an advanced artifact rejection step will be required^{41, 42, 43, 44}.

8. Conclusion

Permutation Disalignment Index (PDI) was proposed in this paper as a new measure of time series coupling, with application to the indirect estimation of the brain connectivity in patients affected by AD and MCI. PDI was tested on simulated two unidirectionally coupled Henon maps time series in order to test its ability to measure the coupling between interacting dynamic systems. The simulation also yielded suggestions on how to tune the PDI parameters. PDI was then applied to real EEG traces recorded during a follow-up study carried out over 7 Alzheimer's Disease (AD) patients and 8 Mild Cognitive Impairment (MCI) subjects. Every pa-

tient was evaluated at time T0 and at time T1, 3 months later. At time T1, 5 out of 8 MCI patients were still diagnosed MCI, whereas the remaining 3 exhibited a conversion from MCI to AD. PDI was compared with the spectral coherence, a common parameter used in literature in the analysis of AD and MCI EEG and with the Dissimilarity Index, a recent symbolic measure of time series dissimilarity. The study consisted in evaluating the overall evolution of PDI, sub-band by sub-band. Both coherence and PDI resulted sensitive to the conversion from MCI to AD, but only PDI resulted specific as it allowed to detect a recurrent behaviour that was peculiar only to converted MCI patients, who indeed showed a significant increase of PDI in delta and theta bands ($p < 0.001$). In conclusion, the intra-subject, longitudinal evaluation of the overall coupling between the EEG signals, through PDI, looks a promising method to discriminate between converted MCI and stable MCI. Future efforts will be focused on extending the study to a larger number of patients that should be monitored periodically and over a long time. Furthermore, PDI will be tested, in conjunction with other EEG features, as a possible input parameter to a system designed for the estimation of the probability that a patient will convert to AD.

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