Relationship between EEG Alpha3/Alpha2 Ratio and the Nucleus Accumbens in Subjects with Mild Cognitive Impairment

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Abstract

Background: The nucleus accumbens (NAc) has been associated with impulsive behavior in subjects with early cognitive impairment; grey matter (GM) changes of basal ganglia have been demonstrated to be involved in Alzheimer’s disease (AD). Moreover, the increase of EEG alpha3/alpha2 ratio has been associated with AD-converters subjects with mild cognitive impairment (MCI).

Objective: To study the association of EEG marker with specific GM changes of NAc in subjects with MCI

Methods: 74 adult subjects with mild cognitive impairment underwent EEG recording and high resolution 3D magnetic resonance imaging (MRI). The alpha3/alpha2 ratio was computed for each subject. Three groups were obtained according to increasing tertiles values of alpha3/alpha2. Grey matter density differences between groups were investigated using a Voxel Based Morphometry technique.

Results: Subjects with lower a3/a2 and middle a3/a2 ratio showed greater grey matter reduction in the NAc bilaterally when compared to subjects with higher a3/a2 ratio.

Conclusion: The a3/a2 ratio was associated with increase of grey matter density inside the NAc in MCI subjects at major risk to develop AD.

Introduction

EEG have been demonstrated a reliable diagnostic tool in dementia research [1,2]. The increase of high alpha relative to low alpha power has been recently demonstrated a reliable EEG marker of hippocampal atrophy as well as conversion of patients with mild cognitive impairment (MCI) in Alzheimer’s disease (AD;[3]).

Impulsive behaviors are frequently described in brain-damaged patients, including patients with AD. More specifically, a significant increase in urgency, lack of premeditation, and lack of perseverance was noted, whereas a decrease in sensation seeking was observed in these patients [4]. Furthermore, this increase of impulsivity on urgency, lack of premeditation, and lack of perseverance was not associated with global cognitive impairment as assessed by the Mini Mental State Examination (MMSE; [5]) or the Mattis Dementia Rating scale [6], whereas lower sensation seeking was associated with a lower score on the MMSE [4,7].

The nucleus accumbens (NAc) is a key component of the neural processes regulating impulsivity [8]. Data from imaging studies [8] suggest the involvement of the NAc in impulsive choice. In particular an increase in the activity of NAc has been associated with decisions involving immediate outcome [9-11], with decisions including intertemporal differences [12], with choices of shorter delays [13] as well as with switching to risk-seeking choices [14].

In the present study the association of EEG indexes with grey matter (GM) changes in NAc has been studied in subjects with MCI. The working hypothesis was that modifications of alpha3/alpha2 power ratio could be underpinned by different deep brain structures. Results show that subjects with higher a3/a2 ratios when compared to subjects with lower and middle a3/a2 ratios showed minor atrophy in the NAc bilaterally. The results are discussed in the light of possible relationship with impulsive behavior in prodromal Alzheimer’s disease patients. The subjects in this study are the same of a group analyzed in a previous paper (Moretti et al., 2012) [15]. The novelty of the present work was that the attention was more focused on the role of the NAc volumetric changes in subjects with mild cognitive impairment. This choice was made considering the growing consideration of recent literature about the function of the NAC as an important hub in behavioral regulation. In this view, the statistical threshold of the analysis was strengthened (uncorrected threshold of p<0.001).

Materials and Methods

Subjects

Seventy-four subjects with MCI were recruited from the memory Clinic of the Scientific Institute for Research and Care (IRCCS) of Alzheimer’s and psychiatric diseases ‘Fatebenefratelli’ in Brescia, Italy. All experimental protocols had been approved by the local ethics committee. Informed consent was obtained from all participants or their caregivers, according to the Code of Ethics of the World Medical Association (Declaration of Helsinki).

Diagnostic criteria

Patients were selected from a prospective study on the natural history of cognitive impairment (the translational outpatient memory clinic—TOMC study) carried out in the outpatient facility of the National Institute for the Research and Care of Alzheimer’s Disease.
were signs of behavioural and/or EEG drowsiness.

positioned according to the 10–20 international systems (Fp1, Fp2, F7, 
deviation, number and [range].

Table 1: Demographic and cognitive characteristics in the whole sample, disaggregated for increased levels of Alpha3/Alpha2 ratio. Numbers denote mean ± standard deviation, number and [range]. *p denotes significance on ANOVA (continuous variables) and chi-square test (dichotomous variables).

|               | ALL      | High     | Middle   | Low      |
|---------------|----------|----------|----------|----------|
| Number of subjects | 74       | 18       | 38       | 18       |
| Age, years    | 69.4 ± 7.6 [52-85] | 70.4 ± 6.7 [60-85] | 66.8 ± 8.2 [52-83] | 70.4 ± 7.4 [57-80] |
| Sex, female   | 51 (%)   | 13 (%)   | 24 (%)   | 14 (%)   |
| Education, years | 7.6 ± 3.9 [3-18] | 6.6 ± 3.6 [4-18] | 7.6 ± 3.7 [3-17] | 8.3 ± 4.7 [3-18] |
| Mini mental state exam | 27.2 ± 1.7 [23-30] | 26.9 ± 1.3 [23-29] | 27 ± 1.7 [24-30] | 27.4 ± 1.2 [23-30] |
| Alpha3/alpha2 | 1.09 ± 0.15 [0.77-1.52] | 1.29 ± 0.14 [1.17-1.52] | 1.08 ± 0.0 * [1.1-1.16] | 0.9 ± 0.1 * [0.77-0.98] |

Analysis of individual frequency bands

All recordings were obtained in the morning with subjects resting comfortably. Vigilance was continuously monitored in order to avoid drowsiness. A digital FFT-based power spectrum analysis (Welch technique, Hanning windowing function, no phase shift) computed – ranging from 2 to 45 Hz – the power density of EEG rhythms with a 0.5 Hz frequency resolution. Two anchor frequencies were selected according to the literature guidelines [30,31], that is, the theta/alpha transition frequency (TF) and the individual alpha frequency (IAF) peak. These anchor frequencies were computed on the power spectra averaged across all recording electrodes. The TF marks the transition frequency between the theta and alpha bands, and represents an estimate of the frequency at which the theta and alpha spectra intersect. TF was computed as the minimum power in the alpha frequency range, since our EEG recordings were performed at rest. The IAF represents the frequency with the maximum power peak within the extended alpha range (5–14 Hz). Based on TF and IAF, we estimated the frequency band range for each subject, as follows: delta from TF-4 to TF-2, theta from TF-2 to TF, low alpha band (alpha1 and alpha2) from TF to IAF, and high alpha band (or alpha3) from IAF to IAF + 2. The alpha1 and alpha2 bands were computed for each subject as follows: alpha1 from TF to the middle point of the TF-IAF range, and alpha2 from such middle point to the IAF peak [22-28]. The mean frequency range computed in MCI subjects considered as a whole are: delta 2.9–4.9 Hz; theta 4.9–6.9 Hz; alpha1 6.9–8.9 Hz; alpha2 8.9–10.9 Hz; alpha3 10.9–12.9 Hz; beta1 12.9–19.2 Hz; beta2 19.2–32.4; gamma 32.4–45. The relative power density for each frequency band was computed as the ratio between the absolute power and the mean power spectra from 2 to 45 Hz. The relative band power at each band was defined as the mean of the relative band power for each frequency bin within

The project was aimed to study the natural history of non-demented persons with apparently primary cognitive deficits, i.e., deficits not due to psychic (anxiety, depression, etc.) or physical (hypothyroidism, vitamin B12 and folate deficiency, uncontrolled heart disease, uncontrolled conditions (diabetes, etc.) in the absence of functional impairment. The selection criteria had the aim to include as much as possible primary prodromal dementia due to neurodegenerative disorders. Demographic and cognitive features of the subjects in study are summarized in Table 1. Selection criteria were extensively described elsewhere [15]. Briefly, patients were rated with a series of standardized diagnostic and severity instruments, including the Mini-Mental State Examination (MMSE; [5]), the Clinical Dementia Rating Scale (CDRS; [16]; the Hachinski Ischemic Scale (HIS; [17]; and the Instrumental and Basic Activities of Daily Living (IADL, BADL; [18]). In addition, patients underwent diagnostic neuroimaging procedures (magnetic resonance imaging, MRI), and laboratory testing to rule out other causes of cognitive impairment. All patients underwent: (i) semi-structured interview with the patient and – whenever possible – with another informant (usually, the patient’s spouse or a child of the patient) by a geriatrician or neurologist; (ii) physical and neurological examinations; (iii) performance-based tests of physical function, gait and balance; (iv) neuropsychological battery assessing memory (Babcock Story Recall – Rey–Osterrieth Complex Figure, Recall – Auditory-Verbal Learning Test, immediate and delayed recall [19]) verbal and non-verbal memory, attention and executive functions (Trail Making Test B, A and B-A; Inverted Motor Learning-Clock Drawing Test; [19]; abstract reasoning thinking (Raven Colored Progressive Matrices; [19]), frontal functions (Inverted Motor Learning); language (Phonological and Semantic Fluency-Token test [19]), and apraxia and visuo-constructional abilities (Rey–Osterrieth Complex Figure, Rey figure copy, Clock Drawing Test; [19]); (v) assessment of depressive symptoms by means of the Center for Epidemiologic Studies Depression Scale (CES-D; [20]). All the neuropsychological tests were standardized on Italian population, thus scores were compared to normative values with age, education and gender corrections in an Italian population. All subjects were right-handed. As the aim of our study was to evaluate the relationship between GM loss and alpha2/alpha3 ratios in MCI subjects, we did not consider the clinical subtype of MCI, i.e., Amnesic MCI, Vascular MCI, Mixed MCI. All subjects were right-handed. As the aim of our study was to evaluate the relationship between GM loss and alpha2/alpha3 ratios in MCI subjects, we did not consider the clinical subtype of MCI, i.e., Amnesic MCI, Vascular MCI, Mixed MCI. All subjects were right-handed. As the aim of our study was to evaluate the relationship between GM loss and alpha2/alpha3 ratios in MCI subjects, we did not consider the clinical subtype of MCI, i.e., Amnesic MCI, Vascular MCI, Mixed MCI. All subjects were right-handed. As the aim of our study was to evaluate the relationship between GM loss and alpha2/alpha3 ratios in MCI subjects, we did not consider the clinical subtype of MCI, i.e., Amnesic MCI, Vascular MCI, Mixed MCI. All subjects were right-handed. As the aim of our study was to evaluate the relationship between GM loss and alpha2/alpha3 ratios in MCI subjects, we did not consider the clinical subtype of MCI, i.e., Amnesic MCI, Vascular MCI, Mixed MCI. All subjects were right-handed. As the aim of our study was to evaluate the relationship between GM loss and alpha2/alpha3 ratios in MCI subjects, we did not consider the clinical subtype of MCI, i.e., Amnesic MCI, Vascular MCI, Mixed MCI. All subjects were right-handed.
that band. The alpha3/alpha2 was computed in all subjects and three groups were obtained according to increasing tertiles values of alpha3/alpha2: low (a3/a2<1) middle (1< a3/a2<1.16) and high (a3/a2>1.17). The three groups of MCI have been demonstrated in previous studies to be different in nature. In particular, the high alpha3/alpha2 EEG power ratio MCI group is at major risk to convert to Alzheimer’s disease [29], as well as to have different pattern of hippocampal atrophy [28] and basal ganglia and thalamus gray matter lesions [29,15] as compared to the other alpha3/alpha2 power ratio MCI groups. Moreover, this group subdivision has been chosen for reason of homogeneity and comparability with the previous studies.

**MRI scans**

For each subject, a high-resolution sagittal T1 weighted volumetric MR scan was acquired by using a 1.0 T Phillips Gyroscan scanner, with a gradient echo 3D technique: TR=20 ms, TE=5 ms, flip angle=30°, field of view =220 mm, acquisition matrix 256 × 256, slice thickness 1.3 mm. The pattern of gray matter atrophy was studied using the Voxel Based Morphometry technique [32].

**Voxel-based morphometry**

3D images were processed through SPM5 software package (Statistical Parametric Mapping, Version 5; Wellcome Department of Imaging Neuroscience, London, UK; http://www.fil.ion.ucl.ac.uk/spm), running on Matlab 7.0.1 (Math-Works, Natick, MA, USA). DICOM files were converted in ANALYZE format image, the extra-cranial voxels were removed and the anterior commissure (AC) was manually set for all images as the origin of the spatial coordinates for an anatomical normalization algorithm implemented in SPM. Converted files were then segmented into gray and white matter and normalized to the GM population templates, generated from the complete image set, using the Diffeomorphic Anatomical Registration using Exponentiated Lie algebra (DARTEL) registration method [32]. This non-linear warping technique minimizes between-subject structural variations.

Spatially normalized images were modulated by the Jacobian determinants derived from the spatial normalization, to ensure that the overall amount of each tissue class was not altered by the spatial normalization procedure. The final voxel resolution after DARTEL was 1.5 × 1.5 × 1.5 mm. Finally each modulated, warped GM image was transformed to MNI space and smoothed with an 8 mm full-width at half-maximum (FWHM) Gaussian kernel filter. The experimenter performing the MRN computations was blinded to the results of previous EEG works, so that there were not biases in the analysis.

**Statistical analysis**

VBM results were assessed at an uncorrected threshold of p < 0.001. This threshold has an important limit in that it allows the type I statistical error. Anyway, a more permissive threshold could be more adequate to this explorative study, in order to avoid the beta (or type II) statistic error, with the risk to neglect interesting results. Anyway, the power of the study was allowed by size of the sample, and by the robust results of the subsequent analyses. The sample was disaggregated into three groups according to increasing values of alpha3/alpha2 ratio: low-a3/a2 (a3/a2<1), middle-a3/a2 (1< a3/a2<1.16), high-a3/a2 (a3/a2>1.17).

Voxel-based analyses were carried out comparing the three patient groups with increasing values of the alpha3/alpha2 ratio (high-a3/a2; middle-a3/a2; low-a3/a2).

Between-group regional differences in GM volumes were assessed by using an ANCOVA model, modelling the effects of groups (high, middle, low), and parametric nuisance covariates (age, gender, education, MMSE scores as covariates). Moreover, the total intracranial volume was introduced in the statistical analysis as a covariate, to avoid the confounding item of the global cortical atrophy. The total intracranial volume (TIV) was computed by manually tracing the entire intracranial cavity on 7 mm thick coronal slices, by the use of the software DISPLAY 1.3 tools. Correlation or regression analyses were not performed. The reason is that the EEG markers represents different patients population (MCI who will convert and MCI who will not), as previously demonstrated [3]. As a consequence, it is not correct to use the regression analysis because significant results should have masked. It would be possible only within each of the three groups individuated by the tertile subdivisions, but the size of each group dose not allows a powerful regression analysis.

All the analyses were restricted to the NAc as regions of interest in order to focus the relationship between there brain areas and EEG markers. It should be possible to perform a computation encompassing other brain areas, but this was beyond the scope of the present work. Moreover, the relationship of EEG markers with hippocampus and amygdala was faced in previous papers [28]. To this purpose, a mask including Caudate Nucleus, Putamen, Globus Pallidus, and NAc was entered into the models as explicit mask. It was manually traced, through the software MRicroN, on the previous template generated from the complete image set. The detection of the anatomical regions was based on the localization of the thalamic nuclei and basal ganglia in histological sections from a human atlas [33].

**Results**

Table 1 shows the sociodemographic characteristics of MCI subgroups. When groups were defined by the tertile values of alpha3/alpha2 the ANOVA analysis showed that there was not statistically significant differences between groups expect for alpha3/alpha2 ratio levels where all Games-Howell post hoc comparisons were significant (p < 0.001) with progressive increasing values between groups.

**EEG alpha3/alpha2 ratio**

- **Low-a3/a2 group:** No regions of GM tissue loss were found when patients with low a3/a2 ratio were compared to those with Middle and High a3/a2 ratio.

- **Middle-a3/a2 group:** Subjects with middle a3/a2 ratio, contrasted to individuals with high a3/a2 ratios, showed more atrophy in the bilateral Accumbens nuclei, though the atrophic area was minimal in the left hemisphere (Figure 1).

No regions of significant GM tissue loss were found in other comparisons in this group.

**Discussion**

**Association between EEG markers and GM changes**

Results show that the increase of alpha3/alpha2 ratio is mostly associated with minor atrophy NAc. These results show some difference with previous studies on AD patients, showing basal ganglia involvement ([34-37]) in that these reported a greater atrophy of basal ganglia. Some possible explanations are: 1) in our study the evaluation of atrophy is EEG-driven, focusing on specific morphostructural features of patients; this aspect could have magnified only some GM patterns; 2) the patterns of atrophy observed in previous studies could be linked to a specific regressive processes due to aging ([38-41]) our
results are about subjects with MCI whereas previous studies were most performed on AD patients; it should be noted that different, perhaps compensatory, mechanisms have been suggested for MCI subjects as compared to AD patients ([42-44]). We have not found any significant difference in NAc VBM analysis between the highest and lowest as well as the middle and the lowest alpha3/alpha 2 ratio MCI groups. This result is somewhat contradictory but it could be explained considering the different nature of the MCI group with highest alpha3/alpha2 ratio. Of note, this group is at major risk to develop AD, so it is a MCI due to AD group. On the contrary, the lowest alpha3/alpha2 ratio group probably is not a neurodegenerative MCI.

Anatomo-physiological relationship between EEG markers and GM changes

Our results show the presence of a specific pattern of GM changes associated with an EEG marker. The novelty emerging from our results is represented by minor atrophy of NAc in MCI patients who will develop AD. The relatively preserved anatomical structure could suggest a state of compensatory hyperfunction of this circuitry, determining both cognitive and psychiatric symptoms of prodromal AD [45]. Indeed, the NAc is associated with the regulation of emotional, impulsivity and fear behaviour control function [8]. We argue that the increase of alpha3/alpha2 ratio is the brain electrical activity marker of these structural changes. The prevalence of an anterior circuit impinging on anterior cingulate cortex, ventral (limbic) striatum and orbito-frontal cortex, relatively spared in pre-AD patients, could be hypothesized as the anatomo-physiological network underpinning the increase of alpha 3 frequency power. On the other hand, the decrease of low alpha (alpha 2 in our analysis) rhythm could be based on the well-known disrupture of the posterior circuit, encompassing hippocampal cortex, posterior cingulate, precuneus, posterior parietal cortices, in MCI subjects [21-29].

Clinical and behavioural implications

Our results show a clear association between the increase of alpha3/alpha2 power ratio and bigger volume inside the NAc in subjects with MCI, suggesting that this network could be hyperactive in patients at major risk to develop AD. These findings confirm a large
The NAc is a key entry point structure for afferent information from the periphery as well as for afferents and efferents of wider cortico-striatal-pallido-thalamo-cortical functionally segregated loops (CSPTC; [49-51]). The NAc is primarily characterized by its strong inputs from limbic structures such as the amygdala, hippocampus, mldine thalamus and certain regions of the prefrontal cortex, as well as from the mesolimbic dopamine system originating in the ventral tegmental area (VTA). Impulsive acts and decisions are related to individual differences in the neural representations of stimuli/events [52]. The NAc plays an important regulatory role in the neural representation of response options, as shown by functional neuroimaging studies in healthy individuals [4]. A voluminous literature on decision-making in humans provides imaging evidence of the involvement of a variety of brain structures in impulsive choice; different subregions of the human prefrontal cortex, cingulate cortex, insula, and the amygdala are the most prominent ones [53-55]. These structures are known to either have direct anatomical connections to the NAc, or are indirectly connected [4]. The selection of immediate reinforcement has been repeatedly reported to be associated with increased activity in the NAC [9]. Imaging studies with various task have implied an association between increased NAc activity and selection of less safe option ([14,56,57]) [12]. Another possible explanation of increase in NAc and ventral striatum hyperactivity could be related to exposure associated to novel environments. A recent study has convincingly demonstrated that a modifications in glutamatergic transmission with an increase of glutamatergic receptorial subtypes occurs in NAc after novelty exposure [58]. A modification in the shape of NAc has been demonstrated also in patients with AD showing sensation-seeking behavior [59]. In this optic the alpha3/alpha2 power ratio could a physiological network marker, related to a specific behavioral disorder, useful to screen out MCI subjects that could develop AD. The results of this paper have to be integrated by studies with a neuropsychiatric assessment and behavioral data to correlate with GM volumetric changes.

Conclusion

The a3/a2 ratio was associated with increase of grey matter density inside the NAC in MCI subjects at major risk to develop AD. Further studies will investigate the salience of this result in early diagnosis and the association of this EEG marker with cognitive impairment in prodromal AD patients.

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