Amyloid burden identifies neuropsychological phenotypes at increased risk of progression to Alzheimer’s disease in mild cognitive impairment patients

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Abstract

Purpose The extent of amyloid burden associated with cognitive impairment in amnestic mild cognitive impairment is unknown. The primary aim of the study was to determine the extent to which amyloid burden is associated to the cognitive impairment. The secondary objective was to test the relationship between amyloid accumulation and memory or cognitive impairment.

Materials and methods In this prospective study 66 participants with amnestic mild cognitive impairment underwent clinical, neuropsychological and PET amyloid imaging tests. Composite scores assessing memory and non-memory domains were used to identify two clinical classes of neuropsychological phenotypes expressing different degree of cognitive impairment. Detection of amyloid status and definition of optimal amyloid ± cutoff for discrimination relied on unsupervised k-means clustering method.

Results Threshold for identifying low and high amyloid retention groups was of SUVr = 1.3. Aβ + participants showed poorer global cognitive and episodic memory performance than subjects with low amyloid deposition. Aβ positivity significantly identified individuals with episodic memory impairment with a sensitivity and specificity of 80 and 79%, (χ² = 21.48; P < 0.00001). Positive and negative predictive values were 82 and 76%, respectively. Amyloid deposition increased linearly as function of memory impairment with a rate of 0.13/point of composite memory score (R = −44, P = 0.0003).

Conclusion The amyloid burden of SUVr = 1.3 allows early identification of subjects with episodic memory impairment which might predict progression from MCI to Alzheimer’s disease.

Trial registration EudraCT 2015-001184-39.

Keywords Mild cognitive impairment · PET imaging · Beta-amyloid · Cognitive trajectory · Memory performance

Introduction

Mild cognitive impairment (MCI) identifies the transitional stage between normal neurocognitive ageing and the progression towards several subtypes of dementia, including Alzheimer’s disease (AD) [1]. To date, there appears to be no single underlying neuropathological condition characterizing MCI, and indeed the clinical syndrome of MCI features a broad spectrum of subtypes, which differentiate from one another based on their underlying aetiology into AD, frontal-temporal dementia, vascular cognitive impairment, dementia with Lewy bodies, Parkinson’s disease, Huntington’s disease, HIV/AIDS, traumatic brain injury and substance abuse [2].

During recent decades, numerous efforts have been made to identify clinical markers to be used as reliable predictive...
markers of disease progression and thus collecting individuals at increased risk to develop AD. To date there are no means to accurately identify those likely to progress from MCI to advanced stages of dementia. Nor, in the latter case, is there a way to establish the aetiology and the pathophysiology of the process responsible for conversion, whether it be AD or other dementing conditions. Similarly, the well-established distinction among amnestic (aMCI), non amnestic (naMCI) or multiple domain MCI (mdMCI) although useful from a clinical point of view does not help to identify MCI converters to AD [3].

The recent availability of different biomarkers provide an additional tool in the definition of the pathological process. According to the evidence accumulated so far the progression from MCI to AD is characterized by a cascade of functional and structural brain and cerebral-spinal fluid (CSF) changes, starting many years before the AD onset, during which biomarkers become sequentially abnormal without clinical evidence of dementia. To date, the most convincing model of progression of cognitive impairment is that offered by Jack and colleagues in which CSF Aβ42 and brain amyloid biomarkers are the first to become abnormal, followed by biomarkers of neurodegeneration, before symptoms of AD become clinically detectable [4]. The dynamic model described by Jack and colleagues has been confirmed by a longitudinal study on dominantly inherited Alzheimer’s subjects that demonstrated the temporal progression of biomarkers changes which are characterized by an early tau and amyloid deposition increase followed, in time sequence, by neuronal dysfunction and neurodegeneration as measured by 18F-Florbetaben, a radiopharmaceutical already approved for detecting brain cortical amyloid deposition [11].

The primary aim of the study was to determine the extent at which amyloid burden is associated to the cognitive impairment as assessed by neuropsychological tests.

The secondary objective was to test the relationship between amyloid accumulation and memory or cognitive impairment.

**Methods**

The present prospective cross-sectional study was conducted at two institutions, the S. Andrea Hospital and Mem Lab & Clinics in La Spezia (Italy) between December 2015 and June 2017. The Nuclear Medicine and Neurology units of S. Andrea Hospital were involved in patient recruitment, clinical evaluation and PET imaging, and Mem Lab & Clinics conducted neuropsychological assessment.

**Patients**

The study enrolled a total of 66 participants, age ≥ 50 years, based on their medical history, clinical manifestations, and neuropsychological assessment.

Subjects inclusion criteria according to Petersen definition for amnestic MCI (aMCI) [12] consisted of Mini-Mental State Examination (MMSE) uncorrected score ≥ 24, Clinical Dementia Rating (CDR) of 0.5, absence of dementia and preserved basic activities of daily living (ADL) [13].

Presence of diseases potentially related to memory impairment, such as normal pressure hydrocephalus, Parkinson’s disease, or progressive supranuclear palsy, major structural abnormalities, signs of major vascular pathology such as intracerebral aneurysm or arteriovenous malformation, infarction, extensive leucoencephalopathy were among the exclusion criteria which also included relevant ischemic processes causing cognitive impairment, in accordance with the NINDS–AIREN criteria [14], clinical history of depression within the past year, ongoing treatment with psychotropic medication (e.g., antidepressants, neuroleptics), drug consumption and alcohol abuse.
Standard patient consent, protocol approvals, and registrations

All participants gave written informed consent after a complete written and verbal description of the study. The study had been previously approved by the regional medical ethics committee, authorized by the Italian Competent Authority (AIFA) and registered in the EudraCT database as non-profit phase III clinical trial (EudraCT number 2015–001184-39).

Clinical assessment

All participants underwent neurologic examinations, neuropsychological assessment and [18F]-Florbetaben PET/CT scan. The age of onset of the first signs of cognitive impairment was tracked back by means of a semi-structured interview to family members. MMSE scores, used for the statistical analysis as a measure of global cognitive status [15], were corrected for age and education levels according to the Italian norms (MMSEC) [16].

In this study MMSE was used as an index of global cognitive performance to identify the most impaired subjects. Among MCI subjects, those with lower cognitive performance were classified as aMCI+ (MMSEC<=24), whereas study participants with MMSEC>24 were defined as aMCI– [12–15]. Clinical severity was determined using the Clinical Dementia Rating (CDR) scale [17].

Neuropsychological assessment

Neuropsychological evaluation included neuropsychiatric interview and a comprehensive battery of cognitive tests carried out within 2 weeks prior to PET scan by certified clinical psychologists, who were blinded to the subjects’ cognitive status.

Participants were administered the following tests: Prose Memory Test (PR) [18], Rey-Osterrieth Complex Figure Test – Copy (RFCTc) [19, 20], Rey-Osterrieth Complex Figure Test - Recall (RCFTd) [20], Category Verbal Fluency (CVF) [18], Digit Symbol Substitution Test (DS) [21, 22], Digit Span forwards (DSf) and backwards (DSb) [23].

Individual scores of each test were Z-transformed with reference to the mean and SD of the whole sample. Results were grouped into Episodic Memory Composite scores (EMCs) and Non-Memory Composite scores (NMCs). Individual EMCs was expressed by averaging Z-score of RCFTd and PR and individual NMCs was the average of Z-scores for RFCTc, CVF, DS, DSf and DSb [24].

PET imaging and preprocessing procedures

PET/CT images were acquired in 3D mode 86 ± 8 min after intravenous injection of 306 ± 29 MBq of [18F]Florbetaben (FBB) (Neuraceq™) on a DISCOVERY TM 710 PET/CT scanner (GE Medical Systems, Milwaukee, WI, USA). PET projection data were iteratively reconstructed using 3-D OSEM algorithm of 8 iterations, 48 subsets, postsmoothed by a Gaussian filter of 3 mm FWHM, and with CT based attenuation correction. Image processing were performed using SPM12 (http://www.fil.ion.ucl.ac.uk/spm/software/SPM12) implemented under Matlab 8.6 (MATLAB R2015b, Mathworks Inc., Natick, MA, USA).

PET/CT images were spatially normalized to standard atlas coordinates in Talairach space using SPM T1 template [25].

PET data were converted to standardized uptake values (SUV) by scaling each image according to the body weight of each subject to the injected dose. Standardized Uptake Value Ratio (SUVr) was generated by dividing all regional SUV by the cerebellar gray matter SUV. For each subject, grey, white matter and cerebrospinal fluid (GM, WM, CSF) compartments were segmented from CT images using the segmentation routine implemented under SPM12 [26].

GM and WM voxels were then labeled according to their location, by use of Talairach Daemon database [27]. For the purpose of this study, we defined six volumes-of-interest (VOI): frontal (including inferior, medial, middle and superior gyrus), parietal (superior, inferior lobe, angular and supramarginal gyrus), temporal (inferior, middle, superior and parahippocampal gyrus), occipital (middle and inferior gyrus), posterior cingulate, and cerebellum. These VOIs were transferred onto the corresponding PET dataset to calculate the SUV mean of each brain region. Amyloid cortical burden (Aβ burden) was calculated as the average SUVr of the area-weighted mean for frontal, parietal, temporal, occipital and cingulate VOIs.

Statistics

Data were analyzed with the JMP statistical software package (SAS, Institute; Cary, NC, USA).

Individual with EMCs or NMCs lower than 10th percentile of positive values (i.e. EMCs and NMCs ≥0) were considered abnormal. Based on the value of Aβ burden, subjects with high Aβ tracer deposition (Aβ+) were set apart from those with low Aβ deposition (Aβ-) by applying the k-means cluster analysis. This method is used for a priori classification of subjects in different groups by calculating the centroid for each group and assigning each subject to the group with the closest centroid [28, 29]. The optimal cut-off to separate Aβ+ from Aβ- was SUVr = 1.30 and corresponded to 90th percentile of Aβ- cluster [29].

The analysis was restricted to two clusters representing Aβ+ or Aβ-patients whose cognitive performance were analyzed both in terms of single test results and Z-transformed EMCs and NMCs.
A linear regression model was used to evaluate the relationship between cortical amyloid deposition and global cognitive, memory and non-memory performance. The slope of regression was used to estimate the rate of amyloid deposition associated to cognitive changes.

Chi-square analyses used to test the extent to which amyloid positivity increases the risk of episodic memory, non-memory cognition and global cognitive performance impairment. Odds ratios (ORs) and their 95% confidence intervals (CIs), positive predictive value (PPV), and negative predictive value (NPV) were calculated to assess the magnitude of associations.

The differences between the mean score of neuropsychological tests in amyloid positive and negative groups were determined with one-way ANOVA.

Continuous data were analyzed using independent t-tests, with degrees of freedom adjusted for inequality of variance where appropriate. For all tests, significance was assumed as $P < 0.05$.

**Results**

Of the 66 participants initially recruited, three patients were excluded due to protocol deviations or to head movement that did not allow images analysis, thus leaving 63 subjects (who completed neuropsychological evaluation and PET imaging) for data analyses. As expected, the study sample included a slightly higher percentage of women (Table 1).

Thirty-four out of 63 (54%) aMCI subjects were classified as Aβ+, whereas 29 of 63 (46%) were identified as Aβ-. The difference between SUVr means of two amyloid clusters, reported in Table 1, was significant (mean ± sd, 1.55 ± 0.14 vs. 1.11 ± 0.07; $F = 251.5; P < 0.0001$).

The mean age of Aβ + patients was significantly higher compared to Aβ- patients ($P = 0.01$); however, no significant correlation was found between amyloid deposition and patient age. No significant differences were found among Aβ- and Aβ + groups for gender and education (Table 1). Figure 1 shows the representative image of Aβ + and Aβ- groups calculated by averaging all PET images obtained from subjects above and below the threshold of SUVr = 1.30, respectively.

As reported in Table 1 MMSE and CDR scores did not differ significantly between Aβ+ and Aβ- clusters. By contrast, the results of all other tests worsened in Aβ+ patients. However, the difference between Aβ+ and Aβ- groups reached the levels of statistical significance only for RCFTc, RCFTd, PR, CVF and DS (Table 1).

Aβ + subjects had lower EMCs and NMCs values than Aβ- ones, with a greater decrease in EMCs than NMCs scores (Table 1). Differences between mean EMCs and NMCs score in Aβ+ and Aβ- subjects were assessed by one-way ANOVA. Analysis of results showed a significant difference of EMCs ($−0.41 ± 0.68$ vs. $0.46 ± 0.74$, $P < 0.0001$) and NMCs ($−0.23 ± 0.72$ vs. $0.26 ± 0.63$, $P = 0.006$; mean ± sd, $P$) scores between groups of Aβ + and Aβ- individuals.

Among 35 subjects with episodic memory impairment 28 (80%) had positive amyloid scan and seven (20%) were classified as amyloid negative while in EMCs- group 6 (21%) were Aβ+ and 22 (79%) were Aβ- ($OR = 14.67$).

Aβ positivity significantly identified individuals with episodic memory impairment with a SS of 80% and a SP of 79%, as compared to Aβ- subjects ($χ^2 = 21.48; P < 0.00001$). The PPV was 82% and NPV was 76% (Table 2).

Of 32 with non-memory cognition impairment 23 (72%) were Aβ+ and 9 (28%) were Aβ-, whereas in NMCs- group 11 (35%) were Aβ+ and 20 (65%) were Aβ- ($OR = 4.65$).

PET amyloid positivity was associated to non-memory cognition impairment with a SS of 72% and a SP of 65% ($χ^2 = 8.39; P = 0.0038$), with a PPV of 68% and a NPV of 69% (Table 2).

Among 25 with global performance impairment 18 (72%) were Aβ+ and 7 (28%) were Aβ-, whereas in MCI+ group 16 (42%) were Aβ+ and 22 (58%) were Aβ- (SS = 72%, SP = 58%, PPV = 53%, NPV = 76%, $OR = 3.54$; $χ^2 = 5.42; P = 0.019$) (Table 2).

The linear regression of individual SUVr values as function of EMCs and NMCs scores is shown in Fig. 2. The analysis confirmed the inverse relationship between Aβ burden and both EMCs and NMCs (Fig. 2).

Interestingly, EMCs showed a stronger correlation with Aβ burden ($R = −0.44; P = 0.0003$) than NMCs ($R = −0.30; P = 0.02$) (Fig. 2).

The computed rate of deposition was estimated to be 0.13 SUVr and 0.10 SUVr per Z-transformed ECMs and NMCs unit (Fig. 2).

The significance of the correlation between amyloid retention levels and episodic memory survived also by using in the analysis the tracer retention levels measured in individual brain lobes (Frontal: $R = −0.41$, $P = 0.0009$; Parietal: $R = −0.39$, $P = 0.0015$; Temporal: $R = −0.45$, $P = 0.0003$; Occipital: $R = −0.48$, $P = 0.0001$; Posterior cingulate: $R = −0.37$, $P = 0.029$).

When correlation between amyloid retention in individual brain lobes and non-memory cognition was evaluated, the association showed a statistically significant lower extent than that measured for EMCs (Frontal: $R = −0.30$, $P = 0.016$; Parietal: $R = −0.26$, $P = 0.037$; Temporal: $R = −0.29$, $P = 0.022$; Occipital: $R = −0.25$, $P = 0.049$; Posterior cingulate: $R = −0.24$, $P = 0.055$).

**Discussion**

In recent years several studies have evidenced the important role of PET imaging assessments [11, 30–32] in providing in vivo measurements of brain amyloid deposition levels,
confirming significantly higher tracer retention in neocortical areas among MCI subjects who progress to AD compared to subjects who remain stable [7, 33]. Moreover, studies specifically including subjects with aMCI and healthy controls have reported conversion rates from MCI to AD between 59% [33] and 82% among the amyloid positive patients [7].

Table 1  Subject characteristics

| Characteristic            | All subjects | Aß+ | Aß− | F-value | P     |
|--------------------------|--------------|-----|-----|---------|-------|
| Demographic data        |              |     |     |         |       |
| N                        | 63           | 34  | 29  |         |       |
| Male                     | 27 (63)      | 13  | 14  |         |       |
| Female                   | 36 (63)      | 21  | 15  |         |       |
| Age, years               | 75.97 ± 6.59 | 76.38 ± 6.03 | 71.52 ± 7.98 | 0.01° |
| Education, years         | 9.97 ± 4.24  | 9.97 ± 4.01 | 10.69 ± 3.92 | NS°   |
| Neuropsychological battery |             |     |     |         |       |
| CDR                      | 0.5 ± 0      | 0.5 ± 0 | 0.5 ± 0 |       |       |
| Rey-Osterrieth Figure Copy | 28.68 ± 7.86 | 22.63 ± 10.51 | 31.34 ± 5.58 | 16.05 | 0.0002* |
| Rey-Osterrieth Figure Recall | 11.96 ± 5.04 | 10.55 ± 4.61 | 15.09 ± 5.73 | 12.12 | 0.0009* |
| Prose Memory             | 8.55 ± 3.26  | 6.71 ± 3.51 | 9.98 ± 2.84 | 16.50 | 0.0001^ |
| Category Verbal Fluency  | 16.92 ± 4.78 | 15.04 ± 4.6 | 18.14 ± 5.95 | 5.44  | 0.02^   |
| Digit Symbol             | 7.64 ± 2.1   | 6.29 ± 2.42 | 8.66 ± 2.58 | 14.02 | 0.0004^ |
| Digit Span forwards      | 4.83 ± 1.07  | 5.04 ± 1.07 | 5 ± 0.92  | 0.02  | NS^     |
| Digit Span backwards     | 3.75 ± 0.86  | 3.75 ± 0.98 | 3.86 ± 0.86 | 0.22  | NS^     |
| MMSE                     | 25.4 ± 3.07  | 24.74 ± 3.52 | 26.16 ± 2.27 | 3.46  | 0.07^   |
| Composite z-scores       |              |     |     |         |       |
| NMCs                     | 0.01 ± 0.61  | −0.23 ± 0.7 | 0.26 ± 0.63 | 8.22  | 0.006^  |
| EMCs                     | −0.04 ± 0.73 | −0.41 ± 0.68 | 0.46 ± 0.74 | 23.85 | <0.0001^ |
| Amyloid imaging          |              |     |     |         |       |
| FBB SUVR                 | 1.34 ± 0.25  | 1.55 ± 0.14 | 1.11 ± 0.07 | 251.47 | <0.0001^ |

Data are presented as mean ± SD; * Determined by Chi-square test; ° Determined by Student’s t test; ^ Determined by One-Way anova. Neuropsychological scores are based on age- and education-adjusted norms obtained from a prior validation study. Aß+= MCI subjects with Aß burden below the threshold of SUVR>1.3; Aß−= MCI subjects with Aß burden below the threshold of SUVR<=1.3; CDR = Clinical Dementia Rating; EMCs = Episodic Memory composite score; NMCs = Non-memory cognition composite score. FBB = [18 F]Florbetaben. SUVR = Standardized uptake value ratio.

Further studies have also attempted to define thresholds of amyloid deposition and SUVR cut-off values to discriminate cognitively normal subjects, with presumably low amyloid retention levels, from cognitively impaired patients in whom high amyloid retention is expected [9, 31, 34]. So far, however, cut-offs are inconsistent across studies, yielding higher

Fig. 1 Axial view of [18F]Florbetaben PET amyloid load in mild cognitive impairment. Average axial slices of mild cognitive impairment subjects with low (Aß−; top panel) and high amyloid load (Aß+; bottom panel). Subjects were classified using k-means clustering. Signal intensity is significantly lower in the grey matter regions of the top images compared to those of the bottom images (P < 0.0001).
values when studies included healthy controls, cognitive impaired subjects and AD patients (SUVr = 1.6) [24], as compared to studies including MCI subjects alone (SUVr = 1.5) [10, 31].

Hence, our study aimed to identify the beginning signs of amyloid accumulation in subjects belonging to a homogenous clinical entity of aMCI subjects and to evaluate potential association with clinical signs of cognitive decline. By means of [18F]Florbetaben PET imaging we were able to discriminate two aMCI subgroups with significantly different Aß retention levels. The elevated levels of FBB uptake were found to be associated to cognitive decline, in particular to a significantly greater decline in episodic memory. Evidence pointed to a threshold of SUVr = 1.30, which was able to divide our cohort into two subpopulations of Aß− (n = 29) and Aß+ (n = 34) with significantly different levels of Aß retention, thus identifying MCI subjects with episodic memory impairment which might progress from MCI to advanced stages of dementia and who most benefit from a closer clinical follow-up or anti-amyloid treatment.

This 1.30 cut-off identified a 54% proportion of Aß+ cases which is consistent with the prevalence of AD neuropathology in stable MCI and in those progressing to AD [35, 36] and to the 54% proportion reported by Jansen et al. in MCI subjects with the same average age as that enrolled in this study (75 years) [37]. Likewise, the mean value of amyloid positive subjects in our cohort, which is consistent with baseline value measured in a group of subjects converted from MCI to AD over 3 years [7].

Previous works reported that impairment of episodic memory domain is the most suitable marker for conversion to early AD [24, 38, 39]. As explained by Coulter et al. in a longitudinal study comparing cognitive changes over time in converter and non-converter amnestic MCI s into Alzheimer’s disease, the individuals progressing to AD show abnormal cognition as early as 2 years prior to the diagnosis of dementia, with episodic memory, one of the most affected domains– declining in an almost linear fashion [40].

### Table 2: Sensitivity and specificity of amyloid accumulation on cognitive decline

| Cognitive status | Aß+ | Aß− | Chi square | P   | Sensitivity (95%CI) | Specificity (95%CI) | OR (95%CI) | PPV (95%CI) | NPV (95%CI) |
|------------------|-----|-----|------------|-----|---------------------|---------------------|------------|-------------|-------------|
| EMCs+            | 28  (80%) | 7   (20%) | <0.00001   | 80% (67–93) | 79% (63–94) | 14.67 (4.3–49.9) | 82% (70–95) | 76% (60–91) |
| EMCs-            | 6   (21%) | 22  (79%) | 21.48      |     | 72% (56–87) | 65% (48–81) | 4.65 (1.6–13.5) | 68% (52–83) | 69% (52–86) |
| NMCs+            | 23  (72%) | 9   (28%) | 0.0038     | 72% (54–90) | 58% (42–74) | 3.54 (1.2–10.5) | 53% (36–70) | 76% (60–91) |
| NMCs-            | 11  (35%) | 20  (65%) | 8.39       |     | 72% (54–90) | 58% (42–74) | 3.54 (1.2–10.5) | 53% (36–70) | 76% (60–91) |
| aMCI+            | 18  (72%) | 7   (28%) | 0.0199     | 72% (54–90) | 58% (42–74) | 3.54 (1.2–10.5) | 53% (36–70) | 76% (60–91) |
| aMCI−            | 16  (42%) | 22  (58%) | 5.42       |     | 72% (54–90) | 58% (42–74) | 3.54 (1.2–10.5) | 53% (36–70) | 76% (60–91) |

Aß+ = MCI subjects with Aß burden below the threshold of SUVr>1.3; Aß− = MCI subjects with Aß burden below the threshold of SUVr<=1.3; EMCs+ = MCI subjects with episodic memory composite score <= 0.125; EMCs− = MCI subjects with episodic memory composite score > 0.125; NMCs+ = MCI subjects with Non-memory composite score <= 0.06; NMCs− = MCI subjects with Non-memory composite score > 0.06; aMCI+ = MCI subjects with MMSE score <= 24; aMCI− = MCI subjects with MMSE score > 24; CI = confidence interval; OR = Odd ratio; PPV = positive predictive value; NPV = negative predictive value;
In our study, the mean cortical florbetaben SUVr was associated with episodic memory decline. Amyloid positivity correctly identified individuals with memory impairment featuring both high sensitivity and specificity (PPV = 82%, NPV = 76%, \( P < 0.00001 \)). Moreover, the amyloid deposition changed linearly with memory decline (episodic memory resulting one of the most affected cognitive domains) with an average increase of 0.13 SUVr per memory composite score. Such observations are in line with previous studies that have reported the linear increase of amyloid deposition in the time interval between the detection of amyloid positivity and the achievement of the average SUVr threshold expected in AD (SUVr>2) [41, 42].

Interestingly, the work of Jack et al. has evidenced a bimodal trend of the deposition curve as function of time. While this was confirmed also in our study, our observations suggest a closer association between increase in amyloid deposition and clinical decline, with the amyloid deposition rate appearing to be a function of episodic memory decline. From a clinical point of view such finding is quite relevant, as it prospects the possibility of (i) reaching an earlier recognition of subjects at risk of progression to AD before neurodegeneration and irreversible related symptoms incur, and (ii) evaluating the efficacy of targeted pharmacological treatments on such patients.

The major limitation of the study lies in the cross-sectional design that does not allow to draw definitive conclusions from this analysis. Our findings should be confirmed by further studies based on longitudinal design.

Another limitation of the present study is that we used CT images for tissue segmentation which is likely to be less accurate than MR-based segmentation. We have chosen this approach on the basis of the recent report on a relatively slight differences between the probability maps of brain tissues obtained from CT and MR images [26]. Comparing 11C-PIB-PET SUV values obtained by correcting partial volume effects with CT and MR-derived probability maps authors did not find significant differences between SUV estimates as shown by the high correlation coefficient reported (R2 = 0.89).

Moreover, if these results were confirmed by other studies, PET based amyloid burden estimates could be assessed also without MR imaging, thus minimizing the patients diagnostic work-up.

In conclusion the present study reports the use of amyloid load as assessed by FBB-PET as a valid approach for objectively dichotomizing MCI individuals in amyloid positive and negative and identifying neuropsychological phenotypes characterized by increased risk of progression to Alzheimer’s disease. Our results have evidenced a significantly greater episodic memory decline in amyloid positive subjects featuring a linear correlation between amyloid load and memory decline in MCI with an average increase of 0.13 SUVr per score of episodic memory decline.

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Author contribution AC conceived the study, participated in its design and coordination, drafted the manuscript and made the final revision.

EG carried out the PET experiments, participating to critical review of data analysis and approved final version of manuscript.

BA, OF, and FF performed PET scanner quality control and image analysis, participating to critical review of data analysis and approved final version of manuscript.

GG performed statistical analysis, participating to critical review of data analysis and approved final version of manuscript.

MR carried out the radiochemistry, participating to critical review of data analysis and approved final version of manuscript.

LM participated in the study design, participating to critical review of data analysis and approved final version of manuscript.

AT participating to patient enrollement and contributed to manuscript revision;

AM, CP, EC, MDB carried out patients’ recruitment, administered neuropsychological tests, participating to critical review of data analysis and approved final version of manuscript.

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Compliance with ethical standards

Conflict of interest None.

Ethical approval All procedures performed in studies involving human participants were in accordance with the ethical standards of the institutional and/or national research committee and with the 1964 Helsinki declaration and its later amendments or comparable ethical standards.

Informed consent Informed consent was obtained from all individual participants included in the study.

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