Do cognitive patterns of brain magnetic activity correlate with hippocampal atrophy in Alzheimer’s disease?

F Maestú, J Arrazola, A Fernández, P G Simos, C Amo, P Gil-Gregorio, S Fernandez, A Papanicolaou, T Ortiz

Background: Many reports support the clinical validity of volumetric MRI measurements in Alzheimer’s disease.

Objective: To integrate functional brain imaging data derived from magnetoencephalography (MEG) and volumetric data in patients with Alzheimer’s disease and in age matched controls.

Methods: MEG data were obtained in the context of a probe-letter memory task. Volumetric measurements were obtained for lateral and mesial temporal lobe regions.

Results: As expected, Alzheimer’s disease patients showed greater hippocampal atrophy than controls bilaterally. MEG derived indices of the degree of activation in left parietal and temporal lobe areas, occurring after 400 ms from stimulus onset, correlated significantly with the relative volume of lateral and mesial temporal regions. In addition, the size of the right hippocampus accounted for a significant portion of the variance in cognitive scores independently of brain activity measures.

Conclusions: These data support the view that there is a relationship between hippocampal atrophy and the degree of neurophysiological activity in the left temporal lobe.
Table 1  Group scores on the mini-mental state examination (Spanish version, maximum score = 35), CAMCOG, and FAST

<table>
<thead>
<tr>
<th>Test</th>
<th>Group</th>
<th>Mean (SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>FAST</td>
<td>Alzheimer</td>
<td>3.5 (0.5)</td>
</tr>
<tr>
<td></td>
<td>Control</td>
<td>1.4 (0.6)*</td>
</tr>
<tr>
<td>MMSE</td>
<td>Alzheimer</td>
<td>20.3 (2.2)</td>
</tr>
<tr>
<td></td>
<td>Control</td>
<td>33.7 (1.4)*</td>
</tr>
<tr>
<td>CAMCOG</td>
<td>Alzheimer</td>
<td>56.2 (6.09)</td>
</tr>
<tr>
<td></td>
<td>Control</td>
<td>87.8 (4.2)*</td>
</tr>
</tbody>
</table>

*p < 0.01 (tests of group differences).

MEG and MRI volumetry in Alzheimer’s disease
disease, and eight age matched cognitively intact control subjects (age 75.62 (1.87) years) participated in the study. MEG data from the same patients have been reported in a previous communication. In addition to the standard battery of neuropsychological tests administered in the clinic, patients and controls received two specialised clinical tests of cognitive function, the Cambridge cognitive examination (CAMCOG) and the mini-mental state examination (MMSE). The Spanish version of the MMSE has a maximum score of 35 rather than 30 as in the English version. Information on the participants’ level of function in every day adaptive activities was obtained using the functional assessment scale (FAST). Table 1 presents the score in each of those test for both groups.

All subjects signed a consent form before participating in the study.

Stimuli and task
A detailed description of the task can be found in a report by Maestú et al., and will be summarised here briefly. Scans were obtained in the context of a letter-probe task in which five letters were presented simultaneously, followed by a series of single letters presented one at a time. The subjects were instructed to hold the items from the initial set in memory and to respond by raising their right index finger upon detecting a letter from that set during the subsequent serial presentation. In all, 250 letter stimuli were used, 50% of which were targets (that is, items included in the immediately preceding set) and 50% were distractors (letters not included in the immediately preceding set).

MEG data collection and analysis
The MEG signal was measured using a 148 channel whole head magnetometer (Magnes® 2500 WH, 4-D Neuroimaging, San Diego, California, USA) in a magnetically shielded room. A minimum of 90 epochs (one second MEG data segments) was used to calculate the average event related magnetic flux waveform (ERF) in each condition. Although different investigators have proposed a variety of source modelling approaches, we relied on the single equivalent current dipole (ECD) source model which is part of the 4D Neuroimaging software.

The intracranial generators (that is, activity sources) of the magnetic signals at successive 4 ms intervals during the course of the ERF waveform were modelled using a finite version of the non-linear Levenberg-Marquardt algorithm. Alternative algorithms hold much promise as tools for magnetic source localisation; however, they have not yet been validated against invasive localisation procedures. Currently, this model is part of the standard analysis protocol in essentially all clinical applications of MEG. Moreover, in a variety of both clinical and research applications, a single activity source is sufficient to account for 90–95% of the variance in the ERF data recorded at a given time point following stimulus onset. Most importantly, the localisation accuracy of the single ECD model for activity sources reflecting cognitive neurophysiological operations is excellent when compared with the results of invasive electrical stimulation mapping studies. The algorithm used in this study searched for the activity source that was most likely to have produced the observed magnetic field distribution at a given time. The location of activity sources was computed with reference to a Cartesian coordinate system, defined by a set of three anatomical landmarks (fiduciary points): the right and left external meatus and the nasion. The position of the magnetometers relative to the subject’s head was determined precisely using five coils, three of which were attached to the fiduciary points and two on the forehead. The coils were activated briefly at the beginning and again at the end of the recording session, and their precise location in three dimensional space was determined using a localisation algorithm built into the system. During the recording session a fiberoptic motion detector was used to ensure that the subject’s head did not change position relative to the sensor.

Activity source solutions were considered as satisfactory upon meeting the following criteria: (1) correlation and goodness of fit ≥ 0.90 between the observed and the best predicted magnetic field distribution; (2) a 95% confidence volume of less than 10 cm³.

In order to identify the anatomical regions where the activity sources were localised, activity source coordinates were overlaid onto T1 weighted magnetic resonance images using the STAR software which is part of the 4D Neuroimaging software. Precise coregistration of the MEG coordinate system onto the MRI was achieved by aligning the MEG fiduciary points with high contrast cod liver capsules (3 mm in diameter) which were fixed to the subject’s nasion and inserted in the external meatus before the MRI scan.

The sum of all acceptable sources localised in a particular brain region, starting at stimulus onset and ending one second later, served as a metric of the degree of stimulus locked activation of that area. The validity of this measure as an index of regional activation has been established in several studies involving neurologically intact volunteers and patients. Further, in order to extract information regarding the relative timing of group differences, the number of activity sources in each area was summed within each of 10 consecutive 100 ms time windows. The following variables, representing the total number of activity sources in response to target stimuli, were associated with group ANOVA (analysis of variance) effects in our previous study, and were considered in the present study: (1) left temporal lobe activity (including activity in the hippocampus) in the 400–500 and 600–700 ms time windows; (2) left parietal activity in the 400–500 and 600–700 ms latency windows.

MRI volumetry
High resolution three dimensional volume scans were acquired from each subject with a 1.5 Tesla scanner (Signa, General Electric (GE), Milwaukee, Wisconsin, USA). T1 weighted images (TR 14.6 ms/TE 3.1 ms/FA 15°) were obtained based on gradient echo (3D-SPGR) sequences with a field of view of 22 x 16 cm, 256 x 194 matrix, one excitation, and 1 mm slice thickness, covering the entire brain and skull in the axial plane. The voxel size was 0.7 mm³. The volumetric measurements were done with the RM tissue volume software developed by GE Medical Systems running on an AdvantEdge Windows 4.0 environment. Regions of interest in each scan were defined using the semi-automatic volume segmentation protocol (3SAYS) which is part of the software. This software allows the user to draw a region of interest in each slice with the mouse, assisted by border attraction and three dimensional automatic paint tools, and measure the volume of the painted region (in cm³).

After the measures have been obtained the software produces a histogram of the voxel value for the region of interest. The following regions of interest were defined: total cranial volume (TCV), total cerebral volume (TCV), right and left hippocampal
volume (RHV, LHV), and right and left lateral surface of the temporal lobe (RTLS, LTLS) (fig 1). In order to determine anatomical landmarks on the lateral temporal surface, the Sylvian fissure was traced and the parieto-occipital fissure and the preoccipital notch were identified and joined by a line, following procedures described by Kidron et al. Then a line was drawn from the posterior end of the Sylvian fissure to the line joining the parieto-occipital fissure and preoccipital notch to provide the posterior demarcations of the temporal lobe. After this the ratio of total cerebral to total cranial volume (relative cerebral volume or CVr) was calculated according to the formula:

\[
CVr = \frac{TCV}{TCrV} \times 100.
\]

In a similar manner we computed the proportion of the total cranial volume represented by: the right hippocampus (RHVr), the left hippocampus (LHVr), the right temporal lobe (RTLSr), and the left temporal lobe (LTLSr). This approach was introduced by Whitell et al. in order to control for individual differences in brain size resulting from normal intersubject variation, as well as variation caused by brain pathology. Only relative measures were used in the statistical analyses in order to control for possible group differences in brain size.

To determine the reliability of the MRI measures, we estimated the degree of agreement between the two neuroradiologists who did the volumetric measurements independently without knowledge of the clinical or MEG results. The mean difference in measurements between the two observers for the left hippocampus was −0.03 cm³ (95% confidence interval −0.14 to 0.08 cm³) and 0.07 cm³ for the right hippocampus (−0.3 to 0.1 cm³). Similar results were found for the left temporal lobe (−0.1 cm³ (−0.98 to 0.75 cm³)) and for the right temporal lobe (0.17 cm³ (−0.5 to 0.2 cm³)). Interrater reliability coefficients were very high (left hippocampus, \( r = 0.99 \); right hippocampus, \( r = 0.98 \); left temporal lobe, \( r = 0.94 \); right temporal lobe, \( r = 0.95 \)). Corresponding coefficients for the parietal lobes were much lower (< 0.70), in part because of the inherent difficulty in defining and visually identifying anatomical borders for this area. Accordingly, parietal volumes were not used in subsequent analyses.

RESULTS
Volumetric MRI analyses: group effects
A series of mean comparisons (Student \( t \) tests) between groups was performed on the MRI volumetric variables. Given a total number of five comparisons (CVr, R HVr, L HVr, RTLSr, LTLSr), an adjusted \( \alpha \) level of 0.05/5 = 0.007 was used to evaluate each \( t \) test by the Bonferroni method. The Alzheimer’s disease group had lower volumes, adjusted for total intracranial volume, than the elderly control group in both the left (\( t(14) = 5.68, \ p < 0.0001 \)) and the right hippocampus (\( t(14) = 5.69, \ p < 0.0001 \)). The two groups did not differ on the other relative volume measures (\( p > 0.2 \)).

Relations between volumetric MRI and MEG data
A series of Pearson correlation coefficients was computed between the MEG and the MRI volumetric variables. We focused on the two volumetric variables that reliably differentiated between groups, namely relative left and relative right hippocampal volumes. Three additional variables were examined for contrast: total brain volume (adjusted for head size) and the relative volume of the entire left and the entire right temporal lobes. Two MEG based activity measures were
used, one representing the number of late activity sources in the left temporal lobe and the other those in the left parietal lobe. Temporal lobe activity sources included those localised in the mesial temporal regions. These sources, however, were not found in all subjects and, given the relatively small group size, were not amenable to separate parametric analyses.

Each variable in the left hemisphere reflected the sum of activity sources in two latency windows (400 to 500 and 600 to 700 ms) which reliably discriminated between the two groups, as reported previously.11 Like the MRI data, the MEG data were normalised on the basis of the total number of activity sources in the entire brain for each participant. In all, 10 correlation coefficients were computed (between each of the two MEG variables and each of the five MRI variables) and evaluated using a Bonferroni corrected α level of 0.05/10 = 0.005. In general, strong positive correlations were found between the two MEG variables and the relative hippocampal volume bilaterally (table 2). The only coefficient that was considered statistically significant, however, was that between left temporal lobe activity (including hippocampal activity) and the relative volume of the left hippocampus, indicating that the smaller the number of late activity sources in the left temporal lobe areas, the greater the atrophy in the mesial aspects of the left temporal lobe.

Owing to the inherent difficulty in determining the anatomical borders of the parietal lobe, inter-rater reliabilities among radiologists were not sufficient to warrant the use of the parietal lobe volume measures in the analyses. Interestingly, however, there was a significant positive correlation between degree of activity in the left parietal lobe and the relative volume of the left temporal lobe. Given that the bulk of parietal activity was noted near the temporoparietal junction, this finding is not surprising, in view of the close links between the left temporal and parietal lobes in language and memory functions.

**Relations between volumetric MRI, MEG, and neuropsychological data**

In a series of stepwise linear regression analyses we examined the relative significance of MEG and MRI volumetric measures as predictors of performance on the two tests of cognitive function (MMSE and CAMCOG) and the scale of everyday adaptive behaviours (FAST). Four variables that were found to differentiate reliably between the two groups were used in these analyses. The design included two volumetric MRI measures (left and right relative hippocampal volume) and the two MEG measures used above (relative degree of activity in the left temporal and parietal lobes).

These analyses indicated that only two variables make significant independent contributions to the variability of each of three behavioural measures: the degree of activity in the left temporal lobe and the relative volume of the right hippocampus. Combined, the two variables accounted for 77–85% of the variance in each of the three measures (table 3).

**DISCUSSION**

In agreement with previous reports,14 patients with Alzheimer's disease consistently showed greater atrophy in the mesial temporal areas in both hemispheres than in other areas. Further, our data show that the degree of left hippocampal atrophy correlates strongly with the magnitude of regional activation of left temporal areas during a short term memory task. The smaller the size of the left hippocampus—adjusted for total intracranial volume—the smaller the number of consecutive activity sources in left temporal areas. Notably this activity was found exclusively during the late stages of neural processing of the target stimuli—that is, between 400 and 700 ms after stimulus onset.15 In a similar manner, the magnitude of late activity in left parietal regions is a strong correlate of the relative volume of the ipsilateral temporal lobe. This finding corroborates previous reports of predominant left hemisphere metabolic dysfunction in Alzheimer’s disease.1 Moreover, the real time functional mapping capability of MEG extends these findings to suggest that this dysfunction is related to neurophysiological operations that are part of the brain mechanism, which at least in part, supports memory function.

We can offer two alternative explanations for the relation between the structural integrity of the mesial temporal cortices and the functional status of temporal and parietal lobe regions. First, it is possible that degenerative changes in the left entorhinal cortex early in the course of the disease impair critical functional connections between the neocortex and the hippocampal formation.16 The existence of a direct functional link between the two parts of the temporal lobe is also supported by a strong association between temporoparietal hypometabolism and hippocampal atrophy.10 A disconnection between the hippocampal complex in the left hemisphere and ipsilateral neocortical areas could seriously impair the

**Table 2 Correlation coefficients between MEG and MRI variables**

<table>
<thead>
<tr>
<th>MRI</th>
<th>L hippocampus</th>
<th>R hippocampus</th>
<th>Brain</th>
<th>L temporal</th>
<th>R temporal</th>
</tr>
</thead>
<tbody>
<tr>
<td>MEG L temporal</td>
<td>0.70†</td>
<td>0.59*</td>
<td>0.26</td>
<td>0.26</td>
<td>0.19</td>
</tr>
<tr>
<td>MEG L parietal</td>
<td>0.65†</td>
<td>0.61*</td>
<td>0.30</td>
<td>0.72†</td>
<td>0.15</td>
</tr>
</tbody>
</table>

* p < 0.025, † p < 0.01, ‡ p < 0.003.

L, left; MEG, magnetoencephalography; MRI, magnetic resonance imaging; R, right.

**Table 3 The relative importance of MEG and MRI variables in predicting cognitive function and adaptive behaviour**

<table>
<thead>
<tr>
<th>MMSE</th>
<th>CAMCOG</th>
<th>FAST</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image-url" alt="Table content" /></td>
<td>0.695 (2)</td>
<td>0.006</td>
</tr>
<tr>
<td><img src="image-url" alt="Table content" /></td>
<td>0.684 (1)</td>
<td>0.001</td>
</tr>
<tr>
<td><img src="image-url" alt="Table content" /></td>
<td>0.819 (1)</td>
<td>0.001</td>
</tr>
<tr>
<td><img src="image-url" alt="Table content" /></td>
<td>0.854 (2)</td>
<td>0.001</td>
</tr>
<tr>
<td><img src="image-url" alt="Table content" /></td>
<td>0.660 (1)</td>
<td>0.003</td>
</tr>
<tr>
<td><img src="image-url" alt="Table content" /></td>
<td>0.684 (1)</td>
<td>0.001</td>
</tr>
<tr>
<td><img src="image-url" alt="Table content" /></td>
<td>0.819 (1)</td>
<td>0.001</td>
</tr>
<tr>
<td><img src="image-url" alt="Table content" /></td>
<td>0.854 (2)</td>
<td>0.001</td>
</tr>
<tr>
<td><img src="image-url" alt="Table content" /></td>
<td>0.660 (1)</td>
<td>0.003</td>
</tr>
<tr>
<td><img src="image-url" alt="Table content" /></td>
<td>0.684 (1)</td>
<td>0.001</td>
</tr>
<tr>
<td><img src="image-url" alt="Table content" /></td>
<td>0.819 (1)</td>
<td>0.001</td>
</tr>
<tr>
<td><img src="image-url" alt="Table content" /></td>
<td>0.854 (2)</td>
<td>0.001</td>
</tr>
</tbody>
</table>

Adj R², adjusted R² value for each predictor variable; CAMCOG: Cambridge cognitive examination; FAST, functional assessment scale; L, left; MEG, magnetoencephalography; MMSE, mini-mental examination (Spanish version); MRI, magnetic resonance imaging; R, right.

p Value: test of the significance of the independent contribution of the first (1) and second variable (2) to enter the regression equation.
functional integrity of the neural circuit involved in memory function. At the behavioural level this may have a major effect on the circuit’s efficiency in accessing stored information, thus impairing memory function. At the same time the hypothesised disconnection could lead to reduced neurophysiological activation in left temporal and parietal areas during the performance of memory tasks, as noted in the present study.

A second and complementary hypothesis derives from the notion that normal short term or working memory function depends upon the functional integrity of all the components of the brain circuit involving the mesial as well as the lateral temporal and parietal lobe areas. According to current theoretical models, working memory involves several component processes, including a phonological loop, visuo-spatial representations, central executive processes, and an episodic buffer. In our study we used a task that poses heavy demands on the phonological loop, a process intimately linked to the temporoparietal areas. If any of these regions is affected by degenerative processes—such as neuronal cell loss, neurofibrillary tangles, and senile plaques that are routinely found in these regions in Alzheimer’s disease—the ability of the circuit to support memory related functions will be impaired.

Given the strong positive correlation between left hippocampal volume and degree of left activity in the ipsilateral temporal lobe, it is not surprising that the variable that makes a significant independent contribution to predicting individual scores on general cognitive/behavioural measures is atrophy in the right hippocampus. This finding is consistent with previous reports of an association between bilateral mesial temporal lobe hypometabolism and performance on cognitive scales like the MMSE.

The fact that hippocampal atrophy improves the value of MEG derived region activation measures in predicting individual scores on the cognitive (MMSE, CAMCOG) and functional (FAST) scales establishes a direct link between cognitive impairment and anatomical-functional measures, and highlights the potential clinical significance of both types of measure for the early diagnosis of Alzheimer’s disease. It should be noted, however, that the widespread concurrent use of both imaging modes is unlikely in the near future, given the relatively small number of MEG centres worldwide.

This study supports the view that the lack of left brain magnetic activity in the left tempo-parietal region in patients with Alzheimer’s disease may be related to the degree of atrophy in mesial temporal lobe structures. The results highlight the importance of assessing the complex brain pathology underlying Alzheimer’s disease using multiple brain imaging modes. In addition, a conservative analytical approach was adopted in order to keep the likelihood of type I error at a minimum. This approach, which was thought necessary in view the large number of post hoc comparisons performed, was designed to keep the rate of false positive results under control. The sensitivity of MEG/MRI measures can be improved in future investigations by increasing sample size in order to enhance power and reduce the rate of type II error.

ACKNOWLEDGEMENTS
This study was supported in part by a grant from 4-D Neuroimaging. We would like to thank Thomas Murphy for technical assistance on MEG data acquisition.

Authors’ affiliations
F Maestu, A Fernández, C Amo, S Fernandez, T Ortiz, Centro de Magnetoencefalografía Dr Pérez Modrego, Universidad Complutense Madrid, Spain
J Arrazola, Department of Neuroradiology, Hospital Universitario San Carlos, Madrid
P G Simos, A Papanicolaou, Department of Neuroradiology, Vivian L Smith Center for Neurologic Research, University of Texas-Houston Medical School, Houston, Texas, USA

Pedro Gil-Gregorio, Departamento de Geriatria, Hospital Universitario San Carlos, Madrid

Competing interests: none declared

REFERENCES
26 Lee I, Kesner RP. Differential contribution of NMDA receptors in hippocampal subregions to spatial working memory. Nat Neurosci 2002;5:162–8

www.jnnp.com
Do cognitive patterns of brain magnetic activity correlate with hippocampal atrophy in Alzheimer's disease?

F Maestú, J Arrazola, A Fernández, P G Simos, C Amo, P Gil-Gregorio, S Fernández, A Papanicolaou and T Ortiz

*J Neurol Neurosurg Psychiatry* 2003 74: 208-212
doi: 10.1136/jnnp.74.2.208

Updated information and services can be found at:
[http://jnnp.bmj.com/content/74/2/208](http://jnnp.bmj.com/content/74/2/208)

These include:

**References**
This article cites 28 articles, 7 of which you can access for free at:
[http://jnnp.bmj.com/content/74/2/208#BIBL](http://jnnp.bmj.com/content/74/2/208#BIBL)

**Email alerting service**
Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

**Topic Collections**
Articles on similar topics can be found in the following collections

- Neuroimaging (389)

**Notes**

To request permissions go to:
[http://group.bmj.com/group/rights-licensing/permissions](http://group.bmj.com/group/rights-licensing/permissions)

To order reprints go to:
[http://journals.bmj.com/cgi/reprintform](http://journals.bmj.com/cgi/reprintform)

To subscribe to BMJ go to:
[http://group.bmj.com/subscribe/](http://group.bmj.com/subscribe/)