Neural correlates of lexical-semantic memory

A voxel-based morphometry study in mild AD, aMCI and normal aging

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Abstract — Neuroanatomical correlations of naming and lexical-semantic memory are not yet fully understood. The most influential approaches share the view that semantic representations reflect the manner in which information has been acquired through perception and action, and that each brain area processes different modalities of semantic representations. Despite these anatomical differences in semantic processing, generalization across different features that have similar semantic significance is one of the main characteristics of human cognition. Methods: We evaluated the brain regions related to naming, and to the semantic generalization, of visually presented drawings of objects from the Boston Naming Test (BNT), which comprises different categories, such as animals, vegetables, tools, food, and furniture. In order to create a model of lesion method, a sample of 48 subjects presenting with a continuous decline both in cognitive functions, including naming skills, and in grey matter density (GMD) was compared to normal young adults with normal aging, amnestic mild cognitive impairment (aMCI) and mild Alzheimer’s disease (AD). Semantic errors on the BNT, as well as naming performance, were correlated with whole brain GMD as measured by voxel-based morphometry (VBM). Results: The areas most strongly related to naming and to semantic errors were the medial temporal structures, thalami, superior and inferior temporal gyri, especially their anterior parts, as well as prefrontal cortices (inferior and superior frontal gyri). Conclusion: The possible role of each of these areas in the lexical-semantic networks was discussed, along with their contribution to the models of semantic memory organization.

Key words: semantic memory, naming, voxel-based morphometry, Alzheimer’s disease, mild cognitive impairment.

Correlatos neurais da memória lexical-semântica: um estudo de morfometria baseada em voxel na doença de Alzheimer, comprometimento cognitivo leve amnésico e envelhecimento normal.

Resumo — As correlações neuroanatômicas de nomeação e memória lexical-semântica não são totalmente entendidas. As abordagens mais influentes compartilham a ideia de que representações refletem a maneira na qual a informação foi adquirida por meio da percepção e ação e que cada área do cérebro processa diferentes modalidades de representações semânticas. Apesar destas diferenças anatômicas no processamento semântico, a generalização de diferentes aspectos que tem significância semântica análoga é uma das principais características da cognição humana. Métodos: Nós avaliámos as regiões cerebrais relacionadas à nomeação e à generalização semântica de desenhos de objetos visualmente apresentados do Teste de Nomeação de Boston (TNB), que compreende diferentes categorias, como animais, vegetais, utensílios, comida e móveis. Para criar um modelo de método de lesão, nós investigamos uma amostra de 48 pessoas que demonstraram um declínio contínuo em funções cognitivas, incluindo habilidades de nomeação e em densidade de substância cinzenta (DSC) em relação a adultos jovens: envelhecimento normal, comprometimento cognitivo leve amnésico (CCLa) e doença de Alzheimer (DA). Nós correlacionamos seus erros semânticos no TNB, assim como seus desempenhos de nomeação, com a DSC de todo o cérebro que foi medido por morfometria baseada em voxel (MBV). Resultados:
Language is one of the most important characteristics that allows us to codify, signify, and retain our experience of the world. Naming the many aspects of our environment is an essential attribute for the evolution of human complex adaptive ability and reveals the capacity to learn and share knowledge. Lexical-semantic memory refers to the storage of this knowledge in the brain by means of patterns of neuronal activity interpreted as linguistic symbols of concrete and abstract concepts. The relationship between brain anatomy and the storage of these patterns of information is not yet well understood. Several hypotheses have been proposed to explain how lexical-semantic memory is processed and stored in the brain, and these have been guided by two main general models: a parallel distributed representation, comprising a homogeneous network of equivalent neuronal units that process every aspect of semantics, and a center processing model, which assumes that all memory elements are encoded in a distributed representation, in a combination of these two theories has been proposed.

One of the main characteristics of human cognition is the capacity to generalize across concepts that have similar semantic significance but not necessarily similar specific (physical or behavioural) attributes. The most striking evidence of deterioration of this generalizing capacity, manifested initially by semantic naming errors production, is semantic dementia (SD), in which there is a degeneration of the anterior portions of the temporal lobes, and is more intense on the left side. These patients have difficulties in naming everyday objects and knowing their properties, with impairment of all kinds of concepts in the context of otherwise well-preserved cognition, including episodic memory. Other diseases associated with lesions in the anterior parts of temporal lobe show the same pattern of loss of knowledge, particularly in Herpes simplex virus encephalitis, stroke, and Alzheimer’s disease (AD). In this sense, as proposed by other authors, the temporal lobe, particularly its anterior part, may constitute a convergence zone for information coming from brain regions responsible for processing different aspects of knowledge. It has also been suggested that the temporal lobe object representation system may be organized hierarchically, with increasing convergence and integration of information occurring along its posterior to anterior axis.

Naming complaints are very common in mentally healthy elderly people. Individuals over the age of seventy attain significantly lower scores on these naming tests compared to scores by young adults. Problems with naming and word finding are even more common in mild cognitive impairment (MCI) and are most common in Alzheimer’s disease (AD). MCI is a clinical entity applied to patients with objective cognitive problems, most commonly in episodic memory, without significant impairment of activities of daily life.

Our aims were to evaluate the brain regions related to naming performance and to spontaneous semantic naming errors on the Boston Naming Test (BNT), regardless of category (animals, vegetables, tools, food, and furniture, etc.). In order to create a model of the lesion method, a sample of subjects presenting with continuous decline both in cognitive functions, including naming skills, and in grey matter density (GMD) were compared to normal young adults with normal aging, amnestic mild cognitive impairment (aMCI), and mild AD. Semantic errors on the BNT were correlated with whole GMD as measured by voxel-based morphometry (VBM). This correlation was also performed for BNT total score (correct responses). We hypothesised that temporal lobes, especially their anterior parts, are related with semantic naming error production in this sample of subjects. The majority of structural neuroimaging studies in patients with language problems have employed volumetric measurements on magnetic resonance imaging (MRI) data sets. This kind of approach has used the region-of-interest method, which depends on a priori choices and can be applied to a selected set of brain structures only. A whole-brain VBM approach on the other hand, has the advantage of not only evaluating the previously hypothesized brain structures, but also potentially revealing unexpected areas of gray matter density changes and their correlation with neuropsychological scores.

**Methods**

A total of 48 subjects older than 50 years [17=aMCI, 15=mild AD treated at the Unit for Neuropsychology and Neurolinguistics (UNICAMP Clinic Hospital), and...
Assessment of naming ability

The sixty-item BNT was translated and culturally adapted version for the Brazilian population by Dr. Cândida Camargo (Psychiatry Institute, University of São Paulo School of Medicine), for which subjects were asked to name the presented pictures, was administered to all subjects. BNT total score was calculated by adding the number of correct spontaneous responses to the number of correct responses after a semantic cue, which consisted of a short explanation about the picture (for example, for mask: it’s part of a carnival costume) or a superordinate category (e.g. for elephant: it’s a kind of animal). The semantic cue was only given if the patient had failed to recognize the picture (for example: dog instead of tree) or if he/she said that they did not know what the picture was.

Semantic errors registered when the spontaneous answer was semantically related to the target word. Two independent researchers performed this classification, and the discords were solved by consensus.

Additional neuropsychological evaluation

All subjects were submitted to tests of verbal fluency (VF) - animals category (score=total number of different animal names/one minute); Mini Mental State Examination (Brazilian version); Rey auditory verbal learning test episodic memory delayed recall (RAVLT-A7) and CAMCOG’s subscale of similarities [pairs of nouns - “In what way are they alike?” for the following pairs: apple/banana, chair/table, shirt/dress, and animal/vegetable (score=correct number responses- zero to two for each pair; maximum score eight)]; visual perception subtests of Luria’s Neuropsychological Investigation (LNI; maximum score twenty); the forward (FDS) and backward digit span (BDS) subtests of the WAIS-R and the Cornell Scale for Depression in Dementia (CSDD).

Data analysis was performed using Systat software 12.0. The Kruskall-Wallis and Mann-Whitney tests were used for inter-group comparisons of demographic and cognitive scores. Statistical significance was considered when p<0.05.

MRI scanning protocol and imaging processing

High-resolution MRI was performed using a 2.0 T scanner (Elscint, Haifa, Israel). T1- and T2-weighted images were acquired in axial, coronal and sagittal planes with thin slices. In addition, volumetric (3D) T1 gradient echo (GRE) images were acquired in the sagittal plane with 1 mm-thick slices (flip angle=35°, time to repeat=22 ms, echo time=9 ms, matrix=256×220, field of view=23 cm). Before pre-processing all scans were checked for scanner artefacts and gross anatomical abnormalities. MRicro was used to convert the original DICOM format to ANALYZE format (www.mricro.com) and set the origin of the coordinate system at the anterior commissure.

SPM8b (Wellcome Department of Imaging Neuroscience, London, England; www.fil.ion.ucl.ac.uk) run on MATLAB 7.5 was used to perform voxel-based morphometry (VBM). For segmentation, the “New Segment” toolbox from SPM8b was employed. The algorithm used for segmentation is based on “Unified Segmentation”, which in turn is based on a mixture of Gaussian and combines image registration, tissue classification and bias correction within one generative model. In order to obtain a more accurate inter-subject alignment the DARTEL (Diffeo-
morphic Anatomical Registration Through Exponential Lie Algebra) registration model was chosen. In addition, to preserve the volume of each tissue (modulation step) the warped images were scaled using the Jacobian determinants. Finally, the normalized, segmented, modulated (Jacobian-corrected) and warped images were smoothed by convolving with an isotropic Gaussian kernel with full width at half maximum of 10 mm to reduce interindividual gyral variation. After this pre-processing, the resulting normalized, modulated and smoothed data was used for statistics treatment.

Voxel-based correlation analysis

Multiple regression analysis using Non-Parametric Mapping (NPM) software (http://www.sph.sc.edu/comd/rorden/npm) to identify brain regions whose GMD values were significantly correlated with the absolute number of semantic errors and BNT total score. Age, education, total intracranial volume and global cognition (as measured by MMSE) were also included in the analysis as dependent variables. Total intracranial volume was obtained by the sum of volumes of grey matter, white matter and cerebrospinal fluid. For multiple regression analysis, the three groups were considered together (normal aging, aMCI and mild AD) to increase data variance and enhance the correlation between cerebral region and psychological function, assuming that the denser the gray matter, the better the naming performance and vice versa.

The results were corrected for multiple comparisons by using Bonferroni Correction, which involves adjustments to the statistical threshold to control for overall familywise error rate (FWE). To control for FWE, permutation analysis was also performed using NPM, with 4000 permutations analyzed.

Results

As shown in Table 1, no significant difference was found among the three groups with regard to age (p=0.17) or education (p=0.31). There was a continuum in neurological performance on all tests, except backwards digit span. With regard to BNT total score, AD patients performed worse than both aMCI patients and controls (p<0.001), while aMCI subjects performed worse than controls on BNT spontaneous answers (without cues; p<0.05). The absolute values of spontaneous errors and total number of semantic errors are shown in Table 1. Regarding brain atrophy, results revealed a continuum among the groups, as shown in a previous study involving the same subjects.

Multiple regression analysis revealed significant correlations between GMD and BNT score, mostly in the thalamus: right lateral dorsal nucleus and left medial dorsal nucleus; bilateral hippocampus; bilateral parahippocampal gyrus; left superior temporal gyrus; left inferior frontal gyrus; bilateral superior frontal gyrus; left middle frontal gyrus, and other areas shown in Table 2 and Figure 1. Areas of correlations with spontaneous errors are shown in Table 3 and Figure 2. Semantic errors were related mainly to the bilateral anterior part of the temporal lobe: super-

### Table 1. Demographic and neuropsychological data.

|                  | AD        | aMCI      | Controls | P       |
|------------------|-----------|-----------|----------|---------|
| Age              | 74.26±6.33| 68.29±9.93| 69.12±7.55| 0.170   |
| Education        | 6.00±5.52 | 5.88±4.32 | 6.87±3.66 | 0.315   |
| MMSE             | 22.93±2.65| 26.41±2.76| 29.12±0.71| <0.0001 |
| BNT-total score  | 39.33±9.98| 50.82±7.66| 53.75±4.18| <0.0001 |
| BNT- spontaneous | 34.87±9.7 | 48.25±9.13| 51.62±5.87| <0.05   |
| Omission errors  | 6.43±5.39 | 2.50±2.65 | 1.62±2.50 | 0.006   |
| Visual paragnosia| 7.87±3.72 | 4.18±4.73 | 2.00±2.19 | <0.0001 |
| Semantic errors  | 10.31±4.06| 4.81±3.16 | 4.43±2.44 | <0.0001 |
| CAMCOG’s Similarities | 4.87±1.74 | 6.93±1.18 | 7.37±1.02 | <0.001  |
| A7-RA VLT       | 1.26±1.28 | 4.17±2.40 | 9.56±3.03 | <0.0001 |
| VF               | 10.60±3.39| 13.64±3.92| 19.43±3.03| <0.0001 |
| VSP-LNI          | 17.20±1.42| 18.76±0.97| 18.81±0.98| 0.002   |
| fDS              | 4.46±1.06 | 4.58±0.79 | 5.06±0.85 | 0.108   |
| bDS              | 3.20±0.77 | 3.11±0.92 | 4.12±1.02 | 0.004   |

Data expressed as mean±SD. MMSE: Mini-Mental State Examination; A7-RA VLT: delayed recall of Rey Auditory Verbal Learning Test; BNT: Boston Naming Test; VF: Verbal Fluency; VSP-LNI: visuospatial perception item of Luria’s neuropsychological investigation; fDS: Forward Digit Span; bDS: backward Digit Span.
**Table 2.** Brain areas exhibiting statistically significant correlation with BNT score.

| Region                                      | Number of voxels in cluster | Talairach coordinates | Z score |
|---------------------------------------------|------------------------------|------------------------|---------|
| Left parahippocampal gyrus                  | 97                           | –28 –40 –5            | 2.82    |
| Left hippocampus                            | 79                           | –29 –15 –18           | 2.35    |
| Right hippocampus                           | 80                           | 34 –12 –20            | 2.11    |
| Right thalamus, pulvinar                    | 87                           | 9 –28 9              | 2.35    |
| Left thalamus, ventral anterior nucleus     | 99                           | –12 –7 13            | 2.17    |
| Left thalamus, lateral dorsal nucleus       | 95                           | –10 –17 17           | 2.09    |
| Left inferior frontal gyrus (BA 44)         | 73                           | –50 5 19            | 2.02    |
| Left superior frontal gyrus (BA 9)          | 84                           | –1 55 25            | 2.07    |
| Left precuneus                              | 96                           | –17 83 40            | 2.00    |
| Left superior frontal gyrus (BA 8)          | 77                           | –17 36 51            | 3.05    |
| Right superior frontal gyrus (BA 6)         | 79                           | 17 26 58            | 2.66    |

BA: Brodmann’s area.

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**Table 3.** Brain areas exhibiting greatest statistically significant correlation with semantic errors.

| Region                                      | Number of voxels in cluster | Talairach coordinates | Z score |
|---------------------------------------------|------------------------------|------------------------|---------|
| Left superior temporal gyrus (BA 38)        | 100                          | –29 6 –28             | 4.32    |
| Right superior temporal gyrus (BA 38)       | 88                           | 38 10 –28             | 3.45    |
| Right middle temporal gyrus (BA 21)         | 109                          | 44 3 –34              | 3.48    |
| Left middle temporal gyrus (BA 21)          | 72                           | –48 1 –21             | 2.90    |
| Left inferior temporal gyrus (BA 20)        | 119                          | –49 –4 –37            | 2.21    |
| Right inferior temporal gyrus (BA 20)       | 83                           | 45 –12 –37            | 3.16    |
| Right parahippocampal gyrus (BA 28)         | 60                           | 16 –3 –13             | 2.48    |
| Left uncus                                  | 54                           | –21 –7 –37            | 3.00    |
| Left globus pallidus                        | 103                          | –21 –3 –6             | 2.67    |
| Right anterior cingulate (BA 25)            | 51                           | 1 10 –3               | 2.33    |
| Left thalamus, medial dorsal nucleus        | 55                           | –1 19 10             | 2.70    |
| Right thalamus, medial dorsal nucleus       | 87                           | 3 –20 6             | 2.39    |
| Left thalamus, lateral dorsal nucleus       | 95                           | –11 –19 14           | 2.44    |
| Right thalamus, lateral dorsal nucleus      | 95                           | 11 –19 14            | 2.09    |
| Right caudate nucleus                       | 84                           | 9 17 1              | 2.40    |
| Left caudate nucleus                        | 86                           | –6 4 1              | 2.36    |
| Left putamen                                | 104                          | –23 –2 1             | 2.81    |
| Left inferior frontal gyrus (BA 44)         | 91                           | –51 9 20             | 2.16    |
| Right inferior frontal gyrus (BA 44)        | 83                           | 52 4 20             | 2.12    |
| Left precuneus (BA 7)                       | 101                          | –3 –76 44            | 2.73    |

BA: Brodmann’s area.
rior temporal gyrus; left inferior temporal gyrus; bilateral dorsomedial thalamic nucleus; bilateral hippocampal and left caudate nucleus.

Discussion

Our results support the hypothesis of a continuum in brain pathology and cognitive decline among the three groups, particularly regarding their spontaneous answers during BNT picture naming, which indicates that our lesion model was satisfactorily tested. Several brain regions were found to be negatively correlated with semantic errors on the BNT (i.e. the more errors made, the lower the GMD in that particular area), and positively correlated with BNT score. A discussion follows on the possible role of each of these areas in the lexical-semantic networks and their contribution to the models of semantic memory organization.

Medial temporal structures such as the hippocampus and parahippocampal gyrus have a well-known role in episodic memory processes. Recently, they have also been associated with lexical-semantic memory. In fact, episodic and semantic memories are highly interactive. It is well established that episodic memory for events encoded during semantic categorization is better remembered than when subjects do not associate the target event with a particular previously learned characteristic, which indicates a close relationship between semantic and episodic memories. It is also possible that, through repetition and rehearsal, new information could be abstracted from its episodic context and represented as semantic memory. In addition, it has been demonstrated that amnesic patients with lesions in the medial temporal lobes have impaired acquisition of new semantic memories. Our results concur with those of a recent VBM study in patients with early AD, which also found strong GMD correlations in the medial temporal structures with naming performance, mainly with the most anterior part of the parahippocampus and other parts of the perirhinal cortex. As proposed by these authors, the primary role of this region is combining the different representations of a given object, as part of a process of multimodal synthesis spread across different cortical areas. Thus, lesion of these brain structures in early AD would isolate the hippocampus from the multisensory input of the neocortex, resulting in reduction of retrieval efficiency, rather than loss of representation.

The role of the thalamus in lexical-semantic memory is less well understood than that of other significant areas demonstrated in our study. Recent electrophysiological and functional neuroimaging studies have established the involvement of the thalamus in the process of feature binding, which results in the recall of the object in semantic memory. Researchers have proposed that the thalamus could modulate the mechanism for semantic object recall via synchronizing of electrical brain rhythms. Kraut et al. (2003) studied a word-word feature-binding task using event-related fMRI. They found two distinct loci of thalamic signal change, one anterior in the dorsomedial nucleus, and the other posterior in the pulvinar. Based on these findings and previous electrophysiological studies, the authors proposed a neural mechanism in which the dorsomedial nucleus is involved in the early search or object generation and activates other prefrontal regions specifically involved in task-related working memory or language functions. Our findings support the idea that the thalamus is directly involved in lexical-semantic memory activities, possibly with an integrative role, since its nuclei were correlated with both BNT total score and semantic errors.

The involvement of neocortical temporal regions in semantic memory is better understood and has been extensively demonstrated. Grossman et al. (2004) studied VBM and confrontation naming in AD, frontotemporal dementia, and corticobasal degeneration, and found a left

![Figure 1](image1.png) **Figure 1.** Areas exhibiting significant correlation with BNT total score, predominantly in left superior frontal gyrus, left inferior frontal gyrus, left anterior temporal pole and bilateral thalamus (p<0.05, FWE corrected).

![Figure 2](image2.png) **Figure 2.** Areas exhibiting significant correlation with spontaneous semantic errors on BNT. All slices are in neurological orientation (left on the left side) p<0.05, FWE corrected.
lateral temporal atrophy as a common source of impaired naming across these patient groups. Another VBM study of semantic dementia showed that ATL activation peaks aligned closely with areas of strongest grey matter reduction, mostly with atrophy of the left anterior temporal lobe. We found correlations, particularly in the anterior parts of the STG, bilaterally but stronger on the left side, and in the anterior parts of the ITG, albeit weaker and less spread out than in the STG. Our findings support the notion that the anterior temporal lobe (ATL), predominantly its superior part, is strongly related to semantic generalization, since the subjects were asked to name pictures of different categories and a close relationship was observed between semantic errors, regardless of their specific categories.

The prefrontal cortex is also related to the lexical-semantic system, often in an asymmetrical manner, with the left side more involved than the right. The left inferior prefrontal cortex (LIPFC) has been regarded as a “semantic working memory system” responsible for retrieving, maintaining, monitoring, and manipulating semantic representations stored elsewhere, as evidenced by functional neuroimaging, transcranial magnetic stimulation, and lesion studies. 

On functional imaging studies, the LIPFC is more active when subjects make semantic judgments regarding words than when they make non-semantic judgments for the same words, and also when they make semantic judgments for line drawings. The role of the LIPFC is crucial when the semantic tasks require cognitive control of semantic or lexical retrieval, particularly during selection among competing alternatives. A study suggests that the LIPFC does not support retrieval of semantic knowledge per se. Rather, this retrieval is done entirely by the posterior neocortex based upon cues presented through bottom-up processes, and the specific role of the LIPFC would be to select those retrieved representations that are task-relevant from among competing, irrelevant representations.

Patients with left prefrontal lesions often have difficulty retrieving words in response to specific cues (e.g. words beginning with a specific letter or names of objects belonging to a specific semantic category), even when there is no aphasia. In such cases, making a semantic error (for example, naming “animal” instead of the target word “dog”) might indicate difficulty in selecting the appropriate phonological response to answer a particular semantic question. In fact, activation of the LIPFC has been elicted by phonological tasks such as discrimination of visually and auditorily presented words with the greatest activation more posteriorly near Broca’s area. These and other studies have also suggested a domain-specificity of the anterior LIPFC (BA 45/47) for controlled semantics and of the posterior LIPFC (BA 44/6) for controlled phonology. However, more recent studies have argued against domain-specificity and for domain-preferentiality in LIPFC. Thus, it may be hypothesized that the LIPFC is activated to the extent that lexical and semantic information must be rehearsed, temporarily stored, and selected in working memory to perform a particular task.

Our study has some limitations including the relatively small sample size and the fact that the BNT is not well balanced in terms of psycholinguistic variables. Despite this factor, the BNT is one of the most widely used naming tests in clinical practice, and it continues to be a well-accepted measure of naming impairment in brain-damaged patients. Notwithstanding the limitations of this study, we found evidence that several brain areas are related to the process of higher-order semantic generalization, particularly the thalamus, medial temporal lobe, prefrontal cortex (left more than right), and bilateral anterior temporal lobes (predominantly STG and ITG).

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