Cognitive fluctuations in connection to dysgraphia: a comparison of Alzheimer’s disease with dementia Lewy bodies

Emanuela Onofri
Marco Mercuri
Giuseppe Donato
Serafino Ricci
Department of Anatomy, Histology, Legal Medicine and Orthopaedics, Sapienza University of Rome, Rome, Italy

Background: The purpose of the present study was to examine the relationship between cognitive impairment and the performance of handwritten scripts presented as “letter-writing” to a close relative by patients with dementia Lewy bodies (DLB), as fluctuations of the symptoms phase, and in a matched group of patients with Alzheimer’s disease (AD). The degree of writing disability and personal, spatial, and temporal orientation was compared in these two groups.

Design and methods: Fourteen simple questions, designed in a form that could be utilized by any general practitioner in order to document the level of cognitive functioning of each patient, were presented to 30 AD patients and 26 DLB patients. The initial cognition test was designated PQ1. The patients were examined on tests of letter-writing ability. Directly after the letter-writing, the list of 14 questions presented in PQ1 was presented again in a repeated procedure that was designated PQ2. The difference between these two measures (PQ1 – PQ2) was designated Δ. This test of letter-writing ability and cognitive performance was administered over 19 days.

Results: Several markedly strong relationships between dysgraphia and several measures of cognitive performance in AD patients and DLB patients were observed, but the deterioration of performance from PQ1 to PQ2 over all test days were markedly significant in AD patients and not significant in DLB patients. It is possible that in graphic expression even by patients diagnosed with moderate to relatively severe AD and DLB there remains some residual capacity for understanding and intention that may be expressed. Furthermore, the deterioration in performance and the differences noted in AD and DLB patients may be due to the different speed at which the process of the protein degradation occurs for functional modification of synapses.

Conclusion: Our method can be used as part of neuropsychological tests to differentiate the diagnosis between AD and DLB.

Keywords: cognition, deficits, correlation, deterioration

Introduction

In Western industrial nations, Alzheimer’s disease (AD) represents the most common form of dementia, at approximately 80%, occupying fourth place among the causes of death (after heart disease and circulatory disorders, cancer, and cerebral hemorrhage). Dementia with Lewy bodies (DLB) is the second most common cause of degenerative dementia after AD.

Worldwide there are currently 25 to 35 million individuals with these illnesses, with five to seven million new cases diagnosed each year, or one new case every 7 seconds. According to the World Health Organization it has been estimated that by the year 2050 the number of people presenting with dementia will be 115.4 million.
The pathology of dementia is associated with memory loss, loss of orientation, inability to focus attention, and loss of speech. In addition, we notice cognitive fluctuations, defined as spontaneous alterations in cognition, attention, and arousal. Fluctuating confusion, accompanied by disturbances of consciousness, is an important clinical symptom, with a frequency of 80%–90% in DLB and 20% in Alzheimer’s disease. In fact, in these patients, cognitive fluctuations and periods of behavioral confusion, inattention, and alternating episodes of lucidity and capable task performance have been described. McKeith identified the “marked amplitude between best and worst performance” as a distinguishing feature of the fluctuations in cognition and consciousness in DLB, contrasting this with “minor day-to-day variations” that can occur in dementia of any cause. The term cognitive fluctuation remains elusive despite several attempts to identify, quantify, and assess the phenomenon.

This fact raises an important question: do the fluctuations occurring in AD patients have a particular quality and specific characteristics different from fluctuations that occur in DLB dementia?

In addition, many authors have observed in patients with AD dementia, a progressive disorganization and degeneration of the various components of handwriting, the diversity and the accuracy of words used, organization, punctuation, the production of grammatically incorrect sentences, the length of the sentences, the morphology of the letters, the amount of written information, spelling, and graphic and spatial layout of letters and their arrangement in texts.

We are not aware of research regarding the degeneration of handwriting in DLB patients. The purpose of the present study was to examine the relationship between cognitive impairment and the performance of handwritten scripts presented as “letter-writing” to a close relative by AD patients, as fluctuations of the symptoms phase, and in a matched group of DLB patients.

Also, it is possible that in graphic expression even by patients diagnosed with moderate to relatively severe AD and DLB there remains some residual capacity for understanding and intention that may be expressed. Concurrently, the relationship between measures of functioning, Mini-Mental State Examination (MMSE), and number of years elapsed since first indication of disorder with dysgraphia and assessments of cognitive performance was assessed (or debut).

Materials and methods

Fifty-six patients (from the region of Lazio, Italy) were selected to participate in the study: 30 patients presenting AD and 26 patients presenting DLB. Patients were diagnosed according to the National Institute of Neurological and Communicative Disorders and Stroke and the Alzheimer’s Disease and Related Disorders Association and the Diagnostic and Statistical Manual of Mental Disorders, 4th edition reference.

In addition, a group of 30 younger, less impaired AD patients (ADli) was included. Patients were excluded if they presented a history of known or suspected cerebrovascular disease, focal neurological signs or brain imaging evidence for alcohol misuse, head trauma, significant psychiatric history preceding the current diagnosis, or any other major physical illness. All of these diagnoses were confirmed and verified by neurologists in the Department of Neurology at the hospital (Gemelli University Polyclinic – service neuropsychology, Roma and the Alzheimer Evaluation Unit ASLRMF and Alzheimer Evaluation Unit ASLRMD); 26 patients were confirmed by the Department of Neurology and Psychiatry, Sapienza Hospital, Rome, Italy, in the Lazio region. Only subjects with the clinical follow-up and complete diagnostic agreement by the three neurologists were included.

A considerable amount of time (regular meetings every second day for 3 months) was invested in each of the patients in order to promote a relationship of trust and understanding, as well as to reduce stress factors that may affect patients’ mood and attentiveness, or, more seriously, induce behaviors that suggest hallucinations or auditory illusions, paranoid delirium, difficulty in recognizing persons, or loss of cognition of time and place. All the procedures were adopted according to discussions and meetings with nearest relatives and caregivers in order to obtain the consent of the patients as well as those of relatives and caregivers (caregivers could be legal representatives) according to the legal practices.

Alzheimer’s disease

For the sixty patients constituting the AD group and ADli, the diagnosis was based on normal or nonspecific electroencephalography and lateral, occipital brain atrophy on magnetic resonance imaging or computed tomography brain scan, with documented progression after serial observations, and on cognitive test procedures and routine blood assays aimed at excluding the presence of other medical conditions associated with other types of dementia.
Dementia with Lewy bodies

Twenty-six DLB patients were included in the study. Diagnosis was confirmed using the international consensus guidelines for DLB, revised in 2005. All patients presented with dementia at the time of their initial evaluation, which usually preceded the onset of motor signs of disorder by 1–2 years. Some patients exhibited cognitive impairment concomitant with mild extrapyramidal signs, which, however, never reached the criteria levels for the diagnosis of idiopathic Parkinson’s disease, and showed no evidence of focal brain lesions on magnetic resonance imaging.

Exclusion criteria were presence of secondary, reversible causes of dementia that were untreated; concomitant neurological or psychiatric illness, substance use and abuse; and history of significant head trauma.

Mini-Mental State Examination

MMSE presents a brief 30-point questionnaire to screen for cognitive impairment and dementia. It estimates the severity of disorder and follows the course over time of cognitive changes in an individual, thereby allowing effective monitoring of an individual’s response to treatment.

Unified Parkinson’s Disease Rating Scale

The Unified Parkinson’s Disease Rating Scale is a rating scale used to follow the longitudinal course of Parkinson’s disease. It is the most commonly used scale in the clinical study of Parkinson’s disease and of DLB.

Clinician assessment of fluctuation (CAF)

CAF is a short scale consisting of a series of screening questions, regarding fluctuating confusion and impaired consciousness during the month prior to the assessment. The frequency and duration of episodes of fluctuating confusion are both rated on a scale of 0–12 (0 representing no fluctuating confusion, 12 representing severe fluctuating confusion).

Cognitive performance assessment

Testing material

A writing pad was used for writing text or drawing figures. Preferably the tablet was vergatina type (flimsy typing paper, tissue type, or absorbent in order to avoid false interpretations of the writings). A ballpoint pen was used throughout. The patients were invited to sit in a comfortable position.

Table 1 Example scoring list of 14 questions presented to a hypothetical AD patient in PQ1 phase and PQ2 phase. These are routine questions which a doctor can ask to the patient

<table>
<thead>
<tr>
<th>Questions</th>
<th>Score</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. What is your name?</td>
<td>0–3</td>
</tr>
<tr>
<td>2. When were you born?</td>
<td>0–2</td>
</tr>
<tr>
<td>3. Where were you born?</td>
<td>0–2</td>
</tr>
<tr>
<td>4. Which city do you live?</td>
<td>0–1</td>
</tr>
<tr>
<td>5. What is your address?</td>
<td>0–1</td>
</tr>
<tr>
<td>6. Who are you living with?</td>
<td>0–1</td>
</tr>
<tr>
<td>7. Have you got children?</td>
<td>0–1</td>
</tr>
<tr>
<td>8. What are their names?</td>
<td>0–2</td>
</tr>
<tr>
<td>9. Why are you here?</td>
<td>0–1</td>
</tr>
<tr>
<td>10. Have you pain?</td>
<td>0–1</td>
</tr>
<tr>
<td>11. Where have you pain?</td>
<td>0–1</td>
</tr>
<tr>
<td>12. How did you get here?</td>
<td>0–1</td>
</tr>
<tr>
<td>13. How much time did you take to get here?</td>
<td>0–1</td>
</tr>
<tr>
<td>14. What day is today?</td>
<td>0–2</td>
</tr>
</tbody>
</table>

Abbreviations: AD, Alzheimer’s disease; PQ, cognition test.

Procedure

In order to test the cognitive performance of AD patients and DLB patients as well as test their spatial and time orientation and residual capacity, a standard collection of 14 simple questions were presented to the patients (Table 1). The authors have designed and presented these questions in a form that could be utilized by any general practitioner in order to document the level of cognitive functioning of each patient. For each correct answer, one point was attributed in proportion to the difficulty of the question. The sum of each test session was represented by PQ; the initial session result was designated PQ1. Following this, each patient was then invited to write a letter to a close relative. On consecutive days of testing, patients were invited to write to either the same relative or another one. The letter-writing task was interrupted when it seemed that the text produced by the patient was substantially (pathologically) confused (when the phrase offered no conceptual association with accompanying text although in the presence of otherwise “correct” syntax) and when it had a sudden lack of readability, disjointedness, and incompleteness in meaning, with intrusions, semantic substitutions, alterations in the spatial organization of handwriting, illegible words, incidence of paraphrases, incapacity to form complete sentences, graphemic substitutions (a grapheme is the smallest semantically distinguishing unit in a written language), omissions, and additions. After this, using a chronometer, the number of minutes that had been reached for each single patient was registered and complete sentences were counted (sentences/minutes = XF). The whole procedure involving the letter-writing graphia task was interrupted after 20 minutes (see Figure 1).
Directly after letter-writing

The list of 14 questions presented in PQ1 was presented again in a repeated procedure that was designated PQ2. The difference between these two measures (PQ1 – PQ2) was designated Δ. These procedures for testing: 14-item test, graphia test, 14-item test were presented in an identical manner every second day over 10 days (days 1, 3, 5, 7, 9, 11, 13, 15, 17, and 19) at the same hour of day on test days in order to hold constant testing procedures over daily curriculum and any clinical interventions that the patients may be subject to.

Statistical analysis

The results consisting of PQ1 and PQ2 scores, XF, and Δ were calculated as means and standard deviations of the AD patient group and the DLB patients over consecutive days of testing. Pearson’s correlation coefficient was used to assess the relationship between XF and PQ1 and XF and Δ for both AD patients and DLB patients for each of the test days. Student’s t-tests were used to test for pairwise differences for each of the parameters.

Results

The clinical characteristics of the participants in the study are presented in Table 2.

AD, ADli, and DLB patients: cognitive performance over PQ1 and PQ2

The cognitive performance of the AD patients and DLB patients deteriorated from PQ1 to PQ2 (Table 3), with t-values reaching significance levels over all 10 days of testing.

The cognitive performance of ADli patients deteriorated from PQ1 to PQ2 (Table 4).

Comparison of AD, ADli with DLB patients: correlation coefficients PQ1 versus XF and XF versus Δ

The correlation coefficients of both the PQ1 performance and XF relationship and the XF and Δ relationship were all positive and highly significant in AD patients while they were positive in ADli patients. In DLB patients the correlation coefficient of the PQ1 performance and XF relationship was positive while the correlation analysis of the XF and Δ relationship was negative (Tables 3 and 4).

Correlation analyses between functional measures

Product moment correlations between the functional estimations of MMSE, debut (years), PQ1, Δ, and PQ2 were

Table 2 Clinical, demographic, and neuropsychological data of the study groups

<table>
<thead>
<tr>
<th></th>
<th>AD</th>
<th>DLB</th>
<th>AD younger</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sex (M/F)</td>
<td>40% M 60% F</td>
<td>45.15% M 54.85% F</td>
<td>35% M 55% F</td>
</tr>
<tr>
<td>Mean age (years)</td>
<td>83.06±6.15</td>
<td>78±6.4</td>
<td>77.4±5.8</td>
</tr>
<tr>
<td>Educational (years)</td>
<td>11.0±3.6</td>
<td>10.3±2.3</td>
<td>11.3±2.6</td>
</tr>
<tr>
<td>Debut (years)</td>
<td>4.43±1.19</td>
<td>3.3±0.6</td>
<td>3.4±1.23</td>
</tr>
<tr>
<td>MMSE</td>
<td>13.75±1.96</td>
<td>14.3±1.5</td>
<td>20.5±1.6</td>
</tr>
<tr>
<td>UPDRS</td>
<td>0</td>
<td>61.87±2.5</td>
<td>0</td>
</tr>
<tr>
<td>CAF</td>
<td>2.5</td>
<td>6.7</td>
<td>0</td>
</tr>
</tbody>
</table>

Notes: Values are reported as mean ± standard deviation; Debut is defined as number of years elapsed since first indication of disorder.

Abbreviations: AD, Alzheimer’s disease; CAF, clinician assessment of fluctuation; DLB, dementia Lewy bodies; F, female; M, male; MMSE, Mini-Mental State Examination; UPDRS, Unified Parkinson’s Disease Rating Scale.
Table 3 The Student’s t-test values between PQ1 and PQ2 scores (means ± SD) and Pearson product moment correlational analyses between 1) PQ1 and XF, 2) XF and D∆ over 10 days of testing (days 1, 3, 5, 7, 9, 11, 13, 15, 17, and 19)

<table>
<thead>
<tr>
<th>Test days</th>
<th>PQ1 AD</th>
<th>PQ1 DLB</th>
<th>PQ2 AD</th>
<th>PQ2 DLB</th>
<th>t-values AD</th>
<th>t-values DLB</th>
<th>R (1) AD</th>
<th>R (1) DLB</th>
<th>R (2) AD</th>
<th>R (2) DLB</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>11.9±4.8</td>
<td>10.6±4.9</td>
<td>4.0±1.9</td>
<td>10.3±4.5</td>
<td>8.25±3</td>
<td>0.24</td>
<td>0.802±3</td>
<td>0.919</td>
<td>0.905±3</td>
<td>0.455</td>
</tr>
<tr>
<td>3</td>
<td>10.5±2.3</td>
<td>10.2±4.5</td>
<td>4.1±2.3</td>
<td>10.2±4.4</td>
<td>7.11±3</td>
<td>0.17</td>
<td>0.785±3</td>
<td>0.877</td>
<td>0.669±3</td>
<td>0.099</td>
</tr>
<tr>
<td>5</td>
<td>9.9±5.1</td>
<td>9.4±4.7</td>
<td>4.8±3.0</td>
<td>9.0±4.5</td>
<td>4.71±3</td>
<td>0.27</td>
<td>0.630±3</td>
<td>0.892</td>
<td>0.680±3</td>
<td>0.179</td>
</tr>
<tr>
<td>7</td>
<td>9.5±4.6</td>
<td>8.0±1.1</td>
<td>5.2±2.6</td>
<td>7.8±3.9</td>
<td>4.99±3</td>
<td>0.22</td>
<td>0.644±3</td>
<td>0.824</td>
<td>0.755±3</td>
<td>0.192</td>
</tr>
<tr>
<td>9</td>
<td>11.8±3.1</td>
<td>8.9±3.4</td>
<td>5.7±2.1</td>
<td>8.7±3.4</td>
<td>9.04±3</td>
<td>0.29</td>
<td>0.732±3</td>
<td>0.861</td>
<td>0.588±3</td>
<td>0.297</td>
</tr>
<tr>
<td>11</td>
<td>8.9±4.4</td>
<td>7.3±3.8</td>
<td>5.0±2.8</td>
<td>7.0±3.8</td>
<td>4.04±3</td>
<td>0.23</td>
<td>0.556±3</td>
<td>0.855</td>
<td>0.646±3</td>
<td>0.081</td>
</tr>
<tr>
<td>13</td>
<td>9.3±4.3</td>
<td>7.4±4.1</td>
<td>5.5±2.7</td>
<td>7.1±4.1</td>
<td>4.08±3</td>
<td>0.24</td>
<td>0.688±3</td>
<td>0.846</td>
<td>0.542±3</td>
<td>0.197</td>
</tr>
<tr>
<td>15</td>
<td>8.9±4.4</td>
<td>7.6±4.2</td>
<td>5.1±2.9</td>
<td>7.5±4.2</td>
<td>3.93±3</td>
<td>0.03</td>
<td>0.498±3</td>
<td>0.918</td>
<td>0.643±3</td>
<td>0.122</td>
</tr>
<tr>
<td>17</td>
<td>9.5±4.0</td>
<td>8.3±3.3</td>
<td>5.4±3.1</td>
<td>8.2±3.4</td>
<td>4.42±3</td>
<td>0.07</td>
<td>0.767±3</td>
<td>0.869</td>
<td>0.547±3</td>
<td>0.176</td>
</tr>
<tr>
<td>19</td>
<td>9.5±5.3</td>
<td>8.3±4.5</td>
<td>4.5±3.4</td>
<td>8.1±4.7</td>
<td>4.26±3</td>
<td>0.19</td>
<td>0.871±3</td>
<td>0.853</td>
<td>0.576±3</td>
<td>-0.142</td>
</tr>
</tbody>
</table>

Notes: *t*-tests (df=54); P<0.0001; *P<0.005; **P<0.002; ***P<0.001.

Abbreviations: AD, Alzheimer’s disease; D∆, difference between the measures of PQ1 and PQ2; DLB, dementia Lewy bodies; PQ1, initial cognition test; PQ2, repeated cognition test; SD, standard deviation; XF, number of complete sentences/minutes reached for each patient.

Discussion

This study shows the distinct features of the cognitive impairment following a letter-writing task (PQ1 – PQ2) and dysgraphia (XF) in a group of AD patients presenting with mild cognitive decline, and dementia (PQ1) in AD patients. The results may be summarized as follows: 1) the deterioration in cognitive performance following a letter-writing task (PQ1 – PQ2) and dysgraphia (XF) in AD patients represent a moderate to relatively severe stage of disorder, and 2) the deterioration in cognitive performance is quite advanced in comparison with both AD and DLB patients.

The results may also be analyzed in the following way: 1) the deterioration in cognitive performance and dysgraphia (XF) in AD patients present with a moderate to severe stage of disorder, and 2) the deterioration in cognitive performance is quite advanced in comparison with both AD and DLB patients.

Comparison AD patients with DLB patients

The cognitive impairment that occurs after the writing performance in AD patients is clearly shown in Figure 2. The blue plot shows the levels of cognitive impairment that occurred before the writing performance in AD and DLB patients (over 10 days of testing) day after day. The red plot emphasizes the change in performance from PQ1 to PQ2 over all test days was markedly strong in AD patients, and it was not significant in DLB patients (Figure 4).

The extent of dysgraphia by the DLB patients was found to be quite advanced in comparison with the AD patients (Figure 5).
In a sample of 52 Japanese patients presenting mild AD and 22 healthy controls, writing ability composed of Kana writing-to-dictation and copying Kanji or dictational Kanji and regional cerebral blood flow using single-photon emission computed tomography were studied.44 Hayashi et al observed that while Kana writing-to-dictation and copying Kanji were preserved in these AD patients, writing to dictated Kanji words was impaired. The impaired writing of dictated Kanji words was associated with dysfunctional cortical activity, predominantly in the left frontal, parietal, and temporal brain regions, consistent with other Japanese dysgraphia studies.35,36

Furthermore, clear differences between AD patients and healthy control individuals have been found for visuomotor task measures, demonstrating large effect size deficits by AD patients, especially with visuomotor task progression through its varying conditions.37,38 They have observed dysgraphia, primarily dysexecutive agraphia, in patients with frontotemporal dementia and parkinsonism linked to chromosome 17. Patients presenting dysexecutive agraphia show not only difficulties in maintaining the effort inherent to writing but appear to lack the ability to organize their thoughts for expression in written text. They seem to be lacking in the complicated functions underlying writing that encompass narrative coherence, selective attention, planning, etc that are disturbed in executive function impairments.

The present observations of dysgraphia associated with deficits in semantic memory that were exacerbated acutely by the writing task appear to fit current notions pertaining to the progressive performance impairments of AD patients within language and cognition domains from a staging perspective.39

The possible relationships between dysgraphia and the motor functioning domain in AD has provided novel insights.

4) MMSE correlated significantly with PQ1, PQ2, and with XF in both groups of patients; furthermore, number of years elapsed since first diagnosis of disorder does not affect the XF (or expression of dysgraphia), PQ1, PQ2, and DΔ. In particular, the nonassociation between MMSE and DΔ in DLB patients was markedly significant ($R^2 = -0.185$).

Dysgraphia occurs during both the earlier as well as the later stages during the clinical course of AD26-28 and is associated with attentional, motor, and memory deficits that develop during disorder progression.29 It has been suggested to be a more sensitive indication of language deficits in AD than anamnia.30,31 Some authors32 have observed that the medial walls of the cerebral hemispheres, notably the cingulate gyri, contain species-specific neuron fields that to date are not well known within the scientific community and yet have been implicated as the underlying cause of such varying conditions as dysgraphia and autism in children and obsessive compulsive disorder and AD in adults.33

Table 4 Younger, less impaired AD patient Student’s t-test values between PQ1 and PQ2 scores (means ± SD) and Pearson product moment correlational analyses between 1) PQ1 and XF, 2) PQ2 and DΔ over 10 days of testing (days 1, 3, 5, 7, 9, 11, 13, 15, 17, and 19)

<table>
<thead>
<tr>
<th>Test days</th>
<th>PQ1 AD</th>
<th>PQ2 AD</th>
<th>t-values AD</th>
<th>R (1) AD</th>
<th>R (2) AD</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>16.0±2.1</td>
<td>10.3±1.8</td>
<td>10.92</td>
<td>0.578 O</td>
<td>0.533 O</td>
</tr>
<tr>
<td>3</td>
<td>15.6±2.5</td>
<td>9.7±2.0</td>
<td>10.29</td>
<td>0.656 O</td>
<td>0.536 O</td>
</tr>
<tr>
<td>5</td>
<td>15.3±2.7</td>
<td>9.7±1.9</td>
<td>10.28</td>
<td>0.616 O</td>
<td>0.578 O</td>
</tr>
<tr>
<td>7</td>
<td>15.3±2.7</td>
<td>10.0±1.8</td>
<td>8.84</td>
<td>0.689 O</td>
<td>0.696 O</td>
</tr>
<tr>
<td>9</td>
<td>15.4±2.6</td>
<td>10.0±1.9</td>
<td>8.84</td>
<td>0.698 O</td>
<td>0.663 O</td>
</tr>
<tr>
<td>11</td>
<td>15.3±2.7</td>
<td>9.5±1.7</td>
<td>9.93</td>
<td>0.602 O</td>
<td>0.536 O</td>
</tr>
<tr>
<td>13</td>
<td>16.9±2.5</td>
<td>9.6±1.5</td>
<td>14.81</td>
<td>0.657 O</td>
<td>0.576 O</td>
</tr>
<tr>
<td>15</td>
<td>15.3±2.7</td>
<td>10.0±1.9</td>
<td>8.84</td>
<td>0.603 O</td>
<td>0.553 O</td>
</tr>
<tr>
<td>17</td>
<td>15.5±2.7</td>
<td>10.0±1.8</td>
<td>8.84</td>
<td>0.651 A</td>
<td>0.635 A</td>
</tr>
<tr>
<td>19</td>
<td>15.4±2.7</td>
<td>9.8±1.9</td>
<td>9.45</td>
<td>0.695 O</td>
<td>0.586 O</td>
</tr>
</tbody>
</table>

Notes: *t*-tests (df=58); $P<0.0001$; *$P<0.005$; **$P<0.002$; ***$P<0.001$.

Abbreviations: AD, Alzheimer’s disease; DΔ, difference between the measures of PQ1 and PQ2; PQ1, initial cognition test; PQ2, repeated cognition test; SD, standard deviation; XF, number of complete sentences/minutes reached for each patient.

Table 5 Pearson product moment correlational analyses between MMSE, debut (years), PQ1, XF, PQ2, and DΔ in the 30 AD patients and 26 DLB patients

<table>
<thead>
<tr>
<th>AD patients</th>
<th>DLB patients</th>
</tr>
</thead>
<tbody>
<tr>
<td>MMSE – PQ1</td>
<td>0.791</td>
</tr>
<tr>
<td>MMSE – XF</td>
<td>0.790</td>
</tr>
<tr>
<td>MMSE – PQ2</td>
<td>0.788</td>
</tr>
<tr>
<td>MMSE – DΔ</td>
<td>0.665</td>
</tr>
<tr>
<td>Debut (years) – PQ1</td>
<td>−0.632</td>
</tr>
<tr>
<td>Debut (years) – XF</td>
<td>−0.584</td>
</tr>
<tr>
<td>Debut (years) – PQ2</td>
<td>−0.604</td>
</tr>
<tr>
<td>Debut (years) – DΔ</td>
<td>−0.554</td>
</tr>
</tbody>
</table>

Note: Debut is defined as number of years elapsed since first indication of disorder.

Abbreviations: AD, Alzheimer’s disease; DΔ, difference between the measures of PQ1 and PQ2; DLB, dementia Lewy bodies; MMSE, Mini-Mental State Examination; PQ1, initial cognition test; PQ2, repeated cognition test; XF, number of complete sentences/minutes reached for each patient.
into the cognitive nature of the disorder, through which mild to moderate stage AD patients (n=59) and healthy elderly controls were tested over an extensive assessment of both the central and peripheral components of writing; the former performed less effectively than controls over a broad spectrum of writing measures. Although a predominantly lexical disorder was observed, there were multiple indications of associated disorders located at different stages in the writing/spelling system (eg, phonological route, graphemic buffer, allographic store, graphic motor patterns). The authors concluded that there exists heterogeneous profiles of dysgraphia with primary signs of writing impairment in AD originating from changes at different points in the brain networks that subserve writing and spelling performance.

In this regard, the possibility of related motor deficits in dysgraphia ought to be considered since there is evidence for altered parietal-motor connections in AD. It has been found too that sensory-motor plasticity is impaired in the motor cortex of AD at an early stage of the disease.

Several have indicated that agraphia is related to severity of Alzheimer’s disease. Hughes et al have demonstrated the relationships between extent of MMSE impairment and patterns of agraphia in AD. MMSE deficits in AD are linked to several comorbidity domains, including cardiovascular, ear, nose and throat, genitourinary, musculoskeletal/integument, and neurologic.

We are not aware of studies that have examined the relationship between cognition and dysgraphia in DLB patients, nor of studies that have compared the cognitive impairment after writing between AD and DLB. Much of the current knowledge regarding the relationship between DLB and AD dementia and cognition is derived from clinical–pathological studies of atypical cases of dementia and specialty clinic samples. For instance clinic-based studies have shown that cortical Lewy body pathology is associated with more rapid decline in cognition; however, this has not been noted in all studies.

Cognitive fluctuations that occur in patients cannot be caused by sudden loss or gain of nerve cells; recent evidence...
suggests that variations in the activity of neural networks, and the neural plasticity is mediated by morphological and functional modification of synapses, a process that depends on both synthesis and degradation of proteins, events targeting a variety of molecules in pre- and postsynaptic compartments.

This study, by analyzing relationships between initial cognitive performance (PQ1), the deterioration in cognitive performance following a letter-writing task (PQ2 = DA), and dysgraphia in AD and DBL patients, suggests that cognitive fluctuations that occur in patients with AD during the performance of handwritten scripts are not present in patients with DBL. During the observation period, the DBL patients have fluctuating confusion, inattention that is independent from the fatigue of the writing task.

May the differences noted in AD and DBL patients be due to the different speed at which the process of the protein degradation occurs? Is it the recovery of cognition in AD and DBL patients allowed to the influence of protein degradation in relation to temporal dynamics in other regions of the brain? The current study has a limitation: cognitive fluctuation was measured on majority of change in orientation, not other cognitive domains, approximately 20 minutes apart.

Conclusion

Our results can provide new insights into the mechanism of cognitive decline in AD and DBL patients and may be useful to develop a method based on dysgraphia in order to differentiate the diagnosis of both types of dementia. In particular, the writing, in association with the cognitive questionnaire, can be used as part of neuropsychological tests to diagnose the difference between AD and DBL.

Disclosure

The authors report no conflicts of interest in this work.

References


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