COGNITIVE IMPAIRMENT IN HUMAN CHRONIC CHAGAS' DISEASE


SUMMARY - We proposed to investigate subclinical cognitive impairment secondary to chronic Chagas' disease (CCD). No similar study was previously done. The neuropsychological performance of 45 chronic Chagasic patients and 26 matched controls (age, education place and years of residency in endemic area) was compared using the Mini Mental State Exam (MMSE), Weschler Memory Scale (WMS) and the Weschler Adult Intelligent Scale (WAIS). Non-parametric tests and Chi2 were used to compare group means and multivariate statistics in two way frequency tables for measures of independence and association of categorical variables with the disease. Results: Chagasic patients showed lower MMSE scores (p<.004), poor orientation (p<.004), and attention (p<.007). Lower WMS MQ were associated with CCD (Chi2 5.9; p<.01; Fisher test p<.02). Lower WAIS IQ were associated with CCD (Chi2 6.3, p<.01; Fisher test p<.01) being the digit symbol (p<.03), picture completion (p<.03), picture arrangement (p<.01) and object assembly (p<.03) subtests the most affected. The impairment in non-verbal reasoning, speed of information processing, problem solving, learning and sequencing observed in chronic Chagas disease patients resembles the cognitive dysfunction associated with white matter disease.

KEY WORDS: Chagas' disease, cognitive dysfunction, cerebral white matter involvement.

Chagas' disease is a South American trypanosomiasis. In Argentina 30% of the population is at risk and 5% is infected. Some reports have shown that chronic Chagas' disease (CCD) affects the peripheral nervous system. Involvement of the alpha spinal motoneurone and peripheral nerve trunks have been described in humans\textsuperscript{11,13}. An experimental mouse model, infected with different \textit{Trypanosoma cruzi} strains, has replicated the clinical observations\textsuperscript{5,6}. These last investigations

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revealed that the spinal motoneurone somae, the posterior root ganglia cells, the nerve axons or their myelin sheets were affected either alone or in combination. Chronic and acute forms of central nervous system (CNS) involvement (Chagas' encephalopathy) have been also described. IgG antineuron and antimyelin antibodies were detected in serum and cerebrospinal fluid in the acute stages as a consequence of an immunologic reaction (local release of neuronal and myelinic antigenic fractions and production of self-antibodies). These immunologic reactions may play an important role in the chronic CNS dysfunction in CCD.

Although most of the patient's with chronic Chagas' disease look bradyphrenic, not any study has assessed the cognitive performance in them. We have begun to study the CNS in these patients in order to investigate possible involvement of it in the chronic stages of the disease. As part of this research we have proposed a systematic study of the neuropsychological functions to seek subclinical cognitive involvement in these patients. In the present paper we discuss the data of the first 45 patients studied. Partial reports of these findings have been presented elsewhere.

**MATERIAL AND METHODS**

We studied the neuropsychological performance on 45 patients with chronic Chagas' disease that did not complain on cognitive loss and it was compared with that of 26 matched controls (age, education, place and years of residency in the same geographical area). CCD patients and controls were free of any other systemic or neurologic disorder that may cause cognitive decline.

Previous history of psychiatric disorders, chronic use of neuroleptics or antidepressants and alcohol or any other drug abuse were exclusion criteria for either group.

The Folstein Minimental State (MMSE), the Weschler Memory Scale (WMS) and the WAIS global test of Intelligence were used to score the cognitive performance.

**RESULTS**

Demographic characteristics of the population are shown in Table 1. Special care was taken in matching both groups. Patients and controls were from the same endemic area where they lived on for almost the same period of time. No differences were observed regarding sex, age and education.

Statistical difference (p<.004) was observed in the MMSE scores between controls (28.8±1.2) and CCD.

<table>
<thead>
<tr>
<th>Number</th>
<th>Patients</th>
<th>Controls</th>
</tr>
</thead>
<tbody>
<tr>
<td>45</td>
<td>26</td>
<td></td>
</tr>
<tr>
<td>Sex (Male)</td>
<td>19 (42%)</td>
<td>10 (39%)</td>
</tr>
<tr>
<td>Age (years)</td>
<td>36.9 ± 10.5</td>
<td>33 ± 10</td>
</tr>
<tr>
<td>Education (years)</td>
<td>7.8 ± 2.5</td>
<td>8.5 ± 3.6</td>
</tr>
<tr>
<td>Residency in endemic area (years)</td>
<td>15.7 ± 8</td>
<td>14.1 ± 10</td>
</tr>
</tbody>
</table>

Wilcoxon Non Parametric test (No differences between groups). @ (Chi2 statistics).

Statistical Analysis. Data were analyzed with the BMDP Statistical Software package. Differences between group means were assessed with the Wilcoxon non-parametric test for continuous variables and the Chi2 for categorical variables. Multivariate statistics in two way frequency tables were used to obtain measures of association and independence between the categorical variables (WAIS IQ and WMS MQ) and the presence or absence of Chagas' infection.

Table 2. WAIS scores in patients and controls.

<table>
<thead>
<tr>
<th></th>
<th>Patients</th>
<th>Controls</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Information</td>
<td>6.67 ± 2</td>
<td>8 ± 3.2</td>
<td></td>
</tr>
<tr>
<td>Comprehension</td>
<td>6.5 ± 2.5</td>
<td>7.7 ± 3.2</td>
<td></td>
</tr>
<tr>
<td>Arithmetics</td>
<td>6.1 ± 2</td>
<td>7.1 ± 2</td>
<td></td>
</tr>
<tr>
<td>Analogies</td>
<td>7.7 ± 3</td>
<td>9 ± 3</td>
<td></td>
</tr>
<tr>
<td>Digit span</td>
<td>6.2 ± 2.6</td>
<td>7.4 ± 2.7</td>
<td></td>
</tr>
<tr>
<td>Vocabulary</td>
<td>7.2 ± 1.8</td>
<td>8 ± 3</td>
<td></td>
</tr>
<tr>
<td>Digit symbols</td>
<td>6.2 ± 2</td>
<td>7.3 ± 2</td>
<td>&lt;.03</td>
</tr>
<tr>
<td>Picture compl.</td>
<td>6.2 ± 1</td>
<td>7.4 ± 2.7</td>
<td>&lt;.03</td>
</tr>
<tr>
<td>Block design</td>
<td>7.4 ± 2.4</td>
<td>8.3 ± 2.4</td>
<td></td>
</tr>
<tr>
<td>Picture arrang.</td>
<td>6.1 ± 2.7</td>
<td>8 ± 3.2</td>
<td>&lt;.01</td>
</tr>
<tr>
<td>Object assembly</td>
<td>6.4 ± 2.4</td>
<td>8 ± 3.1</td>
<td>&lt;.03</td>
</tr>
</tbody>
</table>

Wilcoxon Non Parametric test (differences between groups). Values for each variable are expressed as the x ± SD.
The WMS Associated Learning test showed differences (p<.002) between controls (11.6±2.5) and CCD (9.4±3.01). Only the Performance subtests of the WAIS showed statistical differences between groups (Table 2).

Multivariate analysis in 2 x 2 tables showed differences between controls and patients regarding the WAIS IQ and the WMS MQ. Most of the CCD patients had WAIS IQ lower than 90 (Table 3) and 35% of the controls showed WMS MQ higher than 90 while only 13% of CCD patients did.

The presence and absence of Chagas' infection was correlated with the WAIS IQ (using 90 as cutoff) in two way frequency tables (Table 4). A significant Pearson Chi2 statistic with a Fisher Exact test rejected the null hypothesis of independence of rows and columns. In 2 x 2 tables, the Phi-Cramer coefficient can be interpreted by paying attention to its sign. In this case it is negative; so, the lower values for the categorical variable WAIS IQ (IQ below 90) were associated with the higher code for the categorical variable disease (present=1).

The Memory Quotient MQ was evaluated in the same way and the null hypothesis of independence also rejected (Table 5).

**COMMENTS**

Previous observations have shown that patients with chronic Chagas' disease may have some degree of cognitive impairment. However, none of them could define a cognitive profile in these patients and propose a type of lesion which might underlay the process.

Our findings suggest that chronic Chagas' infection is highly correlated with low cognitive performance. We reduced confounders in the analysis of differences between groups matching them by age, education, place and years of residency in the same endemic region.

Familiar, well-learned, and over-practiced knowledge and skills (such as those generally represented by the WAIS Verbal IQ) did not show differences between healthy and infected individuals. Conversely, the non-verbal reasoning and the ability to solve novel and unfamiliar problems, as well as the capacity to plan new strategies (such as those represented in the WAIS Performance IQ) were impaired in the chagasic patients. Among these subtests, those that are timed measures such as picture completion, digit symbol, picture arrangement and object assembly, were the tests that best distinguished the groups.
CCD patients had less number responses in a time unit rather than increased number of wrong responses. These observations suggest that the differences between groups relay in the slowness of the patients to accomplish with the proposed tasks.

CCD patients were as able as controls in the final design the cubes but slower. As the cube design subtest did not show differences between groups in the execution of the task we can assume that perception and visuospatial planning were not affected and that impaired speed of information processing may be an important factor explaining the difference. This is also supported by the slow performance observed in the digit symbol task and leaves the impaired speed of information processing as the principal factor of the difference between groups.

The low performance observed in non verbal reasoning, orientation, problem solving and sequencing suggest frontal lobe dysfunction. The impairment in associative learning indicates deficits codifying new information. This neuropsychological picture resembles the reported cognitive dysfunction associated with white matter disease.

This last conclusion is in agreement with previous reports about evoked potentials that showed slowness of the central conduction time in some CCD patients when tested by somatosensory and auditive paradigms and with recent observations of Segura et al. (personal communication) who found that central reaction time studied with an oddball paradigm was impaired in some of these patients.

REFERENCES