Cognitive deficits in the Steele-Richardson-Olszewski syndrome (progressive supranuclear palsy)

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SUMMARY Clinical examination of 33 consecutive newly diagnosed cases of Steele-Richardson-Olszewski syndrome revealed evidence of cognitive impairment in 20. Eighteen of the 27 right handed patients who underwent neuropsychological assessment had intellectual impairment (mild in nine and marked in nine patients). The pattern of abnormalities was similar to but more severe than those previously reported in Parkinson’s disease, with particular difficulties in carrying out tests which are believed to be sensitive to frontal lobe dysfunction.

In 1964, Steele, Richardson and Olszewski reported the neuropathological and clinical features of a previously unrecognised syndrome which they called Progressive Supranuclear Palsy. Dementia, usually mild, was present in seven of the original nine patients and the presence of cognitive deficits and personality change have subsequently been confirmed. However, the exact frequency, severity and type of neuropsychological impairment is still controversial. Most studies have relied on clinical examination or impression to diagnose dementia and a large series of consecutive patients has never been studied. In studies in which neuropsychological testing has been carried out it has been claimed that the performance of patients with the Steele-Richardson-Olszewski (SRO) syndrome was only impaired on tests requiring visual scanning ability and that otherwise there was no evidence of intellectual or memory impairment. We therefore reviewed the neuropsychological reports in a group of newly diagnosed cases of SRO syndrome seen at the National Hospitals for Nervous Diseases between 1975 and 1984 to determine the incidence and extent of intellectual impairment.

Methods

The clinical records of 33 consecutive admissions to the National Hospitals for Nervous Diseases with a new diagnosis of SRO syndrome were reviewed retrospectively. All patients had unequivocal evidence of the syndrome, with supranuclear palsy affecting at least downward gaze and some or all of the following features: pseudobulbar palsy, axial and/or limb rigidity, bradykinesia, dementia, pyramidal signs and gait disorder. Neuropsychological assessment was performed on 29 patients, of whom 27 were known to be right-handed and these were selected for further examination. These 27 dextral patients had been studied in the Department of Psychology at the time of diagnosis. All patients were assessed on four verbal subtests (vocabulary, digit span, arithmetic and similarities) of the Wechsler Adult Intelligence Scale (WAIS), 20 were assessed on three performance subtests (picture completion, block design and picture arrangement) of the WAIS and an estimate of premorbid IQ was obtained in 19 patients from the National Adult Reading Test (NART). Seven patients did not complete the three WAIS performance subtests because of visual difficulties (oculomotor abnormalities in four patients, blepharospasm in two patients and one patient did not have her spectacles). The WAIS IQ and subtest scores given are age-corrected.

Verbal and visual memory was examined using a Recognition Memory Test. Twelve patients completed both sections of this test and seven completed the verbal section only because of visual impairment.

Supplementary tests performed by some patients included the Graded Difficulty Naming Test (nine patients), the Unusual Views test (14 patients) and the Fragmented Letters test (11 patients), for perceptual deficits, and two tests of reputed frontal lobe function: verbal fluency (number of words beginning with “s” in 90 seconds) (10 patients) and the Weigl test (11 patients). The number and type of supplementary tests administered to a subject was generally determined by the current neuropsychological testing protocol and not by specific clinical indications.

The clinical and neuropsychological findings in two
patients under the care of one of us (A.J.L) are reported in detail to illustrate some of the commoner features.

CT scans were performed and reported on in the Department of Neuroradiology. EEGs were recorded in the Department of Neurophysiology and were independently reviewed by Dr CJ Fowler and Dr MJG Harrison.

Statistical analyses comprised two-tailed t testing for paired data.

**Patient details**

The 27 patients (11 males and 16 females) were aged from 55 to 78 (median 65, mean 64·4) years of age and the median duration of symptoms before diagnosis was 3 years (range 1 to 9 years). Twenty-one patients had complained of mental symptoms: memory impairment (13 patients), emotional lability (7), depression (6) and slower mentation (4). Personality changes were reported by the relatives of four patients (apathetic or withdrawn: four patients, irritability in one patient).

Evidence of intellectual impairment and memory disturbance was considered to be present on clinical examination in 17 of the 27 patients. Clinical examination of the four patients who were not neuropsychologically assessed showed evidence of intellectual impairment in three (marked in two, mild in one patient).

Eight of the 27 patients were receiving neurotropic medication at the time of neuropsychological assessment (levodopa and decarboxylase inhibitor preparation six patients, benzhexol three patients and amantadine one patient).

CT scans were performed on all patients and showed evidence of cerebral atrophy and ventricular dilatation in 23 of the 27 patients (mild in 15 patients, moderate in seven patients and marked in one patient).

Seventeen patients had had an EEG recorded. None of the records were markedly abnormal. Eight records were normal, six were mildly abnormal (slight slowing or low amplitude alpha rhythm, minor scattered theta activity) and three were more notably abnormal (two showed asymmetrical excess theta activity and one showed non-specific frontal theta activity on hyperventilation.

**Results**

**WAIS testing**

The mean reading, verbal, performance and full-scale IQ scores are given in table 1. For the patients who completed both tests, there was a significant difference between the NART and verbal IQ scores (n = 19 t = 4·46 p < 0·001), the NART and Performance IQ scores (n = 16 t = 6·76 p < 0·001), the NART and full scale IQ scores (n = 17 t = 6·68 p < 0·001) and the verbal and performance IQ scores (n = 20 t = 3·74 p < 0·001).

The discrepancy between the NART and the Full-scale IQ (calculated from the verbal IQ for the three patients who had not attempted the performance scale) score was recorded as an index of deterioration in 19 patients. The mean (+SD) discrepancy was 13·3 (+10·1), six patients were consi-

<table>
<thead>
<tr>
<th>Test</th>
<th>No patients</th>
<th>Score mean</th>
<th>+SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>NART Score: (IQ equivalent)</td>
<td>19</td>
<td>113</td>
<td>+9·33</td>
</tr>
<tr>
<td>Verbal IQ</td>
<td>27</td>
<td>100·5</td>
<td>+15·6</td>
</tr>
<tr>
<td>Performance IQ</td>
<td>20</td>
<td>90·3</td>
<td>+13·8</td>
</tr>
<tr>
<td>Full scale IQ</td>
<td>20</td>
<td>94·3</td>
<td>+13·1</td>
</tr>
<tr>
<td>WAIS Subtest Score: Vocabulary</td>
<td>27</td>
<td>10·33</td>
<td>+2·79</td>
</tr>
<tr>
<td>Digit Span</td>
<td>27</td>
<td>9·19</td>
<td>+2·76</td>
</tr>
<tr>
<td>Arithmetic</td>
<td>27</td>
<td>8·26</td>
<td>+3·28</td>
</tr>
<tr>
<td>Similarities</td>
<td>27</td>
<td>8·26</td>
<td>+3·27</td>
</tr>
<tr>
<td>Picture Completion</td>
<td>21</td>
<td>6·76</td>
<td>+2·10</td>
</tr>
<tr>
<td>Block Design</td>
<td>22</td>
<td>5·91</td>
<td>+2·52</td>
</tr>
<tr>
<td>Picture Arrangement</td>
<td>20</td>
<td>4·21</td>
<td>+3·31</td>
</tr>
</tbody>
</table>

The discrepancy between the NART and the Full-scale IQ (calculated from the verbal IQ for the three patients who had not attempted the performance scale) score was recorded as an index of deterioration in 19 patients. The mean (+SD) discrepancy was 13·3 (+10·1), six patients were consi-
Memory

The verbal version of the Recognition Memory Test was administered to 19 patients and the visual version to 12 patients. On the verbal test the mean score (+SD) was 39 (±7.4), 12 patients scored at an average level or above, three were weak and four scored at a chance level indicating severe memory impairment. The mean score (+SD) on the visual test was 34.4 (±7.6). In only three patients was there a significant selective memory deficit on the visual version of the test and there were no instances of a selective verbal deficit.

Language functions

Mild word-finding difficulties were found in seven of the 25 patients, but there was no evidence of more severe dysphasia or comprehension difficulties. The Graded Difficulty Naming Test was given to nine patients. Two patients scored in the superior range, six in the average range and one in the dull-normal range.

Mental speed

In three patients marked slowness of thought out of proportion to intellectual impairment was noted. One patient was noted to have an involuntary tendency to speed up, often when speaking, but particularly when using her hands.

Perception

Visual Perceptual decoding processes were examined using the Unusual Views test (n = 14) and/or the Fragmented Letters test (n = 11). Nine of the 14 patients scored within normal limits on the Unusual Views test (mean 15-8/20) and nine of 11 patients scored within normal limits on the Fragmented Letters test (mean 17-8/20). The mean (+SD) Performance IQ and Picture Arrangement subtest scores for the 10 patients who scored within normal limits on the Unusual Views and/or Fragmented Letters was 92.7 (+11.9) and 3.9 (+3.48) respectively.

Frontal lobe test

The test of verbal fluency was administered to 10 patients and was failed by seven (all named 14 or less words beginning with “s” in 90 seconds). The Weigl test was administered to 11 patients and was failed by seven, one of whom also displayed reflex grasping. Reflex grasping was found in a further five patients, one of whom also exhibited utilisation behaviour.

Case reports

Case 1 A 59-year-old man presented with a 3 year history of progressive unsteadiness of gait with a tendency to fall backwards, dysarthria and dysphagia. For 6 months he had noticed diplopia in the vertical plane. His wife complained that there had been a gradual onset of mental slowing, emotional liability and inappropriate behaviour. On examination, he was noted to have a staring vacant expression and to hold his head rigid. He walked with a stiff upright gait and tended to lean backwards. Examination of higher mental function showed no definite abnormality although mentation was slow. There was no dysphagia. Ocular movements to command in the vertical plane were absent and following movements were limited to 10° upgaze and 5° downgaze. Horizontal eye movements were full but slow. Convergence was absent, but oculocephalic responses were full. A slurring dysarthria with limited palatal and spastic tongue movements, and an exaggerated gag reflex were present. The marked axial rigidity contrasted with the mild limb rigidity. Power and coordination were normal, but all tendon reflexes were pathologically increased with bilateral extensor planter responses and marked pout and facial reflexes also present. A CT scan showed mild generalised cerebral atrophy and an EEG was within normal limits.

Neuropsychology: NART = 101, verbal IQ = 80, performance IQ = 72, index of deterioration = 25. Analysis of subset scores showed that the highest scores were 8 for arithmetic and 7 for vocabulary. The lowest score was 0 for Picture Arrangement. He scored at chance level on both the verbal and visual versions of the Recognition Memory test (29/50 and 29/50), but scored within normal limits on the Fragmented Letters test for perceptual deficits. He was unable to sort Weigl shapes and perseverated on his first sorting.

Comment: Clinically this patient’s neurobehavioural changes were consistent with a “subcortical” pattern of dementia. Neuropsychological testing showed evidence of marked intellectual impairment, affecting non-verbal skills more than verbal skills, and of memory impairment. There was no evidence of dysphasia or perceptual deficits, but frontal lobe function was impaired.

Case 2 This 64-year-old man gave a 4 year history of progressive slowing and unsteadiness of gait. Three years before admission a diagnosis of Parkinson’s disease had been made and he had been started on Sinemet with little response. His speech became slurred, and he had noticed difficulty in looking down for six months. Examination revealed facial immobility and extreme neck rigidity. His gait was slow and shuffling with a tendency to fall backwards. Testing of higher mental function showed that he was fully orientated, but was slow and inaccurate on mental arithmetic and proverb interpretation was concrete. Eye movements to command and pursuit were restricted to 5° upwards and downwards but lateral movements were full. Convergence was absent. Oculocephalic responses were full. He was dysarthric and tongue movements were slow and spastic. In the limbs there was mild rigidity, hyperreflexia and flexor plantar responses. CT scan showed moderate generalised cerebral atrophy. Test results were: NART = 104, verbal IQ = 102, performance IQ = 84, index of deterioration = 11. Verbal subtest scores ranged from 12 for vocabulary to 5 for similarities. Performance subtest scores ranged from 6 for Picture...
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completion to 0 for Picture arrangement. It was noted that in the latter test he "rationalised" his incorrect responses suggesting that his poor scores on non-verbal tests were not due to visual difficulties. On the verbal version of the Recognition Memory Test he scored at chance level (29/50), and although he could not concentrate long enough to complete the visual part of the test, this also seemed to be impaired. There was no evidence of nominal dysphasia (Graded Difficulty Naming test score was average) and he performed satisfactorily on the unusual views and Fragmented Letters test for perceptual deficits. However, he was completely unable to sort Weigl shapes.

Comment: neuropsychological testing suggested mild intellectual impairment with particular involvement of non-verbal skills. Memory was weak and there was evidence of frontal lobe dysfunction, but dysphasia or impairment of perceptual skills were not detected.

Discussion

In this study we have demonstrated that intellectual impairment can be detected by neuropsychological testing in most patients with SRO syndrome at the time of diagnosis. This conclusion is in agreement with the clinical observations of Steele et al in their seminal paper. There are only a few reports of neuropsychological assessment in patients with the SRO syndrome, but most of these have shown evidence of some intellectual impairment (table 2). Impaired performance on neuropsychological tests might however conceivably result from severe dysarthria or visual difficulties.

Oculomotor abnormalities might lead to difficulties on the WAIS performance but not verbal subtests. Fisk et al found no significant difference between the mean verbal IQ of normal controls and patients with SRO syndrome, and Kimura et al reported that intellectual impairment was unequivocal only on performance subtests which required visual scanning ability. In the present study both verbal and performance IQ scores were impaired when compared to the premorbid IQ. The impairment of verbal IQ cannot be explained by visual difficulties and none of the patients was markedly dysarthric. Furthermore, performance IQ was impaired in patients with normal perceptual decoding abilities suggesting that oculomotor abnormalities were unlikely to be predominantly responsible for the impaired performance on the Picture Arrangement, block design and picture completion subtests. The failure of Fisk et al to find a significant difference between the SRO syndrome patients and controls was probably due to the small numbers studied (four in each group), as the mean verbal IQ for the SRO syndrome patients was 90.8 and 106 for the normal controls in their sample.

The main neuropathological feature of the SRO syndrome is the presence of neurofibrillary tangles and neuronal loss with gliosis in the basal ganglia, brain stem and cerebellar nuclei (1,6). The neurofibrillary tangles are often morphologically different from those in Alzheimer's disease and are not usually accompanied by senile plaques. Both diseases have, however, neuronal loss in the nucleus basalis of Meynert. Although neurofibrillary tangles have been found in the temporal and frontal cortex of patients with the SRO syndrome, cortical changes are infrequent and the disease is characterised by predominant involvement of subcortical structures. In 1974, Albert et al after studying the neurobehavioural changes in five patients with the SRO syndrome and reviewing the literature, proposed that the pattern of intellectual impairment in the SRO syndrome corresponded to a "subcortical dementia" and could be clinically distinguished from that seen in "cortical dementias" such as Alzheimer's disease; the dementia of the SRO syndrome being characterised by forgetfulness, slow mentation, emotional or personality changes, impaired ability to manipulate acquired knowledge and the conspicuous absence of dysphasia, agnosia.

Table 2 Results of neuropsychological assessment in Steele-Richardson-Olszewski syndrome

<table>
<thead>
<tr>
<th>Paper</th>
<th>Patient No</th>
<th>Verbal IQ</th>
<th>Performance IQ</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Messe et al</td>
<td>1</td>
<td>85</td>
<td></td>
<td>Severe impairment</td>
</tr>
<tr>
<td>David et al</td>
<td>1</td>
<td>100</td>
<td>82</td>
<td>Impaired</td>
</tr>
<tr>
<td>Parkes et al</td>
<td>1</td>
<td>85</td>
<td>78</td>
<td></td>
</tr>
<tr>
<td>Dix et al</td>
<td>2</td>
<td>88</td>
<td>93</td>
<td></td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>93</td>
<td>83</td>
<td>Mild dementia</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>77</td>
<td>80</td>
<td>Severe dementia</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>114</td>
<td>111</td>
<td>Mild dementia</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>107</td>
<td>97</td>
<td>Poor memory</td>
</tr>
<tr>
<td></td>
<td>5</td>
<td>107</td>
<td>97</td>
<td>Mild dementia</td>
</tr>
<tr>
<td>Kimura et al</td>
<td>7 mean</td>
<td>92.6</td>
<td>82.1</td>
<td>Impairment only on non-verbal tasks requiring visual scanning ability</td>
</tr>
<tr>
<td></td>
<td>4 mean</td>
<td>90.8</td>
<td>84.3</td>
<td>Mean verbal IQ not significantly different from normal controls</td>
</tr>
<tr>
<td>Jackson et al</td>
<td>7 range</td>
<td>68-119</td>
<td>59-115</td>
<td>Mean IQ slightly below normal values</td>
</tr>
</tbody>
</table>

*Including two patients also included by Kimura et al.
and perceptual abnormalities. Marked dysphasia has been reported in the SRO syndrome but we have confirmed that in the early stages apart from mild word finding difficulties, language functions are usually unimpaired. The deficits we observed on the digit span (which involves immediate memory and the ability to reverse digits), arithmetic (the only timed verbal subtest) and similarities (which involves ability to abstract shared characteristics of the two items) subtests are compatible with observations of Albert et al.' The impaired performance on the picture arrangement subtest in the presence of normal perception suggests that patients with SRO syndrome have a particular difficulty in arranging information in a sequential manner.

Early descriptions of the effect of frontal lobe deficits noted an impairment of "serialisation and synthesising" ability and a similarity between the neurobehavioural changes found in patients with "subcortical dementia" and bilateral frontal lobe damage has been noted. We have found that most patients who were assessed on tests believed to be sensitive to frontal lobe function performed poorly and that clinical signs of frontal lobe damage were common. In animal studies stimulation of lesions of the basal ganglia and the frontal cortex may have similar effects and patients with other diseases affecting the basal ganglia such as Parkinson's disease are impaired on tests of the frontal lobe function. In view of the increasing evidence of the importance of subcortical structures and their cortical projections in human cognitive and perceptual processes, further prospective studies of the performance of SRO syndrome patients on tests of frontal lobe function, and comparison of the pattern of neuropsychometric performance is the SRO syndrome and Alzheimer's disease will clearly be of interest.

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