Magnetic resonance spectroscopic study of parkinsonism related to boxing

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Abstract
Proton magnetic resonance spectroscopy, localised to the lentiform nucleus, was carried out in three ex-professional boxers who developed a parkinsonian syndrome, six patients with idiopathic Parkinson's disease, and six age matched controls. The three ex-boxers all showed a pronounced reduction in the absolute concentration of N-acetylaspartate compared with the patients with idiopathic Parkinson's disease and the control group. This reduction is likely to reflect neuronal loss occurring in the putamen and globus pallidus and supports the hypothesis that the extrapyramidal syndrome that may occur in ex-boxers is a distinct entity from idiopathic Parkinson's disease.

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The development of traumatic encephalopathy in boxers was first described in the medical literature when in 1928 Marland coined the term punch drunk to describe boxers who developed unsteadiness on their feet and who moved and thought more slowly.1 Since then, there have been clinical reports in which the picture of traumatic encephalopathy in boxers has been characterised by evidence of damage to one or more of the pyramidal, extrapyramidal, or cerebellar systems, with associated dementia, psychosis, and personality changes in some cases.2-5 The condition usually has an insidious onset and may present initially with affective disturbances and psychotic symptoms. The development of amnesia with the signs of parkinsonism may then follow. Finally, patients may develop generalised cognitive impairment.6 The clinical course may arrest during any of these stages and the severity of the syndrome correlates to a limited degree with the duration of the boxer's career7 and the total number of bouts.6

The development of parkinsonism is a relatively common feature in those boxers developing the punch drunk syndrome. Roberts8 noted evidence of extrapyramidal involvement in 20 of the 52 cases of traumatic encephalopathy among retired professional boxers that he found in the medical literature. There are, however, few reports in which a parkinsonian syndrome develops as the sole or predominant manifestation of boxing injuries9-9 raising the possibility that parkinsonism occurring in boxers may simply be due to coincidental idiopathic Parkinson's disease rather than from a direct consequence of head trauma.6,7

Postmortem studies in boxers with post-traumatic encephalopathy have shown distinct pathological changes,6-10 Corsellis et al,4 in a study of 15 boxers, described the presence of neurofibrillary tangles spread diffusely through both the cerebral cortex and brainstem, cavum septi pellucidi, cerebellar and cerebral scarring, and degeneration of the substantia nigra. Lewy bodies were not a pathological feature. Although no amyloid plaques were described in this paper, a follow up study of these brains by Roberts et al10 showed diffuse deposition of β protein raising the possibility that the condition is more akin to Alzheimer's disease than idiopathic Parkinson's disease. Although neuronal loss has been well documented in the substantia nigra (predominantly affecting lateral rather than medial areas) there is no information regarding the degree of cell loss in the striatum. The presence of a cavum septum pellucidum during life in these patients may be confirmed by imaging. Computed tomography has shown this abnormality in 18% of boxers compared with its presence in 5% of the general population.2

Proton magnetic resonance spectroscopy (MRS) is a non-invasive method that provides information about the chemical pathology of conditions affecting the CNS. The largest peak visible with MRS is derived principally from N-acetylaspartate (NAA), and amino acid content almost exclusively within neurons and their processes in adult brain.11 Recent work from our group has shown that MRS centres in the lentiform nucleus may be useful in differentiating idiopathic Parkinson's disease from multiple system atrophy,12 a condition with which it may be confused clinically.13,14 We now describe the clinical and MRS findings in three ex-professional boxers who developed a parkinsonian syndrome.

Patients and methods
CASE HISTORIES
Case 1
This patient was first described in the medical literature by Harvey and Newsom Davis in 1974.4 He is a 45 year old man who was noted by his family to have slurring of speech after
being involved in 16 professional bouts (see table for boxing histories). The patient attributed his symptoms to a head injury he had received while in police custody. He did not lose consciousness during this alleged assault. Six months later he started to shuffle when walking and his legs appeared stiff. During his last few bouts he was aware of slowness on his feet and difficulty avoiding trouble in the ring. After retiring, his gait deteriorated; he developed lability of mood and became aggressive. When seen in 1974, he was noted to be cooperative but mentally slow. He had a symmetric extrapyramidal disorder with an expressionless face, generalised poverty of movements, cogwheel rigidity in all four limbs, and an extrapyramidal gait. Tendon reflexes were brisk, with a left extensor plantar. Treatment with a levodopa preparation at that time resulted in some improvement in his gait.

Before his current neurological assessment he had undergone treatment with lithium for a bipolar depressive disorder. On examination he was easily distractable and poorly cooperative. He had reduced facial expression. There was no rest tremor of his limbs and he displayed reduced arm swing on walking. There was bradykinesia and cogwheel rigidity that was worse on the left side. Power was normal. Reflexes were brisk but symmetric with flexor plantar responses. There were no cerebellar signs, no sensory deficit, and sphincter function was normal.

**Case 2**

This 52 year old man fought professionally as a middle and cruiser weight (table). He was involved in a car accident when 28 and received a head injury for which he required 24 stitches. He did not lose consciousness during this event but his injuries were severe enough to prevent him continuing with his boxing career. At the age of 50 he began to drag his left leg and had trouble with his balance. He began to slow down. His speech became slurred and his memory deteriorated. There was slight improvement in his symptoms with levodopa treatment. On examination he had mild bradykinesia, with some cogwheel rigidity and pronounced blepharospasm. He had some slowness of gait with reduced arm swing. There were no cerebellar signs and no sphincter disturbance.

**Case 3**

This 69 year old man first began boxing at the age of 20 when he entered the navy. He boxed for about 18 months before turning professional. He denied being knocked out in any of his amateur or professional bouts (table). At the age of 66 he developed rest tremor of the right arm that soon spread to the other side. Levodopa produced only mild improvement. On examination he was moderately rigid and bradykinetic with bilateral rest tremor. Posture was stooped and he shuffled with short steps. Postural reflexes were impaired. There were no other neurological signs.

We also studied six patients aged 49 to 72 (median 56) years and median duration of disease of seven years with classic, levodopa responsive asymmetric idiopathic Parkinson’s disease and six healthy controls with a median age of 59 years.

The study was approved by the joint ethics committee at the Institute of Neurology and the National Hospital for Neurology and Neurosurgery, London. Informed consent was obtained before each study. Magnetic resonance imaging and MRS were performed on a 1.5 Tesla GE whole body imager with a standard quadrature head coil. The study commenced with a coronal, T2 weighted, spin echo imaging sequence (TR 2000 ms TE 80 ms) (3 mm slices with 1 mm gap, 256 × 256 matrix, echo train length 8).

Water suppressed spectra were obtained from volumes of interest (3-4 ml to 6 ml) centred on the putamen and globus pallidus.

<table>
<thead>
<tr>
<th>Case No</th>
<th>Age</th>
<th>Began boxing</th>
<th>Finished boxing</th>
<th>Amateur bouts</th>
<th>Professional bouts</th>
<th>Knocked out</th>
<th>Control medium (NAA)</th>
<th>Fenastrated septum pellucidum</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>45</td>
<td>14</td>
<td>24</td>
<td>75</td>
<td>25</td>
<td>1</td>
<td>6-1 mmol/l</td>
<td>Present</td>
</tr>
<tr>
<td>2</td>
<td>52</td>
<td>15</td>
<td>27</td>
<td>1</td>
<td>24</td>
<td>3</td>
<td>7-7 mmol/l</td>
<td>Absent</td>
</tr>
<tr>
<td>3</td>
<td>69</td>
<td>20</td>
<td>25</td>
<td>?</td>
<td>30-40</td>
<td>0</td>
<td>5-14 mmol/l</td>
<td>Absent</td>
</tr>
</tbody>
</table>

Control medium (NAA) 9-94 mmol/l; idiopathic Parkinson’s disease, median (NAA) 9-99 mmol/l.
from old pronounced Parkinson’s disease. The Lower spectrum of nucleus the left.

There was a history of predominantly Cr normal. Responsive parkinsonism, who has the concentration from taken in grey matter is (case 2)

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Results

MAGNETIC RESONANCE IMAGING

The three boxes all showed abnormalities on MRI. These consisted of discrete periventricular or subcortical white matter signal lesions. There were no high signal lesions present in the basal ganglia. In one of the three boxes (case 1), there was evidence of cavum septi pellucidi. In three of the patients with idiopathic Parkinson’s disease there were small periventricular or subcortical white matter lesions present on T2 weighted MRI. Similar changes were seen in two of the six controls. There was no evidence of cavum septi pellucidi in any of the patients with idiopathic Parkinson’s disease or controls. In two of the patients with idiopathic Parkinson’s disease there was a moderate degree of low signal extending from the globus pallidus into the medial border of the putamen. This was not present in the controls or in the three ex-boxers.

MAGNETIC RESONANCE SPECTROSCOPY

The table shows the concentrations of NAA from the lentiform nucleus in the three ex-boxers. These were all significantly lower (median 6.1 mmol/l) than the NAA concentrations from the age matched controls (median 9.94 (range 9.26–10.77) mmol/l, P < 0.03) (fig 2). By contrast, there was no significant difference in the concentration of NAA in the idiopathic Parkinson’s disease group (median 9.99 (range 9.0–11.0) mmol/l, P > 0.9) compared with the controls. There was no significant difference in the concentrations of creatine or compounds containing choline between the three groups.

Discussion

The most striking finding in this study was the significant reduction in the concentration of NAA from the putamen and globus pallidus in the ex-professional boxers with a parkinsonian syndrome, compared with the patients with idiopathic Parkinson’s disease and the control group.

In adult brain, NAA is almost exclusively confined to neurons and their processes. An absolute reduction in NAA or a reduction in the NAA/creatine ratio has been seen in many conditions in which there is neuronal or axonal loss although in some disorders this abnormality may partly reverse, suggesting that neuronal dysfunction alone may contribute to a reduced ratio.

There is, to our knowledge, no information in the medical literature describing the corpus striatum in patients with post-traumatic encephalopathy and, in particular, whether there is neuronal loss in this region of the brain. The reduction of NAA that we have shown in this small group of patients with parkinsonism related to a history of professional boxing suggests that neuronal loss may occur in the putamen, globus pallidus, or both in these patients, presumably secondary to head trauma. This hypothesis is supported by the results of an earlier study in which we

\[
\text{Met} = [\text{H}_2\text{O}] \times \text{PI} \times \text{T}_1 \text{cor} \times \text{T}_2 \text{cor} \times \frac{S_{\text{NAA}}}{S_{\text{H}_2\text{O}}} \times 1/2^R
\]

where \( S_{\text{NAA}} \) and \( S_{\text{H}_2\text{O}} \) denote the signal intensities for metabolites and water respectively, \([\text{H}_2\text{O}]\) is the brain water concentration in grey matter taken as 47-2 M, \( \text{T}_1 \text{cor} \) and \( \text{T}_2 \text{cor} \) are \( \text{T}_1 \) and \( \text{T}_2 \) correction values based on published values for the metabolites studied, \( \text{PI} \) is the proton index, and \( R \) (attenuation value) = ((R1 + R2) metabolite − (R1 + R2) water). Statistical analysis was performed with a Mann-Whitney confidence interval and test. Results are expressed as a median value together with the range and P value.

Figure 2 Top: spectrum taken from the right lentiform nucleus of a 52 year old ex-boxer (case 2) who has a four year history of parkinsonism, worse on the left. There is a pronounced reduction in the concentration of NAA. Lower: spectrum taken from the left lentiform nucleus of a 49 year old man with an eight year history of predominantly right sided classic levodopa responsive idiopathic Parkinson’s disease. The concentration of NAA is normal. Cr = creatine; Cho = compounds containing choline.
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found a reduction of NAA from the lentiform nucleus in patients with the striatognal and olivopontocerebellar variants of multiple system atrophy, even in a condition in which neuronal loss is known to occur in the putamen, and to a lesser extent, in the globus pallidus.\(^4\) Neuronal loss within the striatum is not a feature of idiopathic Parkinson’s disease\(^27\) and this in keeping with the finding in the present study of a normal median concentration of NAA in the patients with idiopathic Parkinson’s disease compared with the controls. The reduced concentration of NAA in the post-traumatic patients suggests that the extrapyramidal syndrome which may occur in ex-boxers is an entity distinct from idiopathic Parkinson’s disease.

Prospective studies in patients diagnosed as having idiopathic Parkinson’s disease have shown that up to 24% of patients have an alternative diagnosis at postmortem.\(^14\) The conditions most often misdiagnosed include multiple system atrophy, Steele-Richardson syndrome, and vascular disease, although the pathological changes of post-traumatic encephalopathy have been found at postmortem in the brains of a few patients believed during life to have idiopathic Parkinson’s disease (Parkinson’s Disease Society Brain Bank, Institute of Neurology, unpublished observations). This finding suggests that trauma may play a part in the development of a parkinsonian syndrome in at least some patients diagnosed as having idiopathic Parkinson’s disease. Our finding of a reduction of NAA in the lentiform nucleus in these ex-boxers supports this hypothesis.

Two of the boxers in this study (cases 2 and 3) developed parkinsonism several years after the end of their boxing career. We cannot exclude the possibility that these patients have developed coincidental idiopathic Parkinson’s disease unrelated to their history of head trauma. There are two reasons, however, why this seems unlikely. Firstly, these patients have only ever shown a mild or sustained therapeutic response to levodopa which is very unusual in idiopathic Parkinson’s disease. Secondly, both have shown a reduction of NAA in the basal ganglia with MRS, a finding not typical of idiopathic Parkinson’s disease. How then, could one explain the late development of parkinsonism in these two patients? If head injury from boxing does produce neuronal loss within the striatum, then the degree of cell loss may be an important factor in determining when symptoms develop. If pronounced, parkinsonian features may develop at, or shortly after, the time of injury, as perhaps in case 1. With less severe cell loss, however, the threshold required to produce symptoms may not be reached. With aging, there may be further loss of neurons from the basal ganglia leading to the development of an extrapyramidal syndrome.

In conclusion, proton MRS centred on the lentiform nucleus has shown a reduction of NAA (a neuronal marker) in three ex-professional boxers who developed a parkinsonian syndrome. By contrast, a group of patients with idiopathic Parkinson’s disease showed preserved concentrations of NAA compared with controls. This finding supports the hypothesis that the parkinsonian syndrome occurring in boxers may be a separate clinical entity from idiopathic Parkinson’s disease.

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