SHORT REPORT

Late adult onset chorea with typical pathology of Hallervorden-Spatz syndrome

D A Grimes, A E Lang, C Bergeron

Abstract

Senile chorea is a well recognised but poorly understood clinical entity characterised by a slowly progressive, generalised chorea in elderly people without mental deterioration or a clear underlying cause. The Hallervorden-Spatz syndrome is typically thought of as a paediatric condition with extrapyramidal features and dementia. However, it has been described in adults usually presenting with parkinsonism plus dementia. An elderly woman with slowly progressive chorea without dementia was found at postmortem to have the pathological features originally described by Hallervorden and Spatz. This association has not previously been reported. (J Neurol Neurosurg Psychiatry 2000;69:392–395)

Keywords: Hallervorden-Spatz syndrome; neurodegeneration with brain iron accumulation type 1; senile chorea

The diagnosis of Hallervorden-Spatz syndrome encompasses a considerable range of disorders, all with the core pathological features of iron deposition, axonal spheroids, and gliosis of the pallidum and substantia nigra. It has recently been reclassified as “neurodegeneration with brain iron accumulation type 1 (NBIA-1)” however, most authors continue to use the original terminology. Most cases with this pathology begin in childhood or early adolescence with extrapyramidal dysfunction and dementia, and follow a relentlessly progressive course. Adult onset cases have been described but typically have prominent dementia associated with parkinsonism. Chorea, although commonly described in paediatric cases, is not found in isolation and has not been reported in adults with the pathological features originally described by Hallervorden and Spatz. We report on a patient with the clinical diagnosis of “senile” or late adult onset chorea who was found to have this pathological diagnosis on postmortem examination.

Case report

A 76 year old woman was referred with a 1 year history of unsteady gait and unusual movements of her head and upper limbs that had been noticed by her family. Her medical history included myocardial infarction 20 years previously with mild congestive heart failure, peripheral vascular disease, and arthritis. Her medications were digoxin, furosemide, and ibuprofen. There was no known neuroleptic or toxic exposure, no history of Sydenham’s chorea or rheumatic fever, and no family history of abnormal movements.

Her initial mental status examination showed an alert, fully oriented woman with only mild difficulty in tasks of attention. Her motor examination was normal except for the mild generalised choreiform movements. The rest of her neurological examination, including extraocular movements, was normal except for slight unsteadiness when walking and mild difficulty with tandem gait.

Laboratory investigations including complete blood count, blood smear, thyroid studies, liver function tests, erythrocyte sedimentation rate, ceruloplasmin, antinuclear antibody, and rheumatoid factor were normal. Huntington’s disease molecular genetic testing for the CAG trinucleotide repeat expansion was negative (number of repeats=23 and 21). Neuropsychological testing disclosed only mild decline in cognitive functioning and was considered not to be consistent with any significant degree of dementia. Head MRI, done 2 years after the onset of her chorea, demonstrated decreased signal intensity in the putamen, caudate, substantia nigra, and dentate nuclei bilaterally (fig 1). There were also multiple small non-specific focal white matter lesions seen in the centrum semiovale and periventricular regions bilaterally. She was diagnosed as having “senile chorea” with the MRI findings thought to represent iron deposition in the basal ganglia of unknown cause.

Her chorea slowly increased in severity and 5 years after her first assessment reserpine (2 mg/day) was initiated with moderate improvement in her chorea. Two years later tetrabenazine (75 mg/day) was substituted with further benefit because of concerns that the reserpine may have been exacerbating her increasing postural instability. She required placement in a nursing home 8 years after the onset of her chorea because of the worsening postural stability and her increasing number of falls. She was last examined at this time and had moderate head rocking movements with mild
generalised chorea that increased with movement. She continued to have only minor memory difficulties. She died suddenly at the age of 85, 9 years after the onset of her chorea.

NEUROPATHOLOGICAL FINDINGS

Neuropathological examination was confined to the right half of the brain. External examination of the cerebral convexity showed no significant cortical atrophy but there were two small areas of subarachnoid haemorrhage in the right occipital and superior parietal areas. Coronal sections through the cerebral hemisphere showed a slight flattening of the head of the caudate nucleus. The medial segment of the globus pallidum had a cribiform appearance. The entorhinal cortex was thinner than normal and there was mild atrophy of the amygdala. A small area of cortical infarction was seen underlying the area of subarachnoid haemorrhage in the right superior parietal area. A large haemorrhage measuring 3.5×4.0×8.0 cm was noted in the occipital lobe extending into the posterior temporal area. Horizontal sections through the brainstem were unremarkable with the substantia nigra and locus ceruleus well pigmented. Sagittal sections through the cerebellum were unremarkable, including the dentate nucleus. Microscopic examination showed severe neuronal loss in the globus pallidus, predominantly in the medial segment, with abundant neuroaxonal spheroids and a moderate degree of iron deposition.

Figure 1 T2 weighted MRI.

Figure 2 (A) Neuronal loss, (B) spheroids (arrows), and (C) iron pigment deposition (arrow) in the medial segment of the globus pallidus. (D) Spheroids in the substantia nigra, pars reticulata (arrows). Haematoxylin and eosin luxol fast blue (A, B, and D). Perl’s Prussian blue reaction for iron (C).
Postmortem cases of late adult onset (age>50 y) Hallervorden–Spatz syndrome

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<tr>
<td>Dooling* et al</td>
<td>—</td>
<td>75</td>
<td>Not thought to have primary neurological disorder</td>
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*Limited details given and the patient’s specific pathology not described but was part of large postmortem series of Hallervorden–Spatz syndrome.

the disorder consist of (1) neuronal loss and gliosis affecting mainly the internal segment of the globus pallidus and the pars reticulata of the substantia nigra; (2) widely disseminated rounded or oval non-nuclear "spheroids", especially in the globus pallidus and substantia nigra; (3) intracellular and extracellular iron deposition in the above areas.1 2

The pathological findings in our patient are characteristic of Hallervorden-Spatz syndrome including neuronal loss and gliosis, axonal spheroids, and iron deposition in a typical distribution. Iron deposition can be seen in normal aged brains and has been described in other neurological entities, including heredo-degenerative and storage diseases.3 Axonal spheroids are also found in many conditions and are thought to be the mark of neuroaxonal dystrophies. However, only Hallervorden-Spatz syndrome is thought to have the unique combination of pallidionigral gliosis, iron deposition, and axonal swellings.

Since Hallervorden and Spatz’s original description an increasing number of cases have been reported with “atypical” clinical features such as a cerebellar ataxia, retinitis pigmentosa, and/or a seizure disorder.1 Several cases have also been described in adults over the age of 50 (table), often presenting with a progressive dementia or parkinsonism. Neurofibrillary tangles4 and Lewy bodies8 2 have also been described in association with the pathology of Hallervorden-Spatz syndrome. This range of clinical features and associated pathologies has encouraged the use of the term Hallervorden-Spatz “syndrome” instead of “disease”. However, it has been proposed that the eponymic designation be dropped completely because of the apparent non-specificity of the pathology and for humanitarian reasons related to their involvement with human experimentation in Nazi Germany.9

Senile chorea is an uncommon, apparently sporadic clinical entity, characterised by the presence of a late onset, generalised chorea without dementia. The question of whether it represents a single disease or syndrome had been debated for many years. Some authors had thought that it was due only to late onset Huntington’s disease, with death presumably occurring before the mental changes.10 As the availability of the trinucleotide CAG genetic testing it has become clear that Huntington’s disease may account for up to 50% of cases, but there is still a subgroup of patients for which no cause is identified despite extensive investigations.11-13

The presentation in our patient of isolated chorea is very unusual and has not been associated with Hallervorden-Spatz syndrome pathology. Her premorbid diagnosis was that of “senile chorea”, as she had no identifiable cause for her abnormal movements. Huntington’s disease was ruled out by her negative CAG genetic testing. Extensive laboratory testing found no abnormalities to suggest an alternative diagnosis. Her MRI did not show the typical “eye of the tiger” sign or pronounced low signal in the globus pallidus without the additional inserted bright signal that has now

NEUROPATHOLOGICAL DIAGNOSIS

The neuropathological diagnoses were: Hallervorden-Spatz syndrome; old occipital infarction; recent intracerebral haemorrhage due to amyloid angiopathy; mild Alzheimer’s changes.

Discussion

In 1922, Hallervorden and Spatz described the syndrome and pathological findings that characterise the condition that bears their names.4 It classically presents as a relentlessly progressive disease of childhood and adolescence in which motor symptoms predominate with prominent rigidity, dystonia, intellectual impairment, and/or pyramidal tract signs.5 Choreoathetotic movements or tremor may also be present but these features have not been described in isolation. This “typical” presentation often follows an autosomal recessive pattern of inheritance and has recently been linked to chromosome 20p12.3-p13 in 10 families.7 Pathologically, the main features of

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been well documented in patients with Hallervorden-Spatz syndrome. In fact the abnormal low signal, designating excessive iron deposition, was restricted to the striatum despite the greater extent of pallidal involvement found at necropsy. This discrepancy may in part be due to the fact that the MRI was done early in the course of her disease.

It is difficult to explain the wide variability of clinical features that have been associated with the pathology of Hallervorden-Spatz syndrome. It is highly unlikely that our patient’s findings arose from the same pathophysiological mechanisms as typical paediatric cases. Hallervorden-Spatz syndrome is an umbrella term for a heterogeneous group of neurodegenerative disorders which will become antiquated as further research improves our understanding and classification.