Motor neuron disease (amyotrophic lateral sclerosis) arising from longstanding primary lateral sclerosis

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Abstract
Three men were initially diagnosed as having primary lateral sclerosis (PLS), but eventually developed amyotrophic lateral sclerosis (ALS) after 7-5, 9, and at least 27 years. Non-familial ALS and PLS might be different manifestations of a single disease or constitute completely distinct entities. The clinical diagnosis of PLS predicts a median survival that is four to five times longer than in ALS.

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Primary lateral sclerosis (PLS), defined as non-familial progressive spinobulbar or bulbospinal spasticity—without amyotrophy, fasciculation, optic atrophy, deafness, or pedes cavi—and with no more pronounced sphincter disturbances than urgency of micturition, and not caused by a segmental lesion, is rare. Its status as a nosological entity, separate from the hereditary spastic paraplegias on the one hand and amyotrophic lateral sclerosis (ALS) on the other, remains disputed. Mulder suggested that ALS begins with peripheral weakness, and Pringle and coworkers diagnose PLS on initial central motor deficit.

Gowers described the first patients with progressive spastic paraparesis, eventually complicated by amyotrophy. Spiller reported eight patients of whom bulbar or limb spasticity was the initial sign. This was followed by lower motor signs in six, but remained as the only sign in two. Wilson and Brouwer claimed to have seen similar patients with a long standing spastic paraparesis eventually followed by wasting in the hands.

Our three patients (one with a necropsy) demonstrate that longstanding PLS may change into ALS.

Case reports
Case 1
In 1984, at the age of 45, a previously healthy manager experienced trouble in maintaining his balance, and noticed slowly progressive stiffness of the left leg with cramps. The family history was negative and consanguinity was excluded. Examination disclosed slight weakness of the left thigh muscles, mild spasticity of the legs, knee and ankle cloni, and extensor plantars. Blood chemistry, CSF, and EMG were unremarkable. Magnetic resonance imaging of the cervical spine was normal and PLS was diagnosed. Micturition urgency began in 1987. Examination in 1988 showed definite spastic paraparesis and mild proximal weakness of the legs. Sural and tibial nerve somatosensory evoked potentials (SSEPs) were bilaterally absent and delayed respectively. Visual and brainstem auditory evoked potentials (VEPs, BAEPs), and brain MRIs were normal. By 1990, walking had become troublesome and dysarthria was present; the calves showed fasciculation and some atrophy. The left extensor hallucis muscle was paralytic and the right corneomandibular reflex was now positive. During 1991 and 1992 dysarthria, forced laughter, and generalised weakness progressed and amyotrophy became evident. Examination now disclosed widespread fasciculation of trunk and limb muscles, considerable amyotrophy, most pronounced in the lower legs, and brisk masseter and bilaterally positive corneomandibular reflexes. The diagnosis was changed to that of ALS. Repeat EMG in 1993 showed very low compound muscle action potentials (CMAPs) of the intrinsic foot muscles bilaterally. Denervation activity seemed limited to the anterior tibial muscles, whereas re-innervation activity was seen in the limb musculature. The patient continues to deteriorate slowly.

Case 2
In 1983, at the age of 39, a previously healthy plumber noticed slurring of speech. The family history was negative; no consanguinity was present. Examination of this otherwise healthy right handed man of athletic build disclosed minimal dysarthria, increased tendon jerks, and positive corneomandibular reflexes. Physical examination and extensive study of blood chemistry and CSF were unremarkable. The diagnosis was changed to that of ALS. Repeat EMG in 1993 showed very low compound muscle action potentials (CMAPs) of the intrinsic foot muscles bilaterally. Denervation activity seemed limited to the anterior tibial muscles, whereas re-innervation activity was seen in the limb musculature. The patient continues to deteriorate slowly.
were bilaterally absent; median nerve SSEPs and EMG were normal and PLS was diagnosed. In 1989, atrophy of the interosseous hand muscles became apparent, with fasciculation of wrist and finger extensors. Sensation remained intact. Electromyography showed signs of re-innervation but no evidence of denervation. Conduction velocities were normal. Magnetic resonance imaging showed some thinning of the cervical cord but no brain abnormalities. Amyotrophy spread to involve the forearms and eventually the lower legs. Repeat EMG in 1991 disclosed fibrillation and positive sharp waves in the left anterior tibial muscle; in 1993 very low CMAPs of the extensor digitorum communis muscles were noted, as well as denervation activity in both tibial anterior muscles, the rectus femoris, and the first interosseous muscle, with fasciculations. Re-innervation activity was seen in the distal and proximal limb muscles.

The diagnosis was changed from PLS to ALS. The course remains slowly progressive.

CASE 3
In 1957, a 33 year old office clerk was admitted for analysis of a heavy feeling in his legs and cramps in the calves and proximal leg muscles, progressive complaints that had begun at the age of 27. There was no consanguinity and the family history was negative. On admission, hyperreflexia of arms and legs was found with a patellar clonus, and a left extensor plantar sign; CSF was normal and PLS was diagnosed. His condition deteriorated very slowly. Examination in 1978 showed moderate spasticity of both legs, knee and ankle clonus, Babinski signs, and a slightly decreased vibration sense at the ankles. Fasciculations or atrophy were not present. Cervical myelography was normal. Electromyography showed decreased conduction velocities of both peroneal nerves with no denervation activity. The patient was lost to follow up but was necropsied in 1989, having died of pneumonia.

There were no gross alterations to the brain. The spinal cord and the anterior roots showed mild atrophy. Microscopy showed a severe loss of spinal anterior horn cells, a subtotal at the dorso lumbar and mild at the cervicodorsal level. Mild cell loss was noted in Clarke’s columns. Numerous amylaceous bodies and occasional instances of neuronophagia with considerable reactive gliosis were evident throughout the cord. A pronounced myelin pallor characterised the pyramidal tracts. Macrophages and activated microglial cells immunocytochemically stained with Tal 1 B5 (MHC common framework antigen) antibody (Bodmer; dilution 1:20) on paraffin slides (for method see van den Bergh et al1') were strongly positive in the entire spinal cord (figs 1 and 2), except the posterior funiculi. Lymphocytic infiltration was not present. The diagnosis was changed to ALS.

Discussion
The El Escorial diagnostic criteria4 for ALS include both upper (spasticity, hyperreflexia, extensor plantar signs, increased gag and snout reflexes, and pseudobulbar effect) and lower (asymmetric weakness, atrophy, fasciculation) motor neuron signs. The onset is insidious, its course invariably progressive, usually without sensory involvement (although sensory pathways may be affected19), and survival is inversely related to age at diagnosis.11 12 Clinical diagnostic criteria for PLS include an insidious adult onset of spastic paresis, usually beginning in the legs, without a family history, running a slowly progressive course of at least three years, ultimately leading to a severe spastic spinobulbar paresis, and atrophy of the precentral gyrus on MRI.

The debate—whether PLS is a distinct nosological entity or a forme fruste of ALS—
is still going on. According to Mackay, PLS is simply ALS without lower motor neuron signs, which are bound to appear unless death supervenes. In his series of 70 deceased patients with ALS 11 presented with purely spastic features for several years before muscular atrophy became manifest. Only one patient remained purely spastic until death. He excluded three patients, alive at the time of study, who had had purely spastic paresis for as long as 21 years. The study did not mention EMG. In a thorough clinicopathological study, four patients clinically had PLS. In three of these patients, the time from onset of symptoms until death was short (16–30 months). A fourth patient (case 43), with a spastic spinobulbar paresis, died eight years after onset. Necropsy showed loss of motor cells and of anterior root fibres, but also a multiple myeloma with lesions in several vertebrae, although compression or infiltration of the cord had been excluded.

Younger et al reported three necropsied cases of PLS, with a disease duration of 1, 5–5, and 10 years. These patients showed symmetric demyelination of the corticospinal tracts at all spinal levels, without involvement of anterior or dorsal columns, without gliosis or Betz cell loss in the precentral gyrus, and without decrease of motor neurons in brainstem nuclei or the spinal cord. Fisher reported one necropsied case of chronic bilateral spinobulbar spasticity with a five year survival, finding demyelination of medullary pyramids and lateral corticospinal tracts at all spinal levels, and probably a reduced number of Betz cells in the motor cortex. A second case of pure spastic paresis, with a disease duration of less than two years, showed selective demyelination of the lateral corticospinal tracts without abnormalities of the brain and brainstem. Beal and Richardson described a necropsy of a woman with a 3–5 year history of PLS. This showed a severe loss of Betz cells in the precentral gyrus, atrophic medullary pyramids, a paramedian pontine infarct, and demyelination of anterior and lateral corticospinal tracts at all spinal levels.

Correct diagnosis in patients with a chronic progressive spastic paraparesis, will remain a diagnostic challenge. To avoid unnecessary distress to patients, retention of the diagnosis of PLS may be justified, because of its favourable prognosis compared with ALS with lower motor neuron onset.

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