Focal cervical poliopathy causing juvenile muscular atrophy of distal upper extremity: a pathological study

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SUMMARY A new clinical entity under the name of “juvenile muscular atrophy of unilateral upper extremity” was first described in 1959. Although about 150 cases in Japan, and several additional cases in other countries, have been clinically reported in the literature, the pathology has remained unknown because of the benign course of the disease. The first necropsy findings are reported, obtained from a patient with this disease, who died of lung cancer at the age of 38, 23 years after the onset of the disease. The lesions existed only in the anterior horns of the spinal cord at C5 ~ T1, particularly marked at C7 and C8, showing shrinkage and necrosis, degeneration of various degrees of large and small nerve cells, and mild gliosis. The pathological findings differ from those of reported cases of spinal vascular disorders, but some circulatory insufficiency in the territory of the spinal cord would seem to be suggested, although the underlying aetiology remains unknown.

In 1959, one of the authors (KH)1 reported 12 cases of “juvenile muscular atrophy of unilateral upper extremity,” clinically differentiated from the known types of motor neuron disease. Our accumulated cases were 20 by 19632 and 38 by 19723 allowing further definition of its clinical features. About 150 clinical cases have since been reported in Japan, including many cases reported by Sobue.4 Cases have also been reported from Denmark,5 Holland,6 Singapore,7 and India.8,9 Very recently, similar cases have been reported from Malaysia10 in a letter to this journal.

Clinical features include the following: (1) It occurs predominantly in males of 15–25 years old. (2) Is usually sporadic, though rarely familial. (3) Manifests itself insidiously with muscular weakness and atrophy in the hand without any inducing factors such as infections and trauma, but its progression ceases in 2–3 years, mostly within 1–2 years. (4) The muscular atrophy is limited to the hand and forearm, excluding the brachioradialis muscle, thus showing a characteristic oblique amyotrophy over the forearm. (5) It shows cold paresis of the fingers (weakness exaggerated by cold) and a fine and irregular tremor upon finger extension. (6) It predominates on one side (unilaterally in most cases, asymmetrical in some, and symmetrical in a few). (7) Muscle biopsy and needle electromyography show denervation, and may disclose subclinical changes even on the non-atrophic side. Nerve conduction velocity and cerebrospinal fluid are normal. (8) Except for a very rare and mild sensory abnormality over the dorsum of a hand, it lacks sensory disturbances, Horner’s syndrome, abnormal tendon reflexes, pyramidal tract signs, and urinary disturbances. (9) Computed tomography with intrathecal metrizamide demonstrates a partial atrophy of the lower cervical cord, but routine radiography fails to show cervical spondylosis or disc herniation.

A quarter of a century has elapsed without pathological findings, due to the benign course of the disease. Recently we first obtained necropsy findings from a patient who died of lung cancer 23 years after the onset of this disease.

Case report

The patient was a 38 year old right handed male. The past medical history is non-contributory, without trauma. The family history includes a brother, 5 years younger, with muscular atrophy in the right hand which developed at the age of 13 years but ceased progression in about 6 months.

History of present illness In 1959, at the age of 15 years, the

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Patient noted weakness in his left hand upon braking a motor cycle. His hand grip became weaker thereafter, with atrophy in the hypothenar eminence and the first dorsal interosseus muscle, which stopped progressing in about a year. In 1961, at the age of 17 years, he was examined in another institute, but no clear diagnosis was made other than muscular atrophy. He also noted a fine tremor and cold paresis in his left hand. His condition had remained unchanged and untreated.

In May, 1980, at the age of 35, he noted a mild atrophy in his right first dorsal interosseus muscle and fatigability of the right hand on writing. In August, 1980, he was admitted to the Department of Neurology (Prof Hirayama) in Chiba University.

**Neurological findings on admission** The left forearm and small hand muscles, except for the brachioradialis, showed the characteristically localised atrophy and weakness. The right small hand muscles also showed mild atrophy (fig 1). These were associated with a mild hypotonia at the wrists and hands. The fingers showed asynchronous, irregular, and fine tremors upon extension. The deep tendon reflexes were all symmetrically normal, without Babinski's sign. Ataxia, extrapyramidal signs, sensory disturbances, Horner's sign, and abnormalities in sweating and urination were all absent.

**Laboratory data** The plain cervical spine radiographs showed no abnormalities. The cerebrospinal fluid was normal. Queckenstedt's test was negative. Needle electromyography showed neurogenic changes in the atrophic muscles, more marked on the left. Peripheral motor nerve conduction velocities were normal in the four extremities. A biopsy specimen of the left flexor carpi ulnaris muscle showed neurogenic changes. Myelography disclosed a slightly decreased anteroposterior diameter of the lower cervical cord.

**Progress note** The signs and symptoms had remained unchanged from August, 1980 until September, 1982 at the age of 38, when he developed a dry cough. He was admitted to the Department of Medicine, Matsudo City Hospital, with the diagnosis of a mediastinal tumour. On October 6 he developed neurological symptoms due to cerebral metastases shown on a CT scan. Chemotherapy was started on October 29. His cerebral symptoms progressed further in November and he expired on December 6. In the meantime, $^{99m}$Tc bone scanning of the whole body had demonstrated abnormal uptake areas in the left 12th rib, the 5th lumbar vertebra, and the right hip joint. Lymph node biopsy through a mediastinoscope suggested a squamous cell carcinoma.

**Neuropathological findings**
The necropsy was performed 4 hours after death. General pathology revealed that the left upper pulmonary lobe contained a primary adenocarcinoma. The aorta showed no atheroma except at the bifurcation nor thickening of the wall.
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Discussion

From his clinical case studies of this disease, Hirayama inferred pathological changes at C6, 7, 8, and T1. The present necropsy study demonstrated...
lesions at C5–T1, milder at C5 and C6, giving a good clinicopathological correlation.

Some authors, on clinical grounds, consider this disease as a form of motor neuron disease, as we ourselves once did. However, the present pathological findings do not fall into the category of motor neuron degenerative disease. Although basophilic inclusion bodies are occasionally found in juvenile motor neuron diseases, they were found only at the upper (C5) and the lower extremes of the lesions (T1) in the present case.

The peculiar case of Garcin and Gruner had features both of motor neuron degenerative disease and of a vascular disorder. The muscle atrophy began in the lower extremities and ascended progressively. The most marked lesions were seen in the lumbar anterior horns, with central cavitation surrounded by astrocytes and proliferated vessels, with some remaining intact neurons. Fibrous gliosis involved not only the anterior horns but also their surrounding white matter.

In cases with long survival of an attack of acute anterior poliomyelitis, such as those described by Peers, one expects to see destruction of gray matter proportional to the loss of neurons and dense fibrous gliosis. In contrast, the usual clinical manifestations...
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and the constancy of the levels affected in this disease, along with the relatively mild gliosis seen in our present case, bear no resemblance to the patterns seen in cases of previous poliomyelitis.

A tumour, a syrinx, or even a traumatic lesion in the lower cervical cord can cause muscular atrophy with a similar distribution to that seen in the present disease. However, the present necropsy study did not show such lesions. As for trauma, not only major cervical spinal injuries but also such minor injuries sustained in stretching the arms or dressing may induce cord lesions. In such cases, tetraparesis will rapidly develop, and the lesions are most marked at C6, involving not only the anterior horns but also the adjacent white matter, including the anterior funiculi, parts of the lateral funiculi, and the anterior portion of the posterior funiculi. Repeated minor traumas can also cause cord injuries which are, however, difficult to substantiate. Some have reported delayed onset of neurological manifestations after trauma, but their clinical and pathological findings are entirely different from ours. A further point of difference from cases of delayed traumatic myelopathy is that some of our cases show a familial tendency to the disease.

Among the vascular lesions of the spinal cord, this disease most resembles the “téphromalacie anté-riure” of Marie and Foix with infarcts confined to the anterior horns at C7–T1. However, the latter occurs among elderly persons, and its pathology consists of infarction, resulting from occlusion of the spinal arteries due to syphilitic arteritis or atherosclerosis.

Spinal vascular lesions, segmentally localised at the lower cervical levels, were reported in four out of 21 cases of arteriosclerotic myelopathy in the aged by Gruner et al. In their transverse sections of the cord, the necrotic changes involved not only the anterior horns but also the anterior portion of the posterior funiculi. After a study of spinal cords of the aged, Larese stated that the anterior horn, particularly the neurons in its central portion, was the most sensitive to spinal circulatory disorders. Gilles et al. observed in children that a transient cardiac arrest or arterial hypotension damaged the anterior more than the posterior horns, and the central more than the peripheral portions of the anterior horn. Our case seems to have pathological changes similar to those reported in these cases. However, in these two and other reports, the lesions secondary to systemic circulatory failure are characteristically confined to the lower (lumbar) spinal cord, unlike those in our case.

Necropsy studies in cervical spondylosis by Brain et al. and in disc herniation by Mair et al. showed lesions at mid-cervical levels, involving the anterior portion of the posterior funiculi, the anterior horns, and the adjacent intermediate zone of the lateral funiculi, all lined on the transverse axis of the spinal cord. Breig et al. postulated that forward bending of the neck would flatten the cord from front to back, stretching the vessels sideways; this in turn could induce a secondary vascular disorder. A role of the dentate ligaments in inducing cord lesions was emphasised in cervical spondylosis by Cusick et al. and in multiple sclerosis by Oppenheimer. Yada et al. recently proposed as a cause of cervical myelopathy an over-stretch mechanism associated with compression of the dura and cord against the vertebrae during neck flexion. These hypotheses are interesting when considering the pathogenesis of the present disease, but these lesions, different from our case, involve the white matter, producing long tract signs.

The pathological findings in the present case certainly differ not only from those of other disorders but also from those of reported cases of spinal vascular disorders; but they still seem to suggest a circulatory insufficiency in the lower cervical cord as the most likely pathogenesis. As long as the underlying aetiology remains unknown this disease may be referred to clinico-pathologically as “juvenile focal cervical poliopathy.”

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