Lhermitte’s sign in subacute combined degeneration of the cord

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SUMMARY Lhermitte’s sign is a common early symptom of subacute combined degeneration of the cord occurring in 11 out of 44 patients admitted to the National Hospitals for Nervous Diseases during the decade 1962–71 with this diagnosis. Two patients, in both of whom it was the presenting complaint, are described in detail. It is concluded that, in these cases, Lhermitte’s sign is due to stretching of demyelinated fibres in the posterior columns in the cervical cord, produced by neck flexion. The symptoms disappear after treatment with vitamin B₁₂. The clinical importance of this symptom is emphasized.

It was in 1917 that Marie and Chatelin first described the syndrome that has come to be known as Lhermitte’s sign, in the following words:

‘... le blessé a la sensation qu’un courant électrique parcourt ses bras et même ses jambes, il en est de même lorsque le blessé baisse fortement la tête sur la poitrine, se baisse ou s’accroupit . . .’

These authors thought that the symptomatology, which in their case was due to trauma, was secondary to a radiculopathy, but Babinski and Dubois (1918) considered that it was caused by a cord lesion. Lhermitte (1918) was of the same opinion and later (Lhermitte, Bollak, and Nicolas, 1924) described it in detail in a case of multiple sclerosis. Since that time, Lhermitte’s sign has been described in a variety of other conditions including cervical cord tumours and tuberculosis of the cervical spine (Patrick, 1930), cervical spondylisis (Alajouanine, Thurel, and Papaïoanou, 1949), arachnoiditis (McAlpine, Lumsden, and Acheson, 1965), and radiation myelopathy (Jones, 1964).

The occurrence of Lhermitte’s sign in subacute combined degeneration of the cord was first described by Hassin (1933), but, since then, there have been very few other reports in the literature. Two cases seen recently suggested that the association might not be so unusual and this view was borne out when it was found that 11 out of the 44 patients admitted to the National Hospitals for Nervous Diseases with subacute combined degeneration of the cord between 1962 and 1971, complained of this symptom at some time in the course of the illness.

CASE 1

The patient, a housewife aged 48 years, was seen for the first time in July 1969. For two months, she had noticed momentary, shooting, tingling sensations down the arms and legs when flexing her neck. Two weeks later she complained of paraesthesiae in the hands and feet, followed by clumsiness in the use of her fingers.

Physical examination was negative apart from mild impairment of position and vibration sense in the feet and the ankle jerks were depressed.

Investigations revealed: haemoglobin 12.4 g/100 ml., MCV 109 cu.μ, macrocytes present on the blood film, serum B₁₂ 80 μμg/ml., Schilling test part I 2.5%, part II 10.5%. Test meal showed histamine fast achlorhydria; radiographs of the cervical spine showed changes of spondylisis; the narrowest diameter of the cervical canal was 15.5 mm. Nerve conduction studies were normal apart from an absent sensory action potential in the left lateral popliteal nerve. Barium meal was normal.

The patient was treated with daily injections of 1,000 mcg vitamin B₁₂ for five days, reducing to maintenance dosage thereafter. Her symptoms, including Lhermitte’s sign, disappeared completely within four months and she was in perfect health when last seen in September 1972.
CASE 2

The patient, a male schoolteacher aged 35 years, was seen for the first time in February 1971. For four months, he had noticed electric-shock-like feelings which ran down the backs of his legs when he bent his neck forwards. The sensation was momentary and invariably precipitated by neck flexion. For two months he had been troubled by paraesthesiae from the costal margin downwards and for three weeks by some clumsiness of the hands.

Examination revealed a smooth tongue and widespread vitiligo. There was impairment of joint position sense in the fingers and toes and vibration sense and cutaneous sensibility were defective below the costal margin. The tendon reflexes in the legs were absent.

Investigations revealed: haemoglobin 13.3 g/100 ml., MCV 104 cu.μ; no macrocytes were seen on the blood film, the bone barrow was megaloblastic, serum B₁₂ level was 75 μg/ml., Schilling test part I 0-5%, part II 17-8%. A test meal showed histamine fast achlorhydria. Radiographs of the cervical spine were normal; the narrowest diameter of the cervical canal was 17.0 mm. Nerve conduction studies were normal apart from an absent sensory action potential in the right lateral popliteal nerve.

This patient was also treated with daily injections of 1,000 mcg vitamin B₁₂ for five days, reducing to maintenance dosage thereafter. His symptoms, including Lhermitte’s sign, disappeared within three months and he was perfectly well when last seen in September 1972.

DISCUSSION

In the decade 1962–71, 44 patients with proven subacute combined degeneration of the cord were admitted to the National Hospitals for Nervous Diseases. In 11 of them (25%), including the two described in this report, Lhermitte’s sign was a prominent complaint.

The symptomatology was remarkably similar in all of them and identical with that described in the early papers by Lhermitte (Lhermitte et al., 1924; Lhermitte, Lévy, and Nicolas, 1927). Momentary electric-shock-like feelings were produced by flexion of the neck. The distribution was variable; most commonly the sensations radiated down the back into the legs, but in one patient they were felt in the arms only and in another just down the back. Although most of these patients had signs of damage to pyramidal tracts and posterior columns, as well as those of a peripheral neuropathy, the one sign common to all of them was loss of position and vibration sense in a distribution suggesting a lesion of the posterior columns in the cervical cord. This finding strongly supports the generally accepted view that damage at this site is a prerequisite for the production of the symptom.

Lhermitte et al. (1924), discussing the mechanism in multiple sclerosis, considered that direct pressure on or elongation of sensory fibres stripped of their insulating myelin sheaths, was the cause and thought that it was analogous to Tinel’s sign in peripheral nerve lesions.

In view of the fact that Lhermitte’s sign can be produced by extrinsic lesions impinging on the cervical cord, Alajouanine et al. (1949) thought that demyelination was not an essential requirement, but Jones (1964) in a carefully reasoned paper considered that the latent interval when the response followed trauma or radiation damage strongly suggested that the presence of medullary changes was necessary, as the same mechanical factors which operated when the sensations appeared were, in fact, also present in the earlier and silent period. He also pointed out that, in addition to the fact that the cervical cord may impinge on the vertebrae in front and on the ligamentum flavum behind on precipitate movement of the cervical spine, it also has its posterior fibres elongated on gradual flexion and may be distorted in the coronal plane by the restricting action of the denticulate ligaments. These stresses are greatest at the mid-line of the posterior columns and on the posterolateral fibres of the lateral spinothalamic tracts, which in both cases would explain the common distribution of Lhermitte’s sign in the lumbosacral segments.

In subacute combined degeneration of the spinal cord, Pant, Asbury, and Richardson (1968) pointed out that, although the axons are affected, the most heavily myelinated sheaths, particularly in the posterior columns in the cervical and thoracic cord, appear to be more severely involved in the early lesions and that the changes are potentially reversible.

In subacute combined degeneration of the cord, Lhermitte’s sign is a common early symptom and in the two patients described above was the presenting one. A lesion of the posterior columns was present in all 11 patients with Lhermitte’s sign, which suggests that demyelination at that site is the underlying pathological
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process. In neither of the two cases was there narrowing of the cervical canal, which supports the view of Jones (1964) that stretching of the fibres of the posterior columns on neck flexion is the mechanism by which the demyelinated fibres are stimulated. The disappearance of the symptoms and signs of the posterior column lesions within a few months of the commencement of treatment is explained by remyelination.

Lhermitte's sign is a much more common early symptom of subacute combined degeneration of the cord than is generally realized. Failure to appreciate this fact may lead to an erroneous diagnosis of multiple sclerosis, particularly if the patient is young as in case 2, or of cervical spondylosis in older people. As a consequence, correct treatment with vitamin B12 may be delayed until irreversible spinal cord lesions have developed.

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