Paravertebral muscles in disease of the cervical spine

S B Wharton, K K Chan, J D Pickard, J R Anderson

Abstract

Objectives—Cervical spine disorders are common in the older population. The paravertebral muscles are essential to the support and stabilisation of the cervical spine but have been little studied. The aim was to determine whether pathological changes develop in these muscles in patients with severe cervical spine disease, which, if present, might contribute to the pathogenesis and symptomatology of their disorder.

Methods—Open biopsies of superficial and deep paravertebral muscles were obtained during the course of surgical procedures to alleviate cervical myelopathy. Most of these patients had cervical spondylosis or rheumatoid arthritis involving the cervical spine. The biopsies were compared with muscle obtained at necropsy from patients without a history of cervical spine or neuromuscular disorder.

Results—Muscle from both the study and control groups showed a similar range and severity of abnormalities. In several patients, grouped fibre atrophy suggested chronic partial denervation. Most biopsies showed type 1 fibre predominance and selective type 2 fibre atrophy. Ragged red fibres were a frequent finding and electron microscopy disclosed accumulations of mitochondria, a small proportion of which contained rounded, or longitudinally oriented, single osmiophilic inclusions. Fibres containing core-like areas were also frequent. These pathological features were seen with increasing severity and frequency with increasing age.

Conclusions—The paravertebral cervical muscles develop pathological abnormalities with increasing age with both neurogenic and myopathic features, the pathogenesis of which is probably multifactorial. Such a muscle disorder would be expected to be accompanied by functional impairment which may contribute to the development and symptomatology of cervical spine disease with increasing age.

(J Neurol Neurosurg Psychiatry 1996;61:461–465)

Keywords: paravertebral muscles; cervical spine disease; age related myopathy

Diseases of the cervical and lumbar spine are a common cause of symptoms and loss of work-
Results

Muscle biopsies from both the study and control groups showed a high prevalence of pathological abnormalities. The features described below were seen in both groups with a similar prevalence and severity (fig 1). Most patients showed predominance of type 1 fibres. Selective type 2 fibre atrophy, assessed subjectively, was found in 60%–70% of patients and was, in some patients, very severe. Evidence of denervation, in the form of grouped atrophy and angular atrophic fibres, was seen in four and two patients in the study and control groups respectively.

Ragged red fibres were seen in 60%–70% of patients in both groups (fig 2A). These fibres showed an excess of lipid and stained intensely with NADH-TR. Several patients showed fibres with a negative cytochrome oxidase reaction, particularly when many ragged red fibres were present. Some of the ragged red fibres were themselves negative for cytochrome oxidase activity (fig 2B), although others had a positive rim.

Core-targetoid fibres were seen in 40%–50% of patients in both groups, consisting of rounded areas of absent reactivity on histochemical preparations for NADH-TR (fig 2C). In most of these patients they were present in 10% to 20% of the fibres, ranging from less than 5% to about 30% of fibres. In one patient in the study group, nemaline rods were seen. Scattered necrotic fibres were seen in 40%–50% of patients; these were not associated with inflammation. Inflammation, when present (in 27% of the study group and 13% of controls), was mild and focal, consisting of a few chronic inflammatory cells in the endomysium which did not invade muscle fibres.

Electron microscopy confirmed that ragged red fibres represented fibres in which there were subsarcolemmal and intermyofibrillar accumulations of mitochondria. Most of the accumulated mitochondria did not appear morphologically abnormal, but inclusions were seen in a few. The most usual type consisted of single, rounded, amorphous inclusions, sometimes seen in longitudinal section.

More rarely, an inclusion with internal structure was found (fig 3A). In patients in whom core-targetoid structures were found, electron microscopy showed fibres with central disruption and streaming of the myofibrillar architecture (fig 3B).

As noted, the prevalence and severity of these abnormalities were similar in the study and control groups, and in the study group there was little apparent difference between the patients with rheumatoid arthritis and cervical spondylosis. The prevalence of pathological abnormalities increased with age in both groups, being much less in patients under 60 years of age. This was illustrated by an increase in the mean total score of histological abnormalities with age. The increase in pathological findings with age was also reflected in an increased prevalence of patients with ragged red fibres (study group < 59 years 0%, > 60 years 75%; control group < 59 years 19 to 99) years. Deaths were mostly related to vascular disease or malignancy, with sudden traumatic death accounting for some of the younger patients. None of the patients had a history of cervical spine disease or neuromuscular disease, and there were no patients with rheumatoid arthritis.

In the study group, biopsies were taken at the approximate level of the disease (mostly mid-cervical) from the semispinalis capitis (superficial) and semispinalis cervicis (deep) muscles at the time of surgery to carry out cervical fusion. In the necropsy population, biopsies were taken from the same group of muscles at roughly the mid-cervical level within 48 hours of death (usually within 24 hours). Biopsies were snap frozen in isopentane and stored at −70°C until use. Frozen sections were cut in a cryostat and stained with haematoxylin and eosin, modified Gomori’s trichrome, periodic acid Schiff, and oil red O. Histochemistry was performed for NADH tetrazolium reductase (NADH-TR), cytochrome oxidase, ATPase at pH 9.4, 4.6, and 4.3, and acid phosphatase. Specimens for electron microscopy were fixed in glutaraldehyde, stained with uranyl acetate and lead citrate, and were examined in a Philips EM410 electron microscope. Ragged red fibres were quantified in biopsies from the deep muscle group on a MOP dedicated digitiser for quantitative microscopy (Kontran, Messgerate, GMBH), counting at least 200 fibres.

The prevalence of histological abnormalities with age was compared in the two groups. A score was simply derived for each biopsy by awarding a point for each of four main histological abnormalities—namely, the presence of ragged red fibres, selective type 2 atrophy, core-targetoid fibres, and necrotic fibres, and summating the result for the superficial and deep biopsy (therefore allowing a maximum score of eight). The score does not attempt to weight the histological abnormalities and is simply designed to facilitate comparison between groups.
29%, > 60 years 88%). The mean percentage of ragged red fibres per muscle also increased with age in both the study and control groups, and indeed was greater in the control group at all ages. The range of values for percentage of ragged red fibres per muscle also increased with age, becoming very wide in the older population, and ranging from several patients with no ragged red fibres to patients with considerable numbers (study group < 59 years, mean percentage of ragged red fibres per muscle 0%, range 0%, interquartile range (IQR) 0%; study group > 60 years, mean 1.89%, range 0–12.38%, IQR 0–1.19%; control group < 59 years, mean 0.17%, range 0%–0.77%, IQR 0%–0.19%; control group > 60 years, mean 4.89%, range 0%–34.12%, IQR 1.07%–4.69%).

**Discussion**

We have shown profound pathological abnormalities in both superficial and deep cervical paravertebral muscle groups. The presence of type 1 predominance is perhaps not surprising in a group of muscles with a largely postural function. The presence of denervation in several patients, indicated by angular atrophic fibres and grouped atrophy, can probably be accounted for by nerve root compression due to bony abnormalities of the cervical spine. Core-targetoid fibres, present in a higher proportion of patients, may also reflect denervation. In addition to these findings, however, there is a high prevalence of myopathic features, including type 2 fibre atrophy, ragged red fibres, and scattered necrotic fibres.

The prevalence and severity of these abnormalities was similar in both the study group and the control necropsy population. Indeed, although a slightly higher proportion of the study group showed denervation changes, the prevalence of other abnormalities such as ragged red fibres, was higher in the control group. A necropsy group is, clearly, not the perfect control (an age and sex matched, otherwise well population, biopsied in life with imaging established healthy cervical spines), and some may have had a degree of cervical spondylosis which is common with increasing age. Nevertheless, our control population had no clinical history of cervical spine disease and none had been subjected to operation for such disease, so that any spinal disease present was certainly much less severe than in the study group. This being the case, our data reject the hypothesis that the pathological changes in the muscles are secondary to the cervical spine disease, there being no difference between the
two groups. Rather, in both groups, the pathological abnormalities become more severe and more prevalent with increasing age, so that this seems to be an age related muscle disorder.

Selective type 2 fibre atrophy was a common finding and is a well recognised, non-specific finding with many causes, which, in our patients, might include various combinations of disuse atrophy, intercurrent illness, and drugs (for example, corticosteroids). The age related presence of ragged red fibres is a more intriguing finding. These are, of course, associated with primary mitochondrial diseases. However, ragged red fibres occur in other diseases, not primarily considered to be disorders of mitochondrial DNA such as inclusion body myopathy, in which they are associated with mitochondrial DNA deletions. In this disease the ragged red fibres have been suggested to arise by clonal expansion of mitochondrial DNA with a deletion in regenerating muscle after segmental necrosis. This is a mechanism which may be relevant to our patients, in which there is evidence of muscle fibre damage. Ragged red fibres have also been found in lumbar spinal muscles of patients with progressive lumbar kyphosis and in patients operated on for lumbar spinal disc disease. We have also found them in lumbar spinal muscles (unpublished observations) suggesting that they may be a particular feature of aging in postural muscles. Cytochrome oxidase deficient fibres increase with aging in muscle, and deletions of mitochondrial DNA increase with aging, possibly secondary to prolonged free radical damage. It remains to be determined whether the age related increase in ragged red fibres in the paraspinal muscles in our study is arising secondary to acquired alteration in mitochondrial DNA, or to some other mechanism.

It is also of note that ragged red fibres and core-targetoid fibres, as found in our patients, have both been produced in an experimental model of acute muscle ischaemia. In particular, in this ischaemic model the ragged red fibres ultrastructurally showed mitochondria containing similar, amorphous, single, rounded electron dense inclusions as seen in our patients, a type of inclusion not, to our knowledge, described in mitochondrial cytopathies.

In conclusion, the paraspinal muscles in both our study group and control group showed increasing pathological abnormalities with age. Both myopathic features and features of denervation were seen, so that this age related muscle disorder may have both a myogenic and neurogenic component. Whatever the precise pathogenesis, it seems probable that such striking pathological findings are associated with functional impairment of this group of muscles with increasing age, and thus may contribute to the pathogenesis and symptomatology of cervical spine disorders and to instability of the cervical spine in those with severe bony disease. The state of these muscles is therefore of clinical relevance and methods to improve their strength may be of value in these patients. Non-invasive methods, such as MRI, do provide information as to the state of musculature, and may be of value in the assessment and follow up of some of these patients.

We thank Ms G Ganward for excellent technical work in the preparation of material for electron microscopy and Mr C Burton for help with photomicrography.

NEUROLOGICAL STAMP

Thomas Sydenham (1624–89)

The 17th century physician, Thomas Sydenham, became known as the English Hippocrates. He studied medicine at Oxford where the dentist Robert Boyle and the physician and philosopher John Lock were his friends. John Lock gave a detailed account of trigeminal neuralgia in a series of letters in 1667 and Sydenham made contributions to the same condition. Sydenham’s education was interrupted by the English Civil War. His father and four brothers were serving in Cromwell’s army. Thomas also enlisted and served for four years as Captain of a Troop of Horse after which he returned to Oxford, gaining his MD in 1648. His medical studies continued at Montpellier where Rabelais (1490–1553) had earlier been Professor of Medicine.

Sydenham understood the need for careful bedside observation of clinical phenomena. His attitude towards his contemporaries was indifferent or scornful and he complained bitterly of the opposition and rejection of his own colleagues. His painstaking observations and excellent description of many diseases were recorded in his Observationes Medicæ (Medical Observations) (1676). His fame largely rests on his first hand account of diseases including hysteria and his description of chorea minor. He was a sufferer from gout and his treatise on this subject Tractatus de podagra (1683) is considered a masterpiece. The Dissertatio Epistolariæ (1682) contains his classic account of hysteria. Sydenham’s description of chorea minor, which he saw as a kind of convulsion, is found in his Schedula Monitoria (1686).

Sydenham was one of the first to prescribe iron for anaemia and he popularised the use of quinine yielding cinchona bark introduced from Peru in the treatment of ague or malaria. Syphils he treated by mercurialore, until free salivation occurred and he believed that it was the salivation, rather than the mercury, that wrought the cure. Opium was a favourite drug in the form of a tincture to which he added saffron, cloves, and cinnamon. “Sydenham’s laudanum” as it was called remained a popular remedy for many years. His service to medicine was not to speculate but to lead people back to the bedside. His great success as a physician led Boerhave, when lecturing in Leyden, always to raise his hat on mentioning the name of Sydenham. He had a dignified ethical regard for his patients, holding himself “answerable to God” for their care. This may not have fitted easily into a modern management environment, but his comment “I have consulted my patients’ safety and my own reputation most effectually by doing nothing at all” would, no doubt, have been looked upon more favourably.

After he left All Souls College he moved to London where he spent the rest of his life in private practice. Although he has not been honoured postally, he was honoured in this United States postmark of 1934, 310 years after his death. The postmark contains a spelling error. He died at his house in Pall Mall and was buried in St James’s Church, Piccadilly, where the College of Physicians, in 1810, erected a tablet to his memory.

L F HAAS
Paravertebral muscles in disease of the cervical spine.

S B Wharton, K K Chan, J D Pickard and J R Anderson

*J Neurol Neurosurg Psychiatry* 1996 61: 461-465
doi: 10.1136/jnnp.61.5.461

Updated information and services can be found at:
http://jnnp.bmj.com/content/61/5/461

These include:

Email alerting service

Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

Notes

To request permissions go to:
http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/