Scleroderma “en coup de sabre”: pathological evidence of intracerebral inflammation

J Stone, A J Franks, J A Guthrie, M H Johnson

Abstract
Linear scleroderma “en coup de sabre” (LScs) is associated with neurological complications, the pathogenesis of which is uncertain. A 27 year old woman is reported on who developed epilepsy and focal neurological signs in association with LScs. Brain MRI demonstrated predominantly ipsilateral relapsing and remitting grey and white matter lesions. Analysis of CSF and pathology obtained at brain biopsy provides evidence of an inflammatory process which may be amenable to immunosuppressive treatment.

Keywords: scleroderma “en coup de sabre”; central nervous system inflammation; progressive facial hemiatrophy

Linear scleroderma “en coup de sabre” (LScs) denotes linear scleroderma of the frontoparietal area of the head. It has been reported in association with ipsilateral intracerebral lesions, epilepsy, and ocular complications. Ipsilateral white matter lesions have been seen on CT and MRI in patients with LScs, but there are only two reports of neuropathology. We report a unique case of LScs where intracerebral lesions which correlate with the clinical course have arisen and resolved spontaneously as recorded with MRI. Neuropathological findings, intrathecal production of IgG, and the response to immunosuppressive treatment, provide evidence for a potentially treatable primary intracerebral inflammatory process occurring in association with linear scleroderma.

We discuss these findings in relation to the overlapping and sometimes coexistent condition of progressive facial hemiatrophy (PFHA, also named Parry-Romberg syndrome) in which characteristically the lower half of the face becomes unilaterally atrophic.

Case report
In 1989 a 27 year old right handed woman presented with a 3 month history of weakness of the right arm and leg. Aged 15 she had presented with typical LScs affecting the right frontoparietal area of her scalp. This had subsequently progressed over several years but had been static for 5 years. She had been under regular dermatology review and received no treatment. There was no other significant medical or family history. On examination, there was mild pyramidal weakness of the left arm with exaggerated tendon reflexes. There were no other manifestations of connective tissue disease.

Brain MRI initially showed extensive abnormalities in the right hemisphere comprising multiple high signal foci on T2 weighted images involving grey matter, and superficial and deep white matter. There was enhancement with gadolinium on T1 weighted images and mild focal atrophy. The constituents of CSF were normal but electrophoresis showed local synthesis of IgG with an oligoclonal distribution.

Six months later she presented with secondarily generalised seizures. A repeat MRI examination showed partial resolution of the previous lesions but a new area of high signal in the right frontal lobe. Routine laboratory studies and a comprehensive autoantibody screen were negative.

Over the next 3 years her epilepsy remained well controlled on anticonvulsant drugs. Brain MRI in 1992 showed partial resolution of the previous right hemispheric lesions. In 1993 she began to experience difficulties with handwriting, dysarthria, and jaw spasm. Repeat MRI demonstrated a large new high signal lesion on T2 weighted images in the left thalamus, with marked enhancement on T1 weighted images and a new lesion in the cerebellum (fig 1). Because of concern that this might be neoplastic, an uncomplicated stereotactic biopsy of the left thalamic lesion was performed.

Histology showed a focal inflammatory process centred on blood vessels with infiltration but not necrosis of the vessel walls. Lymphocytes and monocytes infiltrated the adjacent brain, which showed astrocytic enlargement, and in one area foamy macrophages in the neuropil merged with a zone of necrosis. In places lymphocytes surrounded single necrotic neurons. Stains for bacteria, protozoa, and fungi were all negative. Polymerase chain reaction (PCR) based techniques to detect herpes simplex virus, measles, and Epstein-Barr virus were negative. The appearances were interpreted as a low grade vasculitis with associated focal cerebral necrosis (fig 2).

Our patient again improved spontaneously until 1994 when she returned with further
secondarily generalised seizures. Brain MRI showed significant resolution of the left thalamic lesion with new areas of high signal predominantly in the right frontal lobe (fig 3). Further CSF examination showed 12 lymphocytes/mm$^3$ with normal protein and glucose. She continued to have seizures and developed a mild Raynaud’s phenomenon in her left hand. She was treated with pulsed methylprednisolone (1g daily for 3 days). A
nantly ipsilateral, cortical and subcortical lesions seen on MRI were described in a 32 year old woman who also had uveitis, Raynaud’s phenomenon, and intrathecal synthesis of IgG.1 This case demonstrated some spontaneous remission of lesions. Liu et al reported ipsilateral changes in two children, consisting of calcification and abnormal grey and white matter interpreted as a migrational abnormality and dysmyelination.2 Chung et al3 have reported on a 27 year old person with a large ipsilateral calcified cortical and subcortical lesion. Our patient is unusual in showing remitting and remitting lesions that are not all confined to the same side as the skin lesion. Coincidental multiple sclerosis is unlikely given the involvement of grey and white matter.

Neuropathology in LScs has been reported in only two patients. Chung et al3 found leptomeningeal band-like sclerosis, intraparenchymal calcification, and ectatic vessels with gliosis but no evidence of inflammation. Dubeau et al4 reported chronic inflammatory changes with perivascular infiltrates, neuronal loss, and gliosis similar to our patient. Further evidence for an inflammatory basis to both LScs and PFHA comes from serological studies which show high frequencies of positive serum antinuclear antibodies and rheumatoid factor.5

The neuroradiological and pathological correlates of PFHA are similar. Fry et al5 reported on five children with upper facial hemiatrophy who had high signal white matter lesions seen on MRI, predominantly in the frontal lobe (four ipsilateral, one contralateral). Terstegge et al6 reviewed most of the available radiological evidence and reported on a similar patient. They also commented on the presence of ill defined sulci in the presence of ventricular dilatation, suggesting meningeal adhesions. Dupont et al7 reported on four patients with parasagittal PFHA who had focal MRI changes consisting of ipsilateral cortical thickening, gyral enhancement, and blurring of the grey-white interface. Hyperintense lesions on T2 weighted sequences were also seen. Cory et al8 described a child with similar but more severe findings including changes interpreted as unilateral focal infarction in the corpus callosum, diffuse white matter changes, and leptomeningeal enhancement. Goldberg-Stern et al described a child who developed facial hemiatrophy with uveitis and epilepsy. Brain MRI disclosed ipsilateral meningeal and basal ganglia lesions similar to those in our patient. Their patient’s lesions and ophthalmological problems seemed to resolve after starting methotrexate.9 There are case reports of angiographic evidence of ipsilateral reversible vessel calibre changes,10 bilateral intracranial aneurysms,11 carotid dissection,12 and arteriovenous malformation.13 There have been five reports of intracranial neuropathology in PFHA since the condition was first described in 1823. Two early German reports14 15 commented on thickened opaque meninges in association with meningeal neovascularisation and atrophic underlying cortex. Eadie et al16 and Merritt et al17 commented only on

Discussion

Linear scleroderma “en coup de sabre”, its relation with PFHA, and their neurological complications have been discussed by authors since the 19th century. Both have been reported in association with focal contralateral seizures and pyramidal signs, ipsilateral intracranial calcification, and ophthalmological abnormalities. Although a distinction can be drawn between the atrophy of the deeper tissues in PFHA and the more prominent skin induration in LScs,2 4 they are almost certainly overlapping conditions and sometimes coexist. The close nature of these two conditions means that they should be discussed in parallel.

There are several reports of MRI findings in patients with LScs. Hyperintense, predominantly ipsilateral, cortical and subcortical lesions seen on MRI were described in a 32 year old woman who also had uveitis, Raynaud’s phenomenon, and intrathecal synthesis of IgG.1 This case demonstrated some spontaneous remission of lesions. Liu et al reported ipsilateral changes in two children, consisting of calcification and abnormal grey and white matter interpreted as a migrational abnormality and dysmyelination.2 Chung et al3 have reported on a 27 year old person with a large ipsilateral calcified cortical and subcortical lesion. Our patient is unusual in showing remitting and remitting lesions that are not all confined to the same side as the skin lesion. Coincidental multiple sclerosis is unlikely given the involvement of grey and white matter.

Neuropathology in LScs has been reported in only two patients. Chung et al3 found leptomeningeal band-like sclerosis, intraparenchymal calcification, and ectatic vessels with gliosis but no evidence of inflammation. Dubeau et al4 reported chronic inflammatory changes with perivascular infiltrates, neuronal loss, and gliosis similar to our patient. Further evidence for an inflammatory basis to both LScs and PFHA comes from serological studies which show high frequencies of positive serum antinuclear antibodies and rheumatoid factor.5

The neuroradiological and pathological correlates of PFHA are similar. Fry et al5 reported on five children with upper facial hemiatrophy who had high signal white matter lesions seen on MRI, predominantly in the frontal lobe (four ipsilateral, one contralateral). Terstegge et al6 reviewed most of the available radiological evidence and reported on a similar patient. They also commented on the presence of ill defined sulci in the presence of ventricular dilatation, suggesting meningeal adhesions. Dupont et al7 reported on four patients with parasagittal PFHA who had focal MRI changes consisting of ipsilateral cortical thickening, gyral enhancement, and blurring of the grey-white interface. Hyperintense lesions on T2 weighted sequences were also seen. Cory et al8 described a child with similar but more severe findings including changes interpreted as unilateral focal infarction in the corpus callosum, diffuse white matter changes, and leptomeningeal enhancement. Goldberg-Stern et al described a child who developed facial hemiatrophy with uveitis and epilepsy. Brain MRI disclosed ipsilateral meningeal and basal ganglia lesions similar to those in our patient. Their patient’s lesions and ophthalmological problems seemed to resolve after starting methotrexate.9 There are case reports of angiographic evidence of ipsilateral reversible vessel calibre changes,10 bilateral intracranial aneurysms,11 carotid dissection,12 and arteriovenous malformation.13 There have been five reports of intracranial neuropathology in PFHA since the condition was first described in 1823. Two early German reports14 15 commented on thickened opaque meninges in association with meningeal neovascularisation and atrophic underlying cortex. Eadie et al16 and Merritt et al17 commented only on
Scleroderma “en coup de sabre”

...cerebrocerebellar calcification and ventricular dilatation. Wolf and Verity also found leptomenigeal fibrosis, degenerative cortical changes, and atypical small tortuous meningeal vessels interpreted as a microvascular malformation. Wartenberg described ipsilateral grey and white matter perivascular lymphocytic infiltration with marked pial and glial proliferation.

The dominant picture in this cerebral biopsy is of an inflammatory process involving blood vessels and resulting in focal cerebral necrosis. This is in keeping with the intense inflammation of the dermis and subcutaneous tissue seen in the early stages of LSCs. We propose that this case, as with the case of Dubéau et al., supports the hypothesis that there is an early cerebral inflammatory stage in LSCs which can later progress to the end stage pathology found by Chung et al. This challenges their conclusions that the process is developmental and is supported by analogous neuropathological findings in PFHA. The oligoclonal response in the CSF, and the response to immunosuppression in our patient indicate that there is an inflammatory or immune mediated mechanism underlying the associated cerebral process in LSCs and that treatment should be considered when clinically relevant.

Intracerebral abnormality should always be considered in patients with localised scleroderma of the head or PFHA. The investigation of choice is MRI and this may identify further asymptomatic cases. The similarity of intracranial manifestations in these two disorders suggests that they share a similar aetiology and are probably overlapping syndromes.

Consent has been obtained from the patient for the publication of this case report.


