Acquired Brown’s syndrome in a patient with combined lichen sclerosus et atrophicus and morphea

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SUMMARY A 49-year-old woman with generalised lichen sclerosus et atrophicus and morphea developed bilateral Brown’s syndrome. Some of the skin lesions were in the vicinity of the trochlea. A characteristic feature of morphea is subcutaneous fibrosis, so we postulate that the cause of the Brown’s syndrome was mechanical tethering of the superior oblique tendon by deep subdermal fibrosis. Histopathological diagnosis was made from biopsies of similar lesions on the patient’s face.

In 1950 H W Brown1 first published his series of seven cases with the following clinical features: slight downdrift of the affected eye on adduction; limitation of elevation on adduction; widening of the palpebral fissure on adduction; no overaction of the ipsilateral superior oblique; V exo pattern; positive traction test.

He suggested that this syndrome was due to congenital paralysis of the inferior oblique muscle resulting in a short anterior superior oblique tendon sheath. In a further publication2 he divided his cases of superior oblique tendon sheath syndrome into two types—true and simulated cases. The true cases were those with a congenitally short anterior sheath of the superior oblique tendon. These patients had a positive traction test. The simulated cases were all others, either congenital or acquired. The congenital simulated cases were postulated to have a thick posterior tendon or firm attachment of the posterior sheath to the tendon. The acquired cases were of inflammatory origin.

In 1975 Parks and M Brown3 reviewed the theories of true Brown’s syndrome and were unable to demonstrate H W Brown’s findings of a short anterior sheath except in two of the 24 patients in whom they had surgically explored the orbit. They suggested that the usual cause of Brown’s syndrome was a restrictive connective tissue band situated posteriorly and inferiorly to the globe. In 1977 Parks4 suggested that Brown’s syndrome was due to a taut tendon for which tenectomy was effective treatment and remarked that the superior oblique tendon did not have a sheath, but that the anterior half of the tendon actually penetrated Tenon’s capsule, to which it was attached by an elastic connective tissue sleeve.

Whereas true Brown’s syndrome, which often resolves spontaneously or can be treated surgically, has a limited number of causes, acquired Brown’s syndrome is a diverse group that is often difficult to treat, particularly if the syndrome is caused by trauma. Acquired Brown’s syndrome occurs rarely and usually in an older age group than true Brown’s syndrome of childhood.5,6 Non-traumatic inflammatory causes of acquired Brown’s syndrome have been reported to include frontal sinusitis,7 rheumatoid arthritis,8,9 and juvenile chronic arthritis.10 No obvious cause may be found in some cases.11 Traumatic causes are either surgical (following frontal sinus surgery12 or a superior oblique tuck13) or more commonly non-surgical,14 when a sharp instrument enters the upper nasal quadrant of the orbit. Occasional cases have been reported as having other unusual causes such as a secondary deposit in the orbit from a primary carcinoma of the prostate.15

The present case report describes an additional cause of acquired Brown’s syndrome—combined lichen sclerosus et atrophicus and morphea. An understanding of the histological changes in this skin condition indicates one mechanism for the diplopia in some cases of acquired Brown’s syndrome and thereby provides a rational basis for the management of the ophthalmic problems in such cases.
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Case history

A 49-year-old woman with a clinical and histological diagnosis of lichen sclerosis et atrophicus and morphoea was referred from the dermatology clinic in August 1984 because of increasing diplopia. She had first noticed double vision 12 months previously. This had become increasingly noticeable, particularly on looking up and to the left.

One month after the onset of diplopia the patient developed discrete firm white plaques round the right eye extending on to the forehead. Similar plaques became apparent round the left eye. She noticed a bullous lesion on her hip and a large skin lesion on her neck, for which she sought the advice of a dermatologist. A biopsy was taken from the bullous lesion on her right hip, and histological examination indicated a diagnosis of morphoea.

Within a few months she also developed patches of scaly depigmentation on her right cheek and on the front of her head, with a larger area affecting her forehead in an 'en coup de sabre' distribution (Figs. 1A, B). This lesion extended to the midline of the forehead, up into the scalp, where there was associated frontal alopecia, and down over the superonasal quadrant of the orbit to the lid margin nasally with associated loss of cilia and supercilia. There was a thickened area of skin over the region of the right trochlea and the right interpalpebral fissure was narrowed by 1-5 mm in comparison with that on the left. The discrete lesions over the left brow and superonasal orbit became more noticeable with time. Later two further biopsies from her cheek and forehead showed combined lichen sclerosus et atrophicus and morphoea.

She had a corrected visual acuity of 6/4 in each eye and had a compensatory head posture consisting of chin elevation and a small head turn to the left, with which she avoided diplopia in the primary position. She had an exotropia in elevation and an exophoria in the primary position when wearing glasses. There was slight underaction of her right eye on laevo-elevation and minimal underaction of her left eye on dextro-elevation (Fig. 2). Her diplopia was most marked on laevo-elevation. No click was heard or felt over either trochlea. The Hess chart (Fig. 3A) was consistent with a bilateral Brown’s syndrome, more marked on the right than the left. In the right eye a

Fig. 1A

Figs. 1A, B  Distribution of plaques and en coup de sabre lesion of lichen sclerosis et atrophicus. These lie over the region of the trochleae on both sides.
Fig. 2 Nine positions of gaze showing bilateral Brown's syndrome.

Fig. 3 A: Hess chart demonstrates the apparent underaction of the inferior oblique muscles and overaction of the superior oblique muscles of the bilateral Brown's syndrome. August 1984. B: Hess chart shows slight worsening six months later.
forcedductiontestconfirmedmarkedlimitationof
elevationonadductionandslightlimitationof
elevation in abduction. In the left eye there was slight
limitation of elevation in adduction. During the
following six months new skin lesions became
apparent and existing skin lesions progressed. The
diplopia on elevation became more noticeable (Fig.
3B). She continues to be followed up and there have
been no further changes of her acquired Brown’s
syndrome.

HISTOPATHOLOGY
A punch biopsy of the bullous lesion on her right hip
had shown haemorrhagic subepidermal bullae, a
perivascular chronic inflammatory cell infiltrate, and
periadenexal inflammation with thickening of dermal
collagen which was of a hyaline appearance. These
features indicated a histological diagnosis of
morphae.

Two further skin biopsies were examined: one
from the forehead at the margin of the largest (en
coup de sabre) lesion and one from a small lesion on
the right cheek. Both biopsies were similar in appearance
in histological sections (Fig. 4). The epidermis
was atrophic and separated from the reticular dermis
by an area of oedema within the collagen. There was
no evidence of liquefaction of the basal layer of the
epidermis. Follicular plugging was observed, and
there was fibrosis of the dermis extending into the
subdermal fat. A perivascular acute and chronic
inflammatory cell infiltrate was seen, particularly in
the deeper layers of the dermis. Although there was
follicular plugging, the distribution of the inflamma-
tory infiltrate was not consistent with discoid lupus
erthematous. The deep extension of fibrosis, com-
bined with epidermal atrophy, follicular plugging,
and oedema of the papillary dermis indicates a
histological diagnosis of combined lichen sclerosis et
atrophicus and morphae.

Discussion
In this case of bilateral acquired Brown’s syndrome
the diplopia preceded the manifestation of the skin
lesions and increased in severity as the skin lesions
grew in extent. In morphae the fibrosis is believed to
start in the lower dermis,14 which supports the view
that the diplopia was caused by morphae of the
overlying skin.

Ophthalmic problems previously reported to be
associated with morphae are many and various.
They include loss of cilia and supracilia, tarsal
atrophy, iritis, iridopapillary atrophy,17 unilateral
glaucoma,18 heterochromia, 19 atrophy of skin and
muscle including extraocular muscle occurring with
en coup de sabre lesion,20,21 perilimbal vascular
anomaly,22 corneal opacity and fundal changes.23
This patient had loss of cilia and supracilia, raised intra-
ocular pressures, and episcleritis as complications of
her morphae in addition to Brown’s syndrome.

Lichen sclerosus et atrophicus has been described
as occurring on the eyelid,24 producing lid notching
and ectropion.25 Combined lichen sclerosus et
atrophicus and morphae is rare26 and is thought to
be a manifestation of the same disease process; it
tends to behave as morphae alone and is usually
self-limiting or gradually progressive; with no treatment is effective.

The histopathology of skin lesions on the face, clinically similar to those overlying the trochlea, showed the infiltration of deep fibrosis into the subdermal fat which is part of the morphea component of the combined skin disease. We postulate that bands of fibrosis extended into the perisheath region round the trochlea and mechanically limited the passive movement of the superior oblique tendon in elevation in adduction.

The function of the trochlea has been studied by various authors. Helveston et al. have shown from light and electron microscopical examination of eight specimens of trochlea that the tendon is separated from the trochlea by an encircling vascular layer and an outer thin bursa-like space. The tendon is composed of parallel fibres with low adhesion to each other along their length, so that movement through the trochlea occurs by differential sliding of the fibres in a telescoping fashion, with only the central fibres completing the whole excursion. Excess fluid in this bursa or distension of the vascular sheath is postulated to restrict movement through the trochlea, causing an acquired Brown's syndrome which may be associated with a click.

Seve demonstrated from 54 embryological and fetal specimens the existence of fine trabeculae between the tendon and the trochlea, the persistence of which into adulthood may prevent this sliding movement and limit the excursions of the tendon in the congenital form of Brown's syndrome.

Koonnef considered that the orbital connective tissue and the extracocular muscles functioned as a single anatomical entity. He has demonstrated from histological thick sections of the orbit that the superior oblique tendon has many connective tissue bands with the medial aponeurosis of the levator muscle and also large numbers of septa passing to the globe along the course of the tendon from the trochlea to the posterior surface of the eye. Accordingly, it is not surprising that perisheath scarring in the medial upper quadrant should mechanically limit passive movement of the superior oblique tendon to produce a Brown's syndrome.

Repeated peritrochlear steroid injections have been reported by Beck and Hickling to be effective in cases of Brown's syndrome occurring with rheumatoid arthritis and in acquired cases presumed to be due to acute stenosing tenosynovitis. Peritrochlear injection of steroids was not carried out in this case, as morphea rarely responds to local infiltration of steroids or systemic treatment. This is probably because there is already established fibrosis in the deep dermis by the time the skin lesions have appeared. In addition this patient was able to avoid troublesome diplopia by adopting a tolerable compensatory head posture.

This case shows that morphea of the periorbital skin should be added to the list of causes of acquired Brown's syndrome.

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References
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