The Tolosa-Hunt syndrome

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Defining the syndrome
The syndrome of painful ophthalmoplegia involves periorbital or hemicranial pain, combined with ipsilateral ocular motor nerve palsies, oculosympathetic paralysis, and sensory loss in the distribution of the ophthalmic and occasionally the maxillary division of the trigeminal nerve. Various combinations of these cranial nerve palsies may occur, localising the pathological process to the region of the cavernous sinus/superior orbital fissure.

The constellation of findings described may be due to four major causes: trauma, neoplasm, aneurysm, and inflammation. Comprehensive patient evaluation is essential in establishing the correct diagnosis.

Within the last diagnostic category there is a specific subset of patients who develop painful ophthalmoplegia due to a non-specific inflammatory process in the region of the cavernous sinus/superior orbital fissure. Infrequently, they experience involvement of additional cranial nerves ipsilateral to the ophthalmoplegia, including the optic nerve, mandibular branch of trigeminal nerve, and facial nerve. Having a relapsing and remitting course, they respond promptly to systemic corticosteroid therapy.

The diagnostic eponym Tolosa-Hunt syndrome has been applied to these patients,1 and it is this entity which forms the basis of this review.

Historical review
Almost 50 years ago Tolosa2 reported a patient with left orbital pain, ipsilateral progressive visual loss, total left ophthalmoplegia, and reduced sensation over the first division of the trigeminal nerve. Cerebral angiography disclosed narrowing of the intracavernous segment of the left internal carotid artery. Surgical exploration of the left parasellar region was unremarkable, but the patient died 3 days later. At postmortem, granulomatous inflammation of the affected carotid artery and cavernous sinus was found.

Seven years later, Hunt et al3 defined a clinical entity “of somewhat obscure aetiology” on the basis of six patients. In one patient surgical exploration of the parasellar region showed all structures to be “intact and healthy”. The authors thought that this syndrome of painful ophthalmoplegia was caused by an inflammatory lesion in the cavernous sinus, as described by Tolosa. They characterised it by the following diagnostic criteria:

1. Pain may precede the ophthalmoplegia by several days, or may not appear until some time later. It is not a throbbing hemicrania occurring in paroxysms, but a steady pain behind the eye that is often described as “gnawing” or “boring”.
2. Neurological involvement is not confined to the third cranial nerve, but may include the fourth, sixth, and first division of the fifth cranial nerves. Periarterial sympathetic fibres and the optic nerve may be involved.
3. Symptoms last for days to weeks.
4. Spontaneous remissions occur, sometimes with residual neurological deficit.
5. Attacks recur at intervals of months or years.
6. Exhaustive studies, including angiography and surgical exploration, have produced no evidence of involvement of structures outside of the cavernous sinus. There is no systemic reaction.

In addition, Hunt and coworkers reported the therapeutic efficacy of systemic corticosteroids with prompt, dramatic improvement of signs and symptoms in two patients.

In 1966, Smith and Taxdal1 were the first to apply the eponym “Tolosa-Hunt syndrome” to this entity. They added five additional cases, “because there has been no previous report of this syndrome in the ophthalmology literature and to emphasise the use of steroids as a diagnostic test.” The authors stated that “The administration of large doses of systemic steroids for 48 hours produces a dramatic response in painful ophthalmoplegia, which allows prompt differentiation of these cases.”

Since these early descriptions of the Tolosa-Hunt syndrome, numerous reports have confirmed these findings. The clinical syndrome has been found in virtually every continent of the world.11 The non-specific nature of the pathological process has been confirmed,12-14 as has the dramatic clinical response to systemic corticosteroids.10-14

However, over the past quarter century there has been no progress in understanding the pathogenesis of Tolosa-Hunt syndrome. In terms of clinical management some new information is available thanks to advances in neuroimaging techniques. Initially, radiological studies of patients with the syndrome yielded normal results, but subtle cerebral angiographic abnormalities.15-19 Orbital venography often disclosed abnormalities in filling of the superior ophthalmic vein or cavernous sinus, but these techniques are difficult and therefore
Clinical profile of Tolosa-Hunt syndrome

Tolosa-Hunt syndrome can affect people of virtually any age from the 1st to the 8th decades of life, with no sex predilection. Either side may be affected, and the pain may be unilateral or bilateral. Usually acute at onset, the natural course is altered. Using these neuroimaging modalities, cavernous sinus abnormalities have now been described in Tolosa-Hunt syndrome (see below).

Natural history

As Hunt outlined in his diagnostic criteria,1 Tolosa-Hunt syndrome follows an unpredictable course. Usually acute at onset, the patient's symptoms typically last from days to weeks. Before the use of systemic corticosteroids there was clear evidence that spontaneous remissions occurred. However, recurrences are common, taking place in about a half of reported patients, usually at an interval of months or years from the initial attack.11 24 These recurrences may be ipsilateral, contralateral, or rarely, bilateral.

Although Tolosa-Hunt syndrome is a self-limited illness, it does cause considerable morbidity. Infrequently, residual cranial nerve palsies persist. With the institution of corticosteroid therapy, the natural course is altered.
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Neurosurgical biopsy is only rarely employed as a "last resort", it should be considered in patients with rapidly progressive neurological deficits, lack of steroid responsiveness, or persistent abnormalities on neuroimaging studies.

Treatment

Almost 40 years ago Hunt first documented the beneficial effect of corticosteroid therapy in Tolosa-Hunt syndrome. Unfortunately, since then there is little new information as to optimal dosage, duration of treatment, or alternative forms of therapy. It is clear that spontaneous remissions may occur, but there is no doubt that corticosteroids markedly reduce the periorbital pain. No data are available as to whether treatment hastens recovery of the associated cranial nerve palsies. Although steroids are generally tapered over weeks to months, in some cases prolonged therapy may be necessary. As cautioned previously, positive response to steroids has been reported in various paraspinal neoplasms, including chordoma, giant cell tumour, lymphoma, and epidermoid.

Pathology

Tolosa-Hunt syndrome is caused by an inflammatory process. Tolosa originally described non-specific, chronic inflammation with proliferation of fibroblasts and infiltration of the septa and wall of the cavernous sinus with lymphocytes and plasma cells. Hunt et al corroborated these findings, emphasising the lack of necrosis and pointed out that "...such inflammatory changes, in a tight connective tissue, may exert pressure upon the penetrating nerves." Subsequent reports have shown granulomatous inflammation, with epithelioid cells and occasional giant cells. Necrosis may also be seen. There have been no reports of an infectious organism associated with Tolosa-Hunt syndrome.

Aetiology

The aetiology of Tolosa-Hunt syndrome remains unknown. No information is available as to what triggers the inflammatory process in the region of the cavernous sinus/superior orbital fissure. It seems that the syndrome falls within the range of idiopathic orbital inflammation (pseudotumour). Thus, "non-specific" inflammation typically causes an acute orbitopathy. If located more posteriorly, the inflammatory process involves the cavernous sinus/superior orbital fissure, producing the Tolosa-Hunt syndrome. Rarely, the inflammation may spread intracranially.

Differential diagnosis

Because the diagnosis of Tolosa-Hunt syndrome can be made only after other disease processes have been excluded, it behoves the clinician to be familiar with the differential diagnosis of painful ophthalmoplegia. In fact, during the initial patient evaluation there are often no clues in the history or physical examination to distinguish Tolosa-Hunt syndrome from other causes of painful ophthalmoplegia. Therefore, the clinician should be aware of: (1) causes of paraspinal syndrome and (2) other entities producing painful ophthalmoplegia.

Table 1 Diagnostic evaluation of Tolosa-Hunt syndrome

<table>
<thead>
<tr>
<th>Haematological tests:</th>
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<tbody>
<tr>
<td>Complete blood count</td>
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<tr>
<td>Serum chemistry (glucose, electrolytes, liver and renal function)</td>
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<tr>
<td>Erythrocyte sedimentation rate</td>
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<td>CRP</td>
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<td>Haemoglobin A1C</td>
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<td>Fluorescent treponemal antibody test</td>
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<td>Antinuclear antibody</td>
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<td>Anti-SDNA antibody</td>
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<td>Anti-sm antibody</td>
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<tr>
<td>Serum protein electrophoresis</td>
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<tr>
<td>Antinuclear cytoplasmic antibody</td>
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<td>Angiotensin converting enzyme</td>
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Cerebrospinal fluid:

- Opening pressure
- Cell count and differential
- Protein
- Glucose

Culture: bacterial, fungal, mycobacterial

SeroLOGY

- Angiotensin converting enzyme
- Cytology

Neuroradiological studies

- MRI
- CT
- Cerebral angiography

Biopsy:

- Nasopharynx
- Cavernous sinus

Occasional reports have documented a raised erythrocyte sedimentation rate and leukocytosis in the acute stage of Tolosa-Hunt syndrome. Similarly, positive LE cell preparation and antinuclear antibody concentrations have been documented in patients with Tolosa-Hunt syndrome, but there is no convincing evidence that such patients either have or will develop connective tissue disorder.

In general, CSF examination should also be unremarkable, although rarely raised protein and mild pleocytosis have been reported. However, if CSF abnormalities persist, the diagnosis of Tolosa-Hunt syndrome is untenable, and further diagnostic evaluations are required.

As noted above, MRI is the initial diagnostic study to be performed in patients with a disorder of the cavernous sinus or superior orbital fissure. Computed tomography is a very useful adjunct in detecting bone changes (for example, erosion, hyperostosis) as well as (perisellar) calcification. Cerebral angiography has detected abnormalities in the intracavernous carotid artery in patients with Tolosa-Hunt syndrome. These have been described as “segmental narrowing”, “slight irregularity”, or “constriction”, and will resolve with corticosteroid therapy. The role of non-invasive vascular MRI techniques (MR angiography and venography) has not been defined in patients with the syndrome but these may prove to be valuable adjuncts in diagnostic evaluation.

Neurosurgical biopsy is only rarely employed to establish the diagnosis. This can be technically difficult and should only be performed by experienced neurosurgeons. It usually involves biopsy of the dural wall of the cavernous sinus. Although generally a procedure of “last resort”, it should be considered in patients with rapidly progressive neurological symptoms.
Table 2 Causes of parasellar syndrome producing painful ophthalmoplegia

<table>
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<tr>
<th>CAUSES OF PARASELLAR SYNDROME (TABLE 2)</th>
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| Craniofacial trauma may produce painful ophthalmoplegia in various ways: basilar skull fracture with ocular motor nerve damage, intracavernous carotid artery injury with subsequent aneurysm formation, or carotid-cavernous fistula. Various vascular causes may produce painful ophthalmoplegia, the most frequent being an intracavernous carotid artery aneurysm. Rarely, giant aneurysms of the posterior circulation, including the posterior cerebral artery, are associated with various conditions including polythemia, sickle cell disease, vasculitis, pregnancy, dehydration, trauma, and intracranial surgery. Regardless of aetiology, cavernous sinus thrombosis is characterised by orbital congestion, proptosis, eyelid swelling, chemosis, lacrimation, and ophthalmoplegia. Pain around or behind the eye is common. Treatment of cavernous sinus thrombosis usually involves anticoagulant drugs, at times thrombolytic agents, and in septic cases, appropriate antibiotic therapy.

As outlined in table 2, painful ophthalmoplegia may be caused by either contiguous or metastatic spread of a neoplasm. Metastatic involvement of the cavernous sinus/superior orbital fissure is generally due to haematogenous dissemination of neoplastic cells. Occasionally, intracranial neoplastic invasion may occur by intraneuror or perineural spread, as in the case of squamous cell carcinoma producing painful ophthalmoplegia many months after local excision of a facial skin tumour.

Two aspects of neoplastic involvement of the cavernous sinus/superior orbital fissure require emphasis. Firstly, it is important to realise that mode of onset and clinical course are not indicative of the type of lesion causing painful ophthalmoplegia. In their review of parasellar syndrome, Thomas and Yoss state that “A sudden onset of symptoms does not weigh in favor of aneurysmal or against neoplastic origin, and gradual onset is not necessarily indicative of a neoplasm...spontaneous single or multiple remissions of symptoms, even of years duration, are unreliable in predicting the nature of the underlying process.” Secondly, as noted above, high dose corticosteroid therapy may initially improve signs and symptoms due to neoplasms.

Inflammatory causes of painful ophthalmoplegia include those due to a specific infectious agent. It is essential that careful CSF examination be done and that cultures (bacterial, fungal, mycobacterial) be obtained. The potential role of a parasal sinus as a cause of painful ophthalmoplegia requires attention. Sinus disease may lead to cavernous sinus involvement, either via contiguous spread of infection or due to sphenoid sinus mucocyte.

OTHER CAUSES OF PAINFUL OPHTHALMOPLEgia

Table 3 summarises other aetiologies of painful ophthalmoplegia in which there is no involvement of the cavernous sinus/superior orbital fissure.

Various orbital diseases cause painful ophthalmoplegia. Typically, the patient presents with “orbital signs”, including proptosis, conjunctival injection, chemosis, and resistance to retrodisplacement of the globe. In addition, the eye may be displaced within the orbit, and there may be abnormalities of the ocular adnexa (for example, lids, lacrimal gland).

Diabetic ophthalmoplegia typically produces an acute, often painful mononeuropathy in either a known or previously undiagnosed diabetic person. Invariably there is recovery of ocular motor cranial nerve function, usually
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Table 3  Additional causes of painful ophthalmoplegia

<table>
<thead>
<tr>
<th>Category</th>
<th>Causes</th>
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<tbody>
<tr>
<td>Orbital disease</td>
<td>Idiopathic orbital inflammation (pseudotumour)</td>
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<td></td>
<td>Contiguous sinusitis</td>
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<td></td>
<td>Macromycosis or other fungal infection</td>
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<td></td>
<td>Metastatic tumour</td>
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<td></td>
<td>Lymphoma/leukaemia</td>
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<tr>
<td>Diabetic ophthalmoplegia</td>
<td>Mononeuropathy</td>
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<td></td>
<td>Multiple cranial nerve palsies</td>
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<tr>
<td>Posterior fossa aneurysm</td>
<td>Posterior communicating artery</td>
</tr>
<tr>
<td></td>
<td>Basilar artery</td>
</tr>
<tr>
<td>Giant cell arteritis</td>
<td>Ophthalmoplegic migraine</td>
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within 3 months. In addition, there are reports of diabetic patients with simultaneous paralysis of multiple ocular motor nerves. Frequently, these episodes are painful, often recurrent, and not particularly responsive to corticosteroid therapy.

Posterior fossa aneurysms may produce either acute, painful ophthalmoplegia, or may present in a more subacute or chronic fashion. The acute presentation is most often due to an aneurysm in the anterior circulation, typically at the junction of the internal carotid-posterior communicating arteries, whereas the subacute presentation is caused by basal artery aneurysms. In both, cerebral angiography is diagnostic.

Giant cell arteritis may also produce painful ophthalmoplegia. The clinical picture may be one of single or multiple ocular motor nerve palsies. In the only pathological study of ophthalmoplegia occurring in giant cell arteritis, ischaemic necrosis of the extraocular muscles was demonstrated. Examination of the ocular and motor cranial nerves was unremarkable.

Ophthalmoplegic migraine is a poorly understood form of complicated migraine. Typically, this clinical presentation occurs in a child or young adult with periodic headache, who develops an ocular motor cranial nerve palsy at the height of an attack of cephalgia, which is primarily unilateral and in the orbital region. Most often involved is the oculomotor nerve, occasionally the abducens nerve, and rarely the trochlear nerve. The paresis lasts for days to weeks after cessation of a headache; recovery is gradual and tends to be less complete after repeated attacks. A family history of migraine is usually obtained. There are reports of enhancement of the extra-axial portion of the oculomotor nerves of patients with ophthalmoplegic migraine when evaluated with contrast-enhanced MRI.

Conclusion
Although the pathogenetic basis of Tolosa-Hunt syndrome remains unknown, from a practical clinical standpoint it can be regarded as a distinct entity which may be simulated by various other disorders. It cannot be emphasized too strongly that patients suspected of having the syndrome require careful evaluation, appropriate treatment, and scrupulous follow up observation.