Orbital granuloma: a rare manifestation of extrapolumary tuberculosis
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DESCRIPTION
An 86-year-old woman presented with a 2-year history of progressive right eye blindness associated with ocular retraction and ptosis, revealing an intraorbital nodule (figure 1A). She mentioned no other symptoms aside from initial ipsilateral purulent conjunctivitis, which resolved under local treatment. No fever, pulmonary symptoms or significant weight loss were reported. The patient’s main medical history consisted of high-blood pressure, right retinal detachment 30 years previously, left hip replacement and binocular cataract surgery a few years prior. Neither she nor her relatives had a history of/or contact with tuberculosis. She had not travelled to any developing countries.

Despite the ophthalmological anomalies, physical examination did not present any abnormality: no adenopathies and no spinal or neurological anomalies. Brain MRI revealed a right orbital mass of $23\times23\times25$ mm with contrast enhancement after gadolinium injection. Whole-body 18F-fluorodeoxyglucose positron emission tomography scan showed...
hypermetabolism at the orbital lesion (SUVmax, 33; figure 1C) and at another lesion facing the anterior peritoneum (SUVmax, 24; figure 1E). Thoracic CT was normal. Repeated interferon-γ release test (QuantiFERON) and tuberculin skin test were negative, as were Bartonella, Rickettsia and HIV serology. CD4 count was 368/mm³ (32%). Routine biochemistry and haematology, including C reactive protein, were normal. No lymphopenia was noted and no leucocytes were found in the urine. ACE was normal. Intraorbital biopsy was performed twice, revealing a giant cell granuloma with caseous necrosis. Two samples were cultured, without finding any mycobacteria. Nocardia PCR, 16S rRNA PCR, and common bacteriological and mycological cultures were sterile. Mycobacterial PCR was inconclusive.

Finally, a 6-month antituberculosis therapeutic trial was initiated, with a 2-month initial phase using isoniazid (5 mg/kg/day), rifampicin (10 mg/kg/day), ethambutol (15 mg/kg/day) and pyrazinamide (20 mg/kg/day), followed by a 4-month continuation phase (isoniazid and rifampicin). After 6 months of treatment, ocular retraction was observed (figure 1B); intraorbital and peritoneal lesions both resolved (figure 1D, F), but the patient remained blind in the right eye.

Orbital tuberculosis (OT) is an exceptional form of extrapulmonary tuberculosis. It can arise from haematogenous spread or local invasion through the paranasal sinuses. All orbital structures may be involved, from periostitis to dacryoadenitis. Usually, orbital cold tuberculosis has presenting features including proptosis, palpable orbital mass lesions and diplopia, but, as in the present case, orbital sclerosis with enophthalmos can also be found. Bone involvement should be screened for. Although OT is usually found in young patients, it can arise at any age. Synchronous tuberculosis lesions should be screened for as they are commonly found. Biopsy is usually performed. Diagnosis can be difficult to prove, even with modern immunological and molecular strategies, and treatment response can be taken as confirmation of aetiology. There is no consensus on duration of antituberculosis chemotherapy, but, because evolution is usually favourable, a 6-month treatment schedule is reasonable. Oral steroid therapy is rarely reported. Surgery may help in some cases.

Learning points

▸ Orbital tuberculoma is an uncommon form of extrapulmonary tuberculosis.
▸ Interferon-γ release test (QuantiFERON) and tuberculin skin test could be negative in patients with extrapulmonary tuberculosis.
▸ Screening for synchronous tuberculosis lesions usually reveals other disease locations.
▸ Prognosis is usually good and standard 6-month antituberculosis therapy can be prescribed.

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