Transverse Myelitis: diagnosis, treatment and management

Transverse Myelitis is an autoimmune disorder attacking the spinal cord, which presents with a wide variety of symptoms involving sensory, motor and autonomic dysfunction. These symptoms may develop very quickly over a few hours or gradually over a period of several weeks, complicating diagnosis for GPs and Casualty staff. TM is not common – estimated 300 cases p.a. in UK – but it is both debilitating and treatable in the acute phase. Therefore early diagnosis and referral to a specialist neurologist is important. Primary-care and rehabilitation specialists also bear the main role in managing after-care for TM patients, two thirds of whom suffer from long-term sequelae.

Diagnosis and Treatment

Some patients encounter TM with rapid onset of weakness and/or bladder dysfunction. This group are more likely to visit A&E. But other patients start with subacute onset of sensory symptoms and little motor dysfunction. These patients usually visit their GPs with the symptoms listed below that occur in varying combinations and sequence.

Sensory dysfunction: 80-94% of patients will have numbness, paresthesias or bandlike dysesthesias, although these may not all be present initially. A light, uncomfortable banding sensation around the trunk is a particularly good indicator of TM. Adults are more likely to present with numbness or paresthesias (e.g. burning, tingling) with a micthoracnic sensory level, whereas children show a higher frequency of cervical spine involvement. Other sensory symptoms include heightened or diminished sensitivity to temperature and alodinia – pain caused by non-painful stimuli such as light touch or even wearing clothes.

Autonomic dysfunction and weakness: Acute urinary retention is typical at the outset, and most TM patients suffer bladder and bowel dysfunction due to loss of sensation, which is often complete in the acute phase. Most patients develop leg weakness. At the maximal level of deficit, 50% of patients have lost all leg movement. Onset of paralysis tends to be rapidly progressive; complete paralysis can occur within hours.

Timely MRI imaging, CSF analysis and lab tests are used for diagnosis, as well as history and physical examination. TM must be distinguished from compressive lesions first of all, and then from other conditions such as Guillain-Barre Syndrome and MS. Rheumalolgical symptoms must be looked for, as TM may also be a presentation of systemic autoimmune disease e.g. lupus, sarcoidosis etc. First-line treatment is high-dose IV corticosteroids for 3-5 days to reduce inflammation, followed by oral steroids taper. Plasma exchange should be considered for severe TM that is refractory to corticosteroids, or for patients with suspected antibody-mediated disease e.g. Neuromyelitis Optica (NMO). When TM seems to be recurrent (NMO-IgG blood test), immunosuppressives should be considered.

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Prognosis

Most people with TM experience some degree of neurologic recovery but are also left with neurologic deficits. Though recovery is more rapid in the first six months after onset, patients can experience some improvement for up to two years and physiotherapy should not be terminated prematurely. Around a third recover with little or no sequelae, one third are left moderately disabled, and one third are left severely disabled. Bad prognostic indicators include back pain at onset, rapid progression to maximal symptoms within hours, spinal shock, and sensory involvement up to the cervical level. Little improvement in the first 3-6 months makes significant recovery less likely.

TM is not always monophasic; recurrence affects a small percentage of patients. In all cases of recurrence, the potential for an underlying disorder should be investigated. Patients with lesions over three vertebral segments may go on to be diagnosed with NMO. In some patients TM may be first manifestation of MS – brain MRI and oligoclonal CSF bands are usually indicative of MS.

Treatments for MS, NMO and TM are different, hence the need for good diagnosis. Long-term TM sequelae are often subtle but very disabling. Neurologists do not do a very good job of aftercare and it is up to GPs and rehabilitation specialists to work together to manage chronic pain, continence, spasticity, fatigue and depression.