A vanishing callosal lesion on brain imaging in a patient with meningoencephalitis

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Case vignette
A 21-year-old previously healthy man had become progressively unresponsive five days after developing high-grade fever. On examination, he was unconscious (GCS 3/15), with equally reacting normal-sized pupils and normal optic fundi, and marked neck stiffness. His blood investigations showed neutrophil leucocytosis (total white cells 21,250/µl; 88% neutrophils) and serum sodium of 112 mmol/l. Magnetic resonance imaging of the brain was normal except for a non-enhancing focal hyperintensity on T2 and FLAIR sequences with diffusion restriction and low signal on ADC map in the splenium of the corpus callosum (Figure). Cerebrospinal fluid analysis showed polymorphocytic pleocytosis (225 cells/µl; 80% polymorphocytes), elevated protein (99 mg/dl) and a CSF: plasma glucose ratio of 0.28. A diagnosis of pyogenic meningoencephalitis was established. The patient made a complete recovery after a 14-day course of intravenous meropenem and vancomycin. MRI repeated 14 days after completion of antibiotics showed complete resolution of the splenial hyperintensity.

Q: What is the radiological diagnosis?

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A: Reversible splenial lesion syndrome

The corpus callosum is a bundle of commissural fibres that connects the left and right cerebral hemispheres. The posterior end of the corpus callosum is known as the splenium. Lesions in the corpus callosum and the splenium have been associated with distinct neurological disorders. Reversible splenial lesion syndrome (RESLES) is one such disorder.

RESLES as observed in our patient, is a rare clinicoradiological syndrome described in association with antiepileptic drug withdrawal, cerebral infection, high-altitude cerebral oedema (HACE), or metabolic disorders (hypoglycemia and hypernatremia) [1]. Rarely it has been reported in association with systemic lupus erythomatosus, anorexia nervosa, vitamin B12 deficiency and hereditary motor and sensory neuropathy.

The pathogenesis of RESLES is mostly attributed to cytotoxic oedema with the exception of HACE in which vasogenic oedema is thought to be the underlying mechanism. The reason for the predilection for the splenium of the corpus callosum remains obscure, but a relative lack of adrenergic tone and failure of autoregulation at this site is one hypothesised mechanism [2]. No callosal disconnection syndromes have been described in RESLES while complete clinical and radiological resolution with time is usual [1]. It usually carries a good prognosis except in those patients with an underlying severe disorder. The recognition of this distinct radiological appearance should prompt appropriate differential diagnosis and reassurance of a generally good prognosis.

References
