Incidental Cystic Encephalomalacia from a Silent Cerebral Infarct

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Text

An 86 year old lady presented with a two year history of progressive cognitive decline. The ACE-R identified global, memory-predominant cognitive impairment (51/100). There was a history of hypertension and hyperlipidaemia and no other medical history. An MRI brain confirmed hippocampal atrophy consistent with a diagnosis of Alzheimer’s disease. It also revealed an additional striking finding of a multi-cystic abnormality in the left medial frontal lobe (Figure 1). The MRI brain was repeated and findings were unchanged one year later.

Cystic encephalomalacia is a region of septated, cavitated signal change that follows an acquired parenchymal brain injury such as trauma, infection or ischaemia. The presence of cardiovascular risk factors along with the precise conformation of the lesion to a vascular territory (in this case, the left anterior cerebral artery) makes cystic encephalomalacia secondary to a ‘silent’ anterior cerebral artery infarct the most likely aetiology here. Giant perivascular spaces (PVS) or Virchow-Robin spaces, like cystic encephalomalacia exhibit iso-intensity, appear clustered and demonstrate relative sparing of the cortex as seen here. However PVS are benign expansions of normal pial-lined perivascular spaces [1] and so follow the course of more distal penetrating arteries. In addition they are typically found in the mesencephalothalamic region but can rarely occur in anterior superior temporal or even cingulate areas [2].

Porencephalic cysts are rare and characterized by communication with the ventricular system [3]. Neuroglial cysts are smooth, well demarcated lesions distinct from the multi-cystic structure described in our patient. A cystic neoplasm would not follow a precise vascular territory or demonstrate iso-intensity with CSF [3] and after a 12 month interval would be expected to progress both clinically and radiologically. The radiological findings were consistent with incidental cystic encephalomalacia from a silent cerebral infarct and hippocampal atrophy supporting the clinical features of Alzheimer’s disease.

References