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**Abstract.** We describe the first detailed histological description of an excised calcified *Taenia solium* granuloma from a patient who developed recurrent seizures associated with perilesional edema surrounding a calcified cysticercus (PEC). The capsule, around a degenerated cysticercus, contained marked mononuclear infiltrates that extended to adjacent brain, which showed marked astrocytosis, microgliosis, and inflammatory perivascular infiltrates. The presence of large numbers of mononuclear cells supports an inflammatory cause of PEC. Immunosuppression or anti-inflammatory measures may be able to treat and prevent PEC and recurrent seizures.

**INTRODUCTION**

Neurocysticercosis (NCC), the most common cause of adult onset epilepsy worldwide, is caused by the cystic larval form of the human tapeworm, *Taenia solium*.† Seizures, the most frequent manifestation of NCC, arise when viable cysts degenerate during the course of natural infection or as a consequence of treatment with anthelminthic medication.†† Seizures commonly occur in the presence of calcified granulomas and are frequently localized to them.‡‡–†† Moreover, recurrent, transient episodes of perilesional edema (PEC) and enhancement, both maximally centered around the calcifications, have been described.‡‡–‡‡ In a recent prospective study of patients with only calcified granulomas and a history of remote seizures, PEC was found in 50% of those patients with recurrent seizures and 8.7% of asymptomatic controls from the same population.‡‡ Because calcified granulomas are the most common radiological finding in endemic populations, PEC is likely frequent in those patients with epilepsy.‡‡

Understanding the pathophysiology of PEC is hampered by the inability to directly study the implicated calcified foci. We report the first complete histological description of a calcified lesion that was surgically removed after repeated episodes of perilesional edema. The lesion contains intense regions of mononuclear infiltrates, suggesting that the perilesional edema is inflammatory in nature.

**CASE REPORT**

A 23-year-old Brazilian male, who immigrated to the United States at the age of 18 years, presented with generalized tonic–clonic seizures in December 2004. Neuroimaging showed three cystic lesions in the right posterior frontal, left frontal, and left posterior occipital lobes consistent with NCC (image not shown). A Western blot assay for cysticercosis antibodies was positive. The lesion in the left frontal lobe showed ring enhancement with edema consistent with a degenerating cyst.

Administration of phenytoin sodium and oxcarbazepine led to control of seizures.

Eight months later, in August 2005, he experienced seizures of a different semiology, consisting of focal left-sided motor seizures initially involving the tongue and then shoulder followed by arrest. Magnetic resonance imaging (MRI) of the brain showed resolution of the left frontal lobe lesion, but a new area of perilesional edema had developed around the ring enhancing non-calcified lesion in the right posterior frontal lobe (not shown). Despite attempts to optimize antiepileptic drugs (AED) and treatment with corticosteroids, he continued to experience focal seizures.

He was transferred to Lahey Clinic in August of 2006. Computed tomography (CT) and MRI imaging showed a calcified lesion with surrounding edema in the right frontal lobe (RFL) (Figure 1A–C). He was treated for 3 weeks with albendazole and 14 days with corticosteroids, resulting in marked improvement of the perilesional edema 1 month after discharge (Figure 1D). He was seizure-free until March 2007, when he again developed recurrent focal seizures and marked increase in the edema around the lesion in the RFL (Figure 1E), despite continued treatment with anti-seizure medication. Intractable seizures with recurrent episodes of status epilepticus persisted over the next 6 months, despite maximal AED. MRI imaging showed varying amounts of edema that waxed and waned over time (Figure 1F). After extensive evaluation, including mapping of the seizure focus to the RFL, the lesion was excised in July of 2007. The patient was seizure-free for a period of 2 years while on levetiracetam (1,000 mg/day) and valproate (1,000 mg/day) until he again presented with seizures associated with perilesional edema around the third cyst in the left frontal lobe that had evolved into a calcified granuloma.

**DISCUSSION**

This case report is the first complete histological description of a calcified *T. solium* granuloma associated with episodes of perilesional edema, and it suggests that inflammation is directly or indirectly involved (Figures 2A and B, 3, and 4). A previously published report of the histology of a calcified granuloma was limited to identification of calcareous...
corpuscles, characteristic bodies of unknown function but specific to cestodes.

Few prior histological descriptions of calcified granulomas are available. These describe nodules with a fibrous capsule without notable inflammation, a hyalinized or necrotic parasite that may or may not contain recognizable remnants or calcifications. These reports and similar earlier descriptions led to the concept that calcified lesions are inactive, have no accompanying inflammation, and play little role in the pathophysiology of disease.

Not all calcified granulomas seem to be the same. The evolution of the cellular granulomas to inert calcifications is not instantaneous and may be slow, hindered, or incapable of further evolution, which results in varying levels of calcification.

Figure 2. (A and B) Photograph of two histological sections of the excised lesion. P = degenerated parasite; I = inflammation; AC = amorphous material with calcification; C = collagen; CC = calcareous corpuscles; B = brain. The arrowed line shows the extent of the capsule. The lesion is roughly organized into concentric layers (A) consisting of centrally located eosinophilic material containing thick, bright ribbons of membranous-like tissue, which is most likely a degenerated cysticercus. A granular calcified layer, often adjacent to the host capsule, partially surrounds the core. A tissue section from another region (B) contains a dense mass of concentrated calcified calcareous corpuscles in the core, which are characteristic of cestodes. The degenerated parasite mass is surrounded by a dense collagenous wall making up the host capsule. A mononuclear infiltrate consisting primarily of lymphocytes, macrophages, and plasma cells courses through the capsule but is particularly dense adjacent to the brain. Eosinophils are present in relatively small numbers and do not predominate. The surrounding brain is markedly abnormal with reactive gliosis, which is denoted by glial fibrillary acid protein immunostaining (not shown), CD3-positive T cells (Figure 4C), and histiocytosis and microgliosis by KP1 reactivity (Figure 4D). There is extensive inflammatory perivascular cuffing in adjacent brain tissue.

Figure 3. Higher power view of calcareous corpuscles, which are characteristic of cestodes. The very dense and large mass likely is a cause of calcification. The arrow points to one of many calcareous corpuscles.
in a cellular granulomatous milieu. Additionally, MRIs show enhancement in a subset of calcified lesions, suggesting dysfunction of the blood–brain barrier and the presence of ongoing inflammation. The phenomenon of PEC itself and the pathology of the present calcified granuloma also support an inflammatory etiology for PEC, which may be caused by episodic failure of inhibition of an ongoing inflammation, the periodic release of antigen leading to initiation, aggrava-
tion of an ongoing inflammatory response, or a combination of these processes. An inflammatory etiology of PEC suggests that anti-inflammatory and/or immunosuppressive measures may be efficacious in the treatment and prevention of recurrences. In fact, one person with recurrent perilesional edema and seizures caused by calcified foci responded dramatically to methotrexate therapy. However, no therapy has been shown to be efficacious, and it is premature to recommend methotrexate use as a policy. Because calcified foci, whether associated with perilesional edema (as shown in the present report) or not, consist of dead calcified granulomas, anti-parasitic treatment is not indicated. Anti-epileptic drugs should be administered, although formal studies showing efficacy are lacking.

Surgical removal of epileptogenic foci is accepted treatment of some forms of epilepsy. Presumably, removal of calcified cysticercal foci after similar criteria might be beneficial. However, the literature documenting control of seizures in neurocysticercosis by surgical removal of the foci is scant. Surgical excision should rest on the individual findings and circumstances of a particular patient.

Figure 4. (A) Higher power view of degenerated parasite and amorphous calcified material. (B) Higher power view of the capsule wall with mononuclear infiltrate. (C) CD3-positive T cells. (D) KP1-positive macrophages and microglia.

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