Amebic Meningoencephalitis: Spectrum of Imaging Findings


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Imaging features of amebic meningoencephalitis are non-specific and have only rarely been described in previous literature.\(^1\) \(^2\) We describe the imaging of 5 proved cases of amebic meningoencephalitis and try to familiarize both the unsuspecting radiologist and the clinician with the diverse spectrum of neuroimaging findings. To the best of our knowledge, this is the largest single short series describing the CT and MR imaging features of this entity.

**Subjects and Methods**

The imaging findings on initial CT and MR imaging (1.5T unit) of the brain in 5 proved cases of amebic meningoencephalitis were retrospectively analyzed. Both non contrast–enhanced and contrast-enhanced MR imaging were performed. The diagnosis was confirmed at autopsy and histopathologic examination in 4 cases and CSF culture in 1 case. The first 4 cases were of granulomatous amebic meningoencephalitis (GAE) and the fifth case was of primary amebic meningoencephalitis (PAM).

**Case 1**

A 14-year-old presented to the neurosurgery department after she collapsed from an episode of tonic-clonic seizures. She had experienced a similar history of seizures with loss of consciousness 3 months before the present episode. There was no history of opportunistic infections and no other significant medical or travel history. On presentation, she was afebrile and had a mild right hemiparesis with neck rigidity. CSF examination revealed a normal glucose level (63 mg/dL), normal white blood cell count, but an elevated protein count (52 mg/dL). CT examination of the brain revealed an ill-defined hypoattenuated lesion in the left temporal-parietal area. MR imaging revealed multifocal involvement of the brain. (Fig 1A-C). The largest lesion in the left parietal lobe resembled a mass lesion and showed a linear pattern of enhancement (Fig 1D). A diagnosis of central nervous system (CNS) lymphoma was considered and a brain biopsy was performed.

The biopsy revealed necrotizing vasculitis with perivascular lymphocytic infiltrate and plasma cells suggestive of infection with rare organisms. Subsequently, the child developed signs of increased intracranial tension, was unresponsive to stimuli, and died 12 days after admission. A partial autopsy and brain examination revealed that a large part of the left cerebral hemisphere on the convexity was necrotic and soft. Findings of microscopy revealed acute and chronic nongranulomatous vasculitis, thrombosis, fibrinoid necrosis, and a large number of single and groups of round large organisms, 15–25 \(\mu\)m, with vacuolated cytoplasm and single nuclei. In addition, axon retraction balls and reactive astrocytes were seen in the involved brain. The meninges were chronically inflamed and showed acute necrosis and a few organisms (Fig 1E, -F) consistent with a diagnosis of GAE.

**Case 2**

A 10-year-old boy presented with a history of bifrontal headaches for 2 and a half months, with vomiting, but with no history of any febrile illness. On examination, no cranial nerve involvement was found. There was mild neck rigidity and bilateral papilledema. Blood biochemistry values were within normal limits. Previous CT and MR imaging showed a hypointense masslike lesion on T1-weighted images with isointense signal intensity on T2-weighted images (Fig 2A, -B). Postcontrast images showed a poorly enhancing lesion. A chronic infective etiology was considered, and a CSF examination revealed an elevated protein count (64 mg/dL). A CSF smear was negative for fungal infection, but culture revealed *Acanthamoeba* organisms. The patient was treated with mannitol, dexamethasone, ketoconazole, and rifampicin. He showed symptomatic improvement and was discharged. Follow-up MR imaging after a gap of approximately 2 months, however, showed persistence of the masslike lesion (Fig 2C).

**Case 3**

A 19-year-old woman presented with a history of headache for 6 months, vomiting for 1 month, and unsteadiness of gait for 15 days. Her history was not significant. Fundoscopy revealed bilateral gross papilledema with positive neck stiffness. CSF analysis revealed the following values: proteins, 60 mg/dL; glucose, 91 mg/dL; and an adenosine deaminase level, <1 U/L. CT of the brain revealed multiple focal oval lesions (Fig 3A, -B) with basilar exudates. Multiple T2 hyperintense oval lesions with perilesional edema and intense postcontrast enhancement were seen on MR imaging (Fig 3C, -D). The patient was started on empiric antitubercular treatment, and antiedema measures were instituted. A biopsy was planned; however, the patient had a recurrent seizure and sustained cardiorespiratory arrest from which she could not be revived. At autopsy, there were multiple discrete hemorrhagic areas in the brain, with a large cystic necrotic area in the left parietal lobe, corresponding with the imaging findings (Fig 3E). Microscopy revealed acute vasculitis, thrombosis, fibrinoid necrosis, and a large number of single and groups of round large organisms consistent with amebic trophozoites on an inflammatory background (Fig 3F).

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**CASE REPORT**

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**SUMMARY:** Primary amebic meningoencephalitis and granulomatous amebic meningoencephalitis are central nervous system infections caused by free-living amebae. We describe the neuroimaging findings in 5 such cases on CT and MR imaging. A spectrum of findings was seen in the form of multifocal parenchymal lesions, pseudotumoral lesions, meningeal exudates, hemorrhagic infarcts, and necrosis in the brain. Familiarity with the imaging findings is important for the diagnosis and management of this nearly universally fatal disease.
Case 4
A 38-year-old man presented in our neurosurgical emergency department with deteriorating vision and persistent vomiting for a week. There was history of headache and occasional vomiting for about 2 months. On examination, the patient was in altered sensorium with papilledema and visual field defects. A contrast-enhanced CT (Fig 4A) and MR imaging (Fig 4B–D) of the brain were performed. The patient underwent surgery with a presumed diagnosis of a mass lesion. Subsequently, his condition deteriorated, and he died on the seventh postoperative day. At autopsy, the pathologic findings were consistent with GAE.

Case 5
A 40-year-old man presented with fever, headache, and vomiting for a week and altered sensorium for 3 days. The headache was holocranial in nature with associated nonprojectile vomiting. His history was not significant. On examination, the patient was febrile with positive neck rigidity. There was no papilledema. The CSF sample showed the following values: proteins, 95 mg/dL; glucose, 27 mg/dL, with random blood glucose of 91 mg/dL. CT of the brain revealed exudates in the perimesencephalic cistern, with right basal ganglia infarction (Fig 5A, -B). The possibility of tubercular or fungal meningitis versus a vasculitis was considered. The patient was started on empiric antitubercular treatment and ceftriaxone; however, he developed refractory hypotension and died. An autopsy revealed a softened brain with multiple infarcts in the basal ganglia, deep cerebellar white matter, and the corticomedullary junctions of both parietooccipital lobes. There was also meningeal thickening and gross exudates, especially in the basilar cisterns. On microscopy, the exudates were positive for the characteristic amebic trophozoites.

Discussion
Rare CNS infections in humans caused by free-living amebae of the genera Naegleria or Acanthamoeba were first described in 1965 by Fowler and Carter and in 1966 by Butt. Naegleria and Acanthamoeba organisms are responsible for causing primary PAM and GAE, respectively, having distinctive epidemiology, pattern of presentation, clinical course, pathology, and imaging findings.
**Fig 2.** A 10-year-old boy with headache and vomiting for 2 and a half months (GAE).

A, Contrast-enhanced CT shows an ill-defined pseudotumoral lesion in the left frontotemporal lobe (arrows), with a patchy linear type of enhancement.

B, T2-weighted MR image shows that the mass lesion is isointense to gray matter (arrows). No perilesional edema or any significant mass effect is seen. There were no other focal lesions.

C, Two-month follow-up contrast-enhanced MR image reveals persistence of the mass lesion, with no significant contrast enhancement.

**Fig 3.** A 19-year-old woman with holocranial headaches for 6 months, vomiting for 1 month, and an unsteady gait for 15 days (GAE).

Non contrast–enhanced CT (A) and contrast-enhanced CT (B) scans reveal multiple hyperattenuated oval lesions on the non contrast–enhanced CT scan (arrows), suggestive of hemorrhage. These lesions show intense enhancement on contrast-enhanced CT (arrows). A large nonenhancing cystic lesion is seen in the left parietal lobe (arrowheads).

Axial T2-weighted MR image (C) shows multiple hyperintense oval lesions with perilesional edema and the cystic lesion in the left parietal lobe. Intense postcontrast enhancement of the lesions is seen on the sagittal contrast-enhanced MR image (D).

Coronal cut section of the brain (E) shows multiple hemorrhagic lesions (arrows). Photomicrograph (F) shows the characteristic rounded amebic trophozoites (curved arrows) on an inflammatory background (H&E, original magnification ×40).
GAE (Cases 1–4)

GAE is a subacute to chronic infection caused by Acanthamoeba and also Leptomyxida organisms. Acanthamoeba species are found in all types of environment. Cases have also been associated with amebic keratitis from contaminated contact lens solutions or with hematogenous spread from a primary source of infection, either a pulmonary focus or a skin ulcer. GAE is known to occur in patients who are debilitated or immunocompromised by AIDS, chemotherapy, or steroid therapy. The clinical course is characterized by a long duration of focal neurologic symptoms unlike the rapid progression and fulminant course of patients with PAM. All our 4 patients with GAE had a prolonged duration of focal symptoms (eg, seizures) present for an average of 2–3 months. None of our patients had an underlying immune compromise or any significant history such as a long illness or steroid use. Findings of enzyme-linked immunosorbent assay testing for HIV status performed for 2 patients were negative. All 4 patients were from a low socioeconomic status, and possibly GAE can occur in this subset of individuals who are otherwise immunocompetent. Leptomyxida order species are, however, known to cause GAE in immunocompetent hosts.

Pathologically, the macroscopic appearance in GAE is one of focal edema of the cerebral hemispheres. Multifocal parenchymal lesions with involvement of posterior cranial fossa structures, the diencephalon, and the thalamus are seen. The presence of trophozoites and cysts, along with a chronic granulomatous reaction containing multinucleated giant cells, is characteristic. Microscopically, there is evidence of leptomenigitis, most prominent adjacent to the parenchymal lesions. There may be severe necrotizing angitis. GAE is diagnosed by identifying Acanthamoeba trophozoites or cysts in CSF/brain biopsy. Cultures of brain tissues or CSF can also reveal Acanthamoeba organisms.

The pathologic literature addresses in detail the findings of GAE, but, to our knowledge, little is written about the imaging findings. Sporadic case reports describing the CT characteristics as multiple enhancing lesions involving the cerebral cortex and underlying white matter with mild mass effect are available. On MR imaging, multifocal lesions showing T2 hyperintensity and a heterogeneous or ringlike pattern of enhancement, with a predilection for the diencephalon, thalamus, brain stem, and posterior fossa structures, are seen. A case report by Kidney and Kim described multiple punctate focal areas of enhancement bilaterally throughout the cerebellar hemispheres as well as a few scattered supratentorial lesions. The random corticomedullary location of most of the lesions as seen in our patients (Figs 1B and 3C) and also in another case report of GAE in an HIV patient supports a hematogenous mode of spread. Intralesional hemorrhage was considered an important diagnostic feature by these authors. In 2 of our patients, there was evidence of intralesional hemorrhage (Figs 3A and 4B), which was confirmed on the macroscopic examination of the cut surface of the brain specimen. Such hemorrhagic lesions can be explained by the necrotizing angitis described in severe cases of GAE and can be an important clue in the diagnosis. The lesions of GAE are thought to represent focal areas of cerebritis or microabscesses. The differential diagnosis includes infarcts from septic emboli, ab-
scesses, toxoplasmosis fungal granuloma, or neoplastic lesions.\textsuperscript{5,10} Solitary space-occupying lesions have also been described in cases of GAE,\textsuperscript{2,10} mimicking a low-grade glioma or lymphoma. This type of imaging was demonstrated in our second patient (Fig 2B, -C).

Imaging findings of GAE are, therefore, either in the form of a multifocal pattern with discrete focal lesions in the corticomедullary junction, as a larger solitary masslike lesion,\textsuperscript{11} or a combination of the 2 findings. The masslike lesion (pseudотumoral pattern) often demonstrates a linear and superficial gyriform pattern of enhancement, which could be a useful indicator of the diagnosis. This pattern possibly represents a combination of enhancement in the overlying inflamed meninges covered with exudates and the actual enhancement of the underlying cortex. The amebae possibly infiltrate along the pial vessels into the brain substance leading to an inflammatory response with border zone encephalitis and microabscess formation. The lesion may even suggest a large vessel infarction, except for the history, inordinate amount of associated edema, and multiplicity of the lesions.

Although no consistently effective treatment exists, surgical removal of granulomatous mass lesions, combined with chemotherapy such as amphotericin B or clotrimazole, provides the best chance of survival as seen in our second patient.

PAM (Case 5)

PAM is caused by Naegleria fowleri, with most patients having an acute onset of symptoms with rapid progression and almost always fatal meningoencephalitis within 48–72 hours. The affected patients are children and young adults with no immunologic compromise. The mode of entry of these thermophilic organisms is through the olfactory tract during contact with contaminated water. In our series, none of the patients revealed such a history, and it is most likely that other means, such as inhalation of dust or soil containing the amebic cysts, was the port of entry.

Pathologic changes in cases of PAM are typified by the extensive damage to the brain parenchyma, ependyma, and meninges.\textsuperscript{9,12} Congestion of the meningeal vessels, edematous cortex with herniation of uncus and cerebellum are other features. Microscopically, there is a purulent leptomeningeal exudate with hemorrhage and necrosis throughout the cerebral hemispheres, brain stem, cerebellum, and upper spinal cord.\textsuperscript{9} Our fifth patient demonstrated these typical findings with extensive purulent leptomeningeal exudates mostly along the perimesencephalic cistern. The exudates contained mainly polymorphonuclear cells but were positive for the characteristic amebae, confirming the diagnosis of PAM.

Imaging features of PAM are nonspecific and, to our knowledge, have rarely been described previously. Findings of CT and MR imaging may be normal early in the disease, with evidence of brain edema and basilar meningeal enhancement subsequent-

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