Neuro and intraocular cysticercosis: A clinicopathological case report

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Abstract: Cysticercosis is one of the most common parasitic infestations in humans. Neurocysticercosis (NCC) is the commonest parasitosis of the central nervous system (CNS), endemic in developing countries and is also seen in developed nations with high rates of immigration from prevalent areas. Co-infection of ocular tissue is also significant and often ophthalmologists come across these lesions in their clinical practice. The mode of treatment in NCC and that of intraocular cysticercosis is somewhat different as far as antiparasitic treatment is concerned. This case report highlights the proper management of this parasitic infestation which involved both eye and brain.

Keywords: neurocysticercosis, endemic, scolex, corticosteroid

Introduction
Cysticercosis is a serious public health problem in several developing countries.1–3 It is estimated that 50 million people worldwide are infected with the taeniasis/cysticercosis complex and 50,000 die from cysticercosis annually.2–4 Most infected individuals are aged between 20–50 years and debilitation from cysticercosis represents an all-around financial blow, with considerable human suffering. Human and animal cysticercosis cause a loss of more than US$3–4 billion annually, which are created by measures for control, medical intervention, and lost production.3,4

Humans are definitive hosts and pigs are the intermediate host for Taenia solium.1,3,5,6 In cysticercosis, humans become the intermediate host by ingesting the eggs of T. solium from contaminated food and water. After entering the intestinal wall, the embryo invades the blood stream and can lodge in various organs such as the central nervous system (CNS), eye, skeletal muscles, and subcutaneous tissue.5,6 The ocular manifestations can be destructive because the cysticercus lesion gradually increases in size, leading to blindness in 3 to 5 years. The death of the parasite causes a noticeable release of toxic yield, leading to intense inflammatory reactions and eye damage.6 Initial medical treatment of intraocular cysticercosis with antihelminthic drugs like albendazole or praziquantel is not known to be remedial. Early surgical removal of the parasite is the treatment of choice.5,6 We present a case of a young lady with intraocular cysticercosis with co-infected CNS.

Case report
A 29-year-old female presented to a tertiary institute of northeast India in January 2009 with complaints of seeing floaters and black spots in her left eye (OS) for past 20 days. She was referred from elsewhere with exudative retinal detachment and vitritis in
the OS. She had a past history of convulsion in October 2008 and was put on an antiepileptic, carbamazepine. She was nondiabetic and nonhypertensive. She was non vegetarian and had consumed pork previously.

On examination, her visual acuity was 20/20, N6 in the right eye (OD) and 20/20–3, N10 in the OS. Anterior segment in both eyes were unremarkable. Vitreous reaction in the OS was 2+ cells. Fundus examination of the OS revealed a colloid cyst of 4.5 × 5.5 disc diameter in the inferior temporal quadrant of the retina (Figure 1). There was associated retinal detachment in the lower half of the retina. B-scan ultrasound in the OS showed one cystic lesion, having a spherical curved echo corresponding to the cyst wall located in the inferotemporal retina with high reflective encircling spot scolex. An A-scan ultrasound demonstrated two high amplitude echoes corresponding to the cyst wall; a high reflective echo was seen when the beam passed from side to side of the scolex in real mode. Vitreous echoes were also seen (Figure 2). Magnetic resonance imaging (MRI) diagnosis showed multiple small coin-shaped degenerative lesions in the both cerebral hemispheres with focal edema (Figure 3). The patient was diagnosed with neurocysticercosis with an intraocular cysticercus cyst in the OS. Routine blood examination showed a high eosinophil count. Pars plana vitrectomy plus intraocular cyst removal with fluid gas exchange and an endolaser and silicone oil injection was done. Postoperative vision in OS was 20/30, N8, and the patient is regularly followed up. A course of oral albendazole (400 mg twice daily for six weeks) and steroid (1 mg/kg body weight in tapering dose for six weeks) was prescribed. The cyst was dissected out of the retina and was submitted to pathology laboratory in 10% formalin. After overnight fixation, the cyst became much more transparent. A parasitologist examined the cyst and found it to be in toto globular, transparent, and thin walled, with a small eccentrically located white area representing the scolex inside it. A gross photograph was taken and the cyst was measured (Figure 4). A histopathological section in hematoxylin and eosin stain showed trilaminated corrugated cell wall structures with well defined protoscolex (Figure 5a) and adjoining suckers and hooklets consistent of cysticercosis (Figure 5b).

Discussion

Human cysticercosis is ubiquitous worldwide including northeast India where consumption of semi- or improperly cooked pork, contaminated salad, and inadequate hygiene...
are all possible causes of taeniasis and thereby can cause autoinfection.\textsuperscript{2,3} Food habits, poor hygiene, autoinfection, or traveling patterns may be responsible for their distribution.\textsuperscript{1–3} Clinical presentations of cysticercosis in human may involve the CNS, eye, or other viscera.\textsuperscript{4} Ocular involvements include cysts in eyelids, extraocular muscles, orbit, conjunctiva, anterior chamber, uvea, retina–vitreous, and optic nerve.\textsuperscript{2,3,5,6} Although there are various reports of isolated ocular and intraocular cysticercosis in the literature, there are few reports of cysticercosis involving the eye and brain (Table 1).

Our patient was a young lady who had a complaint in the left eye and was diagnosed with inferior exudative retinal detachment with subretinal cyst in the left eye. Prior to diagnosis, she had a history of convulsion and headache, which may have been due to the parasite’s involvement in the CNS. After removal of the intraocular cysticercus cyst, she was given a course of oral albendazole and steroid.

The intraocular cysticerci are removed either through an external sclerotomy with choroidal incision or via pars plana vitrectomy.\textsuperscript{6} Trans-scleral approaches are used for

\begin{table}[h]
\centering
\begin{tabular}{|l|l|l|l|l|}
\hline
\textbf{Sl No} & \textbf{Author} & \textbf{Reference} & \textbf{No of cases} & \textbf{Location} \\
\hline
1 & Chadha & 13 & 1 & Intravitreal cysticercosis with exudative RD \\
\hline
2 & Rani & 14 & 1 & Ocular-tuberculosis, acquired cysticercosis of the eye and brain \\
\hline
3 & Jethani & 15 & 2 & Vertical dyopia and nystagmus \\
\hline
\end{tabular}
\caption{Neurocysticercosis with intraocular involvement}
\end{table}

\textbf{Abbreviation:} RD, retinal detachment.
subretinal cyst positioned anterior to the equator. The pars plana vitrectomy or transvitreal approach is used for intravitreal cyst and subretinal cyst located posterior to the equator. Vitreous cysts are more common than retinal or subretinal cysts and the inferotemporal subretinal cyst is the most commonly affected quadrant in the retina.6 The parasite may pierce the retina via choroidal circulation; the large choroidal vessels have a larger flow rate and subsequently the parasite migrates to subretinal space and then into the vitreous cavity. The passage of the endoparasite may incite inflammation and can even result in the formation of a chorioretinal scar.6 The cyst, in gross examination, was very typical with an shining eccentric scolex. Histopathological examination of the parasite showed the scolex and suckers. During follow-up, our patient maintained good visual acuity. Convulsive episodes were reduced during that time. The genesis of convulsion in a patient with neurocysticercosis may be due to inflammation, cerebral tissue gliosis, and preference of the cyst to travel to the cerebral hemisphere in general and frontal and temporal lobes in particular.6,9 Albendazole and praziquantel are the principal antiparasitic drugs used to treat neurocysticercosis.1,5,6,9 Numerous trials showed both these drugs reduce the number of cyst and frequency of seizures.3 Chronic epilepsy is usually associated with calcified granuloma. Cysts that are active or undergoing degeneration (colloidal cysts) are the most epileptogenic.9–11 If there are both intraocular and CNS cysticercus cysts, then the complete intraocular cyst must be removed surgically first, followed by antiparasitic medication and with corticosteroids. We do not want to kill the intraocular live parasite initially because it could induce severe intraocular reactions and thereby cause eventual blindness.

Combined intravitreal surgery by removing the subretinal cysticercus cyst and subsequent medical treatment for CNS parasitosis was safe and effective.

Disclosures
The authors report no conflicts of interest in this work.

References