CASE REPORT

Hydatid brain cyst: A delayed diagnosis in a rural setting during COVID-19

C J Opperman,1,2 BSc Hons (Micro), MB ChB, FC Path (SA) Micro, MMed Path (Micro); J M N Enslin,3 BPhysT, MB ChB, FC Neurosurg (SA), MMed (Neurosurg); J Nuttall,1 MB ChB, FC Paed (SA), MSc (Med) (Paed); A J Brink,4 MB ChB, MMed (Med Micro); S P da Fonseca,4 MB BCh; H D Tootla,1,6 MB ChB, FC Path (SA) Micro, MMed Path (Micro)

1 Microbiology Laboratory, National Health Laboratory Service, Groot Schuur Hospital, Cape Town, South Africa; Division of Medical Microbiology, Department of Pathology, Faculty of Health Sciences, University of Cape Town, South Africa
2 Green Point Tuberculosis Laboratory, National Health Laboratory Service, Cape Town, South Africa
3 Division of Neurosurgery, Department of Surgery, Faculty of Health Sciences, University of Cape Town and Red Cross War Memorial Children's Hospital, Cape Town, South Africa
4 Paediatric Infectious Diseases Unit, Department of Paediatrics and Child Health, Faculty of Health Sciences, University of Cape Town and Red Cross War Memorial Children's Hospital, Cape Town, South Africa
5 Community Service Medical Officer, National Department of Health, Central Karoo District, Western Cape Province, South Africa
6 National Health Laboratory Service, Red Cross War Memorial Children's Hospital, Cape Town, South Africa

Corresponding author: C J Opperman (stefanopperman1@gmail.com)

A previously healthy 10-year-old girl, living in a sheep-farming community in South Africa with exposure to dogs, presented to her local hospital with generalised tonic-clonic seizures. The initial clinical assessment and laboratory work-up were unremarkable. When she presented with further seizures 6 months later, attempts to arrange neuroimaging and specialist assessment were unsuccessful owing to restrictions on routine healthcare services during the SARS-CoV-2 nationwide lockdown. Subsequently, 11 months after her first presentation, she developed focal neurological signs suggestive of raised intracranial pressure. A brain computed tomography scan revealed a left-sided cerebral cyst and imminent tonsillar herniation. An emergency burr-hole procedure was performed to relieve the raised intracranial pressure, followed by definitive neurosurgical excision of cysts. Hydatid protoscolices and hooklets were seen on microscopy of cyst fluid, and treatment with albendazole and praziquantel was initiated. While her infection was treated successfully, long-term sequelae including permanent blindness and hemiparesis could potentially have been prevented with early neuroimaging and surgical intervention.

Hydatid disease is a neglected zoonosis caused by tapeworms in the genus Echinococcus. Cystic echinococcosis, caused by E. granulosus, is most frequently associated with human disease.1 It is endemic in South Africa (SA) and worldwide, but is often missed despite clues such as residence in a farming community, or exposure to dogs or farm animals.2 Dogs are usually the definitive host, with the tapeworm residing in the small intestine and shedding infective ova in the faeces. These infective ova are ingested by an intermediate host, usually a sheep, goats or swine. The ingested ova hatch in the small intestine of the intermediate host, releasing oncospheres that penetrate the intestinal wall and migrate to various organs where they develop into cysts. The definitive host is infected after ingesting cyst-containing organs of the intermediate host. Protoscolices in these cysts evaginate in the small intestine of the definitive host and develop into the adult tapeworm, completing the life cycle. Humans are accidentally infected after ingesting infective ova, usually through exposure to dogs (Fig. 1).1

Hydatid cysts may occur in a single organ or in multiple organs. The liver is the most common site of infection, followed by the lungs.3 Other rarer sites of infection include the heart, kidneys, intestine, bone, bladder and brain.4-6 Brain cysts account for 1 - 4% of disease, with 50 - 75% of these cases presenting during childhood and 18% of all cases having multisite involvement.6,7 As such, a high index of suspicion is required to make the diagnosis of hydatid brain cysts, and timeous neuroimaging is required.

In otherwise healthy and neurologically intact individuals presenting with new-onset seizures, early neuroimaging should also be considered to exclude infective intracranial lesions, which have commonly been described as a cause for seizures in low-income countries (LICs) and low- to middle-income countries (LMICs).6,9-10 Lumbar puncture, if indicated, should also be deferred until such lesions have been excluded, owing to the risk of brain herniation from raised intracranial pressure (ICP). A noteworthy hurdle compounding exclusion of infective intracranial lesions and raised ICP is that neuroimaging and specialist care outside of major cities are sparsely distributed in LICs and LMICs, including SA.11 However, every effort should be made to facilitate early neuroimaging to potentially prevent devastating complications of untreated infections in the brain. Clinicians need to be aware of the differences in aetiology of seizures in LICs and LMICs v. high-income countries (HICs), and should not be misled by guidelines developed for HICs suggesting delayed or no neuroimaging for new-onset seizures.12

Case report
A 10-year-old girl, living in a sheep-farming community in SA and exposed to dogs, presented to her district hospital after experiencing a new-onset generalised tonic-clonic seizure. There was no history of substance abuse or head trauma, and no family history of epilepsy. Clinically she was haemodynamically stable, did not have a fever or signs of meningitis, and had no focal neurological...
...white matter were observed. Once surgical removal of the cysts was completed, the brain surface, dura and soft tissue were irrigated with hypertonic saline, followed by normal saline. Cyst fluid and tissue were sent for microbiological and histological investigation, respectively. Direct light microscopy of the cyst fluid revealed hooklets and protoscolices of *Echinococcus* (Fig. 3A and B), while histological examination included visualisation of the laminated membrane of the cyst wall and a single hooklet. Post-neurosurgery concerns included a posterior cerebral artery infarct, hygromas and pneumocranium, which were monitored with neuroimaging and treated conservatively without intervention. Cerebral oedema was managed with dexamethasone, from which the patient was slowly weaned over 2 weeks. Albendazole and praziquantel for 6 months were used to treat the *E. granulosus* infection, and sodium valproate was continued for ongoing convulsions. Chest and abdominal radiographs, abdominal ultrasound and an echocardiogram were performed to search for liver and lung cysts or cysts at other sites. These investigations were all negative. *Echinococcus* serology (IgG) using an enzyme-linked immunosorbent assay was negative.

**Outcome and follow-up**

Although the hydatid cysts were surgically removed and the patient was commenced on antiparasitic therapy, she had devastating neurological sequelae as a result of untreated complicated infection and delayed diagnosis. She initially had swallowing difficulties and aspiration, which subsequently resolved. However, her right-sided weakness remained, and she developed permanent blindness as a result of bilateral optic nerve atrophy due to sustained raised ICP. She was transferred back to her regional hospital for ongoing support and rehabilitation once she had stabilised. Marked brain improvement (Fig. 2C and D) was noted on CT scans on discharge from the tertiary paediatric hospital, and repeat neuroimaging (magnetic resonance imaging) in 3 months was scheduled.

**Ethical considerations**

All protected health information was removed or anonymised, and individual parent consent was obtained.

**Discussion**

The lifetime risk of having a seizure in the general population is high, with half occurring in childhood, and most occurring...
The aetiology of seizures is also different in LICs and LMICs compared with HICs, with a high burden of parasitic diseases causing new-onset seizures in otherwise healthy individuals in LICs and LMICs. Further clues for such zoonotic infections include residence in a farming community and exposure to dogs and farm animals. Consideration of hydatid disease should prompt early radiological intervention to prevent complications of untreated disease. Additionally, where lumbar puncture is indicated, neuroimaging should be considered first to exclude raised ICP from intracranial lesions such as neurocysticercosis or hydatid brain cysts, which could potentially lead to brain herniation and death.

Lumbar puncture was not performed in our patient, and hydatid disease was not considered as a potential cause of her seizures despite the fact that she lived in a farming community. Unfortunately, neuroimaging was not facilitated after her first seizure. After her second seizure, attempts were made to arrange neuroimaging at her closest regional hospital, 250 km away. These were unsuccessful owing to strict restrictions on healthcare services during the initial SARS-CoV-2 nationwide lockdown. Neuroimaging was only eventually performed in this child as an emergency procedure after she presented with severe acute neurological deterioration and features of raised ICP requiring an emergency lifesaving neurosurgical intervention.

The SARS-CoV-2 pandemic, necessitating task shifting and diversion of the healthcare workforce to prioritise essential and emergency services, resulted in an unfortunate additional barrier to accessing investigations that in this case may have led to earlier medical and neurosurgical intervention and prevention of disabling neurological sequelae. This case demonstrates the need for appropriate early neuroimaging in otherwise healthy individuals who present with new-onset seizures, but specialist care and neuroimaging facilities are sparsely distributed in LICs and LMICs, and when present are often limited to major towns or cities. Accessibility, affordability, long travel distances and extended waiting times for such healthcare resources glaringly reveal the disparity between healthcare systems in well-resourced v. poorly-resourced areas.

Diagnosis of hydatid brain cysts relies on a high index of suspicion, using epidemiological clues such as exposure to dogs or farm animals, and clues from clinical presentation such as seizures, suggesting intracranial pathology. Both solid and cystic intracranial lesions may clinically present very similarly, and neuroimaging is critical to classify lesions broadly into solid or cystic. Further broad radiological traits such as tumour nodules, ring enhancement and calcification may also provide clues that support a potential diagnosis. Radiologically, hydatid cysts are non-enhancing, well-
defined, circumscribed spherical lesions that have a thin wall with smooth margins. Calcification, surrounding oedema or rim enhancement are usually not present. In Africa, hydatid cysts should be considered for any cerebral cyst where no tumour nodule is noted on imaging, and any surgery performed should take this differential diagnosis into account, noting differences in surgical techniques required. Although perilesional oedema is not usually present with hydatid cysts, if it is present, it may indicate a secondary infection or cyst rupture. Smaller ’daughter cysts’ within the main cyst can also sometimes be seen, and additional smaller cysts surrounding the main cyst can indicate prior rupture of the main cyst. Unlike liver and lung hydatid cysts, Echinococcus serology can be misleading in the presence of brain cysts and is usually falsely negative owing to minimal immune responses elicited within this privileged site.

Visualisation of hydatid elements such as protoscolices (parasite larvae) and hooklets on microscopy of cyst fluid post neurosurgical intervention confirms the diagnosis. We treated our patient with combination anti-helminthic therapy using albendazole and praziquantel. Evidence supporting combination anti-hydad therapy is mostly limited to hydatid disease in the liver, lung and peritoneum. The combination of albendazole with praziquantel has demonstrated a greater reduction in the number and/or size of cysts and increased scolicidal activity compared with albendazole alone. These factors are important to consider when managing cysts in organs such as the brain and spine, where recurrence due to spillage or cyst rupture can have devastating consequences. Praziquantel inhibits secondary cyst formation and prevents the vesicular development of protoscolices by an unknown mechanism of action. Albendazole prevents the polymerisation of tubulin into microtubules, blocks the uptake of parasite glucose and results in the depletion of its glycogen stores. The outcome is parasite starvation and inhibition of ova production in the adult cestode. The combination of these two drugs also increases the active metabolite of albendazole (albendazole sulphone) in the cyst fluid compared with patients receiving albendazole alone.

Resection of cysts remains the main treatment modality for brain and spinal hydatid cysts, and combining surgical interventions with albendazole has also shown better outcomes for treatment of hydatid disease than surgery alone. Surgery is complicated owing to the thin wall of the cyst, which is prone to rupture. Spontaneous rupture, or iatrogenic rupture of the cyst during neurosurgical intervention, may result in dissemination with formation of secondary brain cysts or anaphylactic shock. Postoperative mortality is estimated to be ~10% and increases when the cyst is ruptured during surgery. Aspiration of hydatid cysts via a burr-hole is not recommended, but the initial lifesaving procedure to relieve the intracranial pressure in our case outweighed the risk of seeding and secondary complications. Secondary infection (15%), subdural haematoma (15%), postoperative epilepsy (12%) and hydrocephalus (8%) are frequent postoperative complications, with blindness reported rarely. Our patient’s blindness and posterior cerebral artery territory infarct with neurological sequelae was probably a result of brain herniation secondary to raised intracranial pressure prior to neurosurgical intervention, and could potentially have been avoided with earlier detection and intervention.

Teaching points

- Hydatid disease is endemic in SA and is often overlooked despite epidemiological clues such as residence in a farming community or exposure to dogs or other farm animals.
- Hydatid brain cysts or other intracranial infections should be considered in otherwise healthy individuals with unprovoked new-onset seizures in LICs and LMICs, and early neuroimaging is critical.
- Echinococcus serology (IgG) can be falsely negative with hydatid brain cysts. Visualisation of hydatid elements such as protoscolices (parasite larvae) and hooklets on microscopy of cyst fluid post neurosurgical intervention confirms the diagnosis.
- Management of hydatid brain cysts needs to be individualised, depending on the extent of disease and whether the cysts are anatomically amenable to neurosurgery. Ideally, cysts should be carefully resected using a surgical technique to preserve as much healthy brain tissue as possible. During surgery, it is critical to prevent rupture and spillage of cyst fluid, which can result in serious complications.
- Combination anti-helminthic therapy with albendazole and praziquantel is recommended.
- Untreated hydatid brain cysts can be life-threatening and can cause devastating and extensive neurological deficits.

Conclusions

The permanent loss of vision and profound disability in a previously healthy young girl highlight the severity of complications due to untreated intracranial infections, and in particular hydatid brain cysts. Hydatid disease is endemic in SA and yet still often not considered. Although hydatid brain cysts are relatively uncommon, new-onset seizures in an otherwise healthy and neurologically intact child should prompt consideration of infective intracranial lesions, and referral for appropriate early neuroimaging. Clues suggestive of zoonotic infection, such as residence or occupation in a farming community or exposure to dogs and other animals, are a useful tool in primary care assessment. The SARS-CoV-2 pandemic placed additional constraints on our already heavily burdened healthcare system, with a devastating outcome in this case. A strategy should be to put procedures in place to protect routine healthcare services during further waves of the SARS-CoV-2 pandemic and other unforeseen threats to the healthcare system.

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