Neurobrucellosis Presenting with Unilateral Abducens Nerve Palsy

Abstract

One of the rare complications of brucellosis is neurobrucellosis. There have been numerous reports showing clinical forms of brucellosis affecting CNS, such as cranial nerve involvement, myelitis, vascular disease, radiculoneuritis, meningitis, meningoencephalitis, and demyelinating disease. In this case report, we introduce a 2.5 yr old girl with unilateral abducens nerve palsy referred to Baghiyatallah Hospital Outpatient Clinic, Tehran, Iran in June 2015.

Keywords: Neurobrucellosis; Abducens nerve palsy; Neuroimaging

Introduction

Brucella, an intercellular bacterium causing chronic granulomatous infection, resembles tuberculosis and demands a compounded and extensive antibiotic treatment (1). Brucellosis is a zoonotic disease, endemic to certain geographic locations of the world including the Middle East, Central Asia, and Mediterranean countries (2, 3). In humans, this infection is acquired from animals. It occurs because of consuming infected dairy products, coming to direct contact with infected animal organs, and inhaling aerosolized infected particles (1, 4).

Neurobrucellosis (NB) is reported to affect 5%-10% of brucellosis cases with presenting symptoms of the peripheral and central nervous system (5). Several clinical symptoms and imaging anomalies of NB can imitate other neurologic diseases (2, 6).

Case report

The patient was a 2.5-yr-old girl referred to Baghiyatallah Hospital Outpatient Clinic, Tehran, Iran in June 2015 with left eye isotropia for 3 days, fever and occasional vomiting for 20 days. She was treated by amoxicilline-clavulanic acid, acetaminophen and ibuprofen. She had the history of consuming fresh un-posturized milk.

In physical exam, we found low grade fever (38 degree of centigrade) and internal deviation of left eye (Figure 1). Other systemic and neurologic exams were normal. Informed consent was taken from patients parents and Ethics Committee of the hospital approved the study.

Neuroimaging results

Brain MRI: There was a round hyperintense lesion at parasagittal cortex of left parieto-occipital area on T1-weighted images (Figure 2). There was another small hyper intense lesion at frontal subcortical white matter, near the anterior horn of left lateral ventricle on T1-weighted images as well (Figure 3). Orbital MRI was
Hematogenous dissemination may happen because of ingesting contaminated products, resulting in the absorption of the bacteria by the reticuloendothelial system, and ultimately, involvement of other organs (7). Brucellosis is a disease that manifests itself in various forms, although fever has been a consistent feature, strongly unpleasant odor and constitutional symptoms generally appear as a result. Mostly, hepatosplenomegaly and lymphadenopathy can be present. Hepatitis, osteoarticular disease and genitourinary system involvement are commonly seen in brucellosis but central nervous system involvement is associated only in 5%-7% of cases most studied (1).

Neurobrucellosis is rarely seen in children, the symptoms may include fever, headache, vomiting, fatigue, depression, back pain, muscle tension and spasms. In addition to such symptoms, meningeal signs, absent deep tendon reflexes (DTR) or upward plantar reflex plus increased DTR and other symptoms of systemic brucellosis can be seen. Factors that could deteriorate the condition include sensorial or motor anomalies at different degrees, cranial nerve retention, convulsions, cerebellar dysfunction, coma and brain abscesses (8, 9). Development of NB is likely at any stage of the disease (5). Reports have shown numerous clinical forms of brucellosis affecting the CNS. Such forms may include radiculoneuritis, myelitis, cranial nerve involvement, meningoencephalitis, and demyelinating or vascular disease (3, 5, 7, 10).

Chronic meningoencephalitis is among the most common forms of NB that affects the CNS, in which high protein levels and increased lymphocytes are detected in the CSF (11). Due to the slow-growing property of Brucella bacterium, CSF and blood cultures can show negative results. Therefore, serological methods are generally carried out to optimize the diagnosis. Detecting Brucella antibody in CSF is conclusive (4). The major clinical appearance of NB in children is depicted as acute meningitis or meningoencephalitis. Usually, in CSF assessment, brucellosis may not be included thus, it could make the diagnosis of the condition difficult (12, 13). Radiologic correlation in NB has been claimed variable (2).

Discussion
Brucella is considered as a crucial human pathogen, and is local to the under developed countries of the world.

Laboratory findings
CBC: WBC= 11600 (Poly: 44.2%, Lymph: 46%, Mono: 9%), Hb= 10.4, HCT=31.6, Platelet= 244000
ESR=17
CRP= Negative
Widal test: Negative
Wright Test: positive (titer →1.640)
2ME: Positive (titer →1.640)
Coombs wright: Positive (titer→1.1280)
Toxoplasma- IgM: Borderline (1.1)
Toxoplasma- IgG: Negative
CMV (IgM): Negative
EBV(IgM): Negative
HIV (Ag and Ab): Negative
Hbs-Ab: Positive (vaccination has been performed)
CSF Analysis: RBC=5, WBC= 8[Polymorph: 20%, Lymph: 80%], Glucose: 47 mg/dL, Protein: 43.6 mg/dL
CSF Culture: Negative
Blood Culture: Negative
Urine Culture: Negative
ANA= Negative
C3, C4, CH50= Normal

Treatment was started by the diagnosis of neurobrucellosis with the following protocol:
1) Intravenous gentamycine (30 mg BID) for 10 days,
2) Oral rifampin (150 mg per day) for 6 weeks, 3) Oral trimethoprim-sulfimatesoxazole (200-40 mg TDS) for 6 weeks.

Fever was discontinued 3 days later and she was discharged from hospital 10 days after starting the antibiotic protocol.

In outpatient follow up visiting 7 days, six and nine months later, there was no fever recurrence or eye deviation. Physical and neurologic exams were normal. Lab findings after six- months were wright test: 1.80, 2ME: 1.40 and after nine- months were wright test: Negative, 2ME: negative.
intensities, and vascular abnormalities. Demyelination of CNS is considered as a rare manifestation of NB (5, 11, 14).

The white matter alterations in NB were a consequence of demyelination; also confirmed by pathologic study (15). This claim was bolstered with autopsy evidence that demyelination in a patient with NB resembled the lesions of multiple sclerosis (6, 16). The underlying reason for the changes in white matter is yet to be known, but they may be resulted from an autoimmune reaction (2). Not only the white matter involvement of NB resembles that of MS or Vasculitis, but also it can imitate that of other inflammatory or infectious diseases such as acute disseminated encephalomyelitis or Lyme disease (2, 17). Due to basal meningitis, involvement of one or more cranial nerves is observed in more than half of NB cases. The vestibulocochlear nerve is the cranial nerve most commonly involved in NB (3). It is very rare to see isolated cranial nerve involvement in NB; only a few isolated abducens nerve palsies (6, 18, 19) have been reported. The pathogenesis of abducens nerve palsy is hypothetical. Possible etiologies include extension of meningeal infection and possible vasculitis processes (18).

**In conclusion,** NB, as a disease, can be treated with a favorable result. The diagnosis of this disease is contingent mainly upon high clinical attention in endemic countries. It is presented diversely in clinical or radiological diagnoses, particularly in young patients with neurological abnormalities. Imaging results of NB can potentially be misleading as they are varied and can imitate properties of other demyelinating, infectious, or inflammatory conditions.

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**Authors’ contribution**
Andisheh M; Substantial web research, Follow-up interviews with the patient
Amirsalari S; Final diagnosis and treatment, Post-treatment follow-up procedures with the patient, Final approvals
Torkaman M; Pharmacologic consultations
Sabzechian M; Oncology consultations
Afsharpaiman Sh; Infectious diseases consultations
All authors agreed to be accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved.

**Conflict of interest**
No conflict of interest.

**Fig 1.** Left side abducens nerve palsy

**Fig 2.** Round hyperintense lesion at left parasagitalparieto-occipital area in brain MRI
to Brucella spp. Eur J Pediatr 2006; 165:726–727.
8. Rangel Guerra R, Martinez HR, Leon Flores L. Neurobrucellosis. Report of five cases and literature review. Rev Inwest Clin 1982;34:62–8.
9. Mugerwa RD, D’Arbela PG. Brucella meningitis; a case report and review of the literature. East African Med J 1976;53:266–9.
10. Bashir R, Al-Kawi MZ, Harder EJ, Jinkins J. Nervous system brucellosis: diagnosis and treatment. Neurology 1985; 35:1576–1581.
11. Al Deeb SM, Yaqub BA, Sharif HS, Phadge JG. Neurobrucellosis: clinical characteristics, diagnosis, and outcome. Neurology 1989; 39:498–501.
12. Lubani MM, Dudin KI, Araj GF, Manandhar DS, Rashid FY. Neurobrucellosis in children. Pediatr Infect Dis J 1989;8:79–82.
13. Santini C, Baiocchi P, Berardelli A, Venditti M, Serra P. A case of brain abscess due to Brucella melitensis. Clin Infect Dis 1994;19:977–8.
14. Koussa S, Chemaly R. Neurobrucellosis presenting with diffuse cerebral white matter lesions. Eur Neurol 2003; 50:121–123.
15. Fincham RW, Sahs AL, Joynt RJ. Protean manifestation of nervous system brucellosis. Case histories and a wide variety of clinical forms. JAMA 1963; 184:269–275
16. Marconi G. Su un caso di sclerosi multipla acuta insorta dopo un’infezione da Brucella abortus. Riv Patol Nerv Ment 1966; 87:548–565.
17. Bussone G, La Mantia L, Grazzi L, Lamperti E, Salmaggi A, Strada L. Neurobrucellosis mimicking multiple sclerosis: a case report. Eur Neurol 1989; 29:238–240.
18. KarakurumGöksel B, Yerdelen D, Karataş M, et al. Abducens nerve palsy and optic neuritis as initial manifestation in brucellosis. Scand J Infect Dis 2006; 38:721–725.
19. Yılmaz M, Ozaras R, Mert A, Ozturk R, Tabak F. Abducent nerve palsy during treatment of brucellosis. Clin Neurol Neurosurg 2003; 105:218–220.

Fig 3. Round hyperintense frontal subcortical lesion in brain MRI

References
1. Pappas G, Akritidis N, Bosilkovski M, Tsianos E. Brucellosis. N Engl J Med 2005; 352:2325–2336.
2. Al-Sous MW, Bohlega S, Al-Kawi MZ, Alwatban J, McLean DR. Neurobrucellosis: clinical and neuroimaging correlation. AJNR Am J Neuroradiol 2004; 25:395–401
3. Bingöl A, Togaylsikay C. Neurobrucellosis as an exceptional cause of transient ischemic attacks. Eur J Neurol 2006; 13:544–548.
4. Adaletli I, Albayram S, Gurses B, et al. Vasculopathic changes in the cerebral arterial system with neurobrucellosis. Am J Neuroradiol 2006; 27:384–386.
5. Shakir RA, Al-Din AS, Araj GF, Lulu AR, Mousa AR, Saadah MA. Clinical categories of neurobrucellosis. A report on 19 cases. Brain 1987; 110:213–223.
6. Esra Özkavukcu, Zeynep Tuncay, Ferda Selçuk, İlhan Erden: An unusual case of neurobrucellosis presenting with unilateral abducens nerve palsy: clinical and MRI findings; Turkish Society of Radiology 2009
7. Tena D, González-Praetorius A, LópezAlonso A, Peña JL, Pérez-Pomata MT, Bisquert J. Acute meningitis due to Brucella spp. Eur J Pediatr 2006; 165:726–727.