Chronic meningitis as the first presentation of sarcoidosis: an uncommon finding

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Abstract
We describe a 40-year-old woman presenting with headache, nausea, episodic amnesia and blurred optic disc. Brain MRI disclosed diffuse leptomeningeal enhancement. CSF analysis showed aseptic meningitis with elevated ACE level. Neurosarcoidosis was diagnosed based on granulomatosis changes on tissue biopsy.

Introduction
Sarcoidosis is a chronic granulomatous systemic disease of unknown etiology, with many different clinical presentations [1,2]. Non-caseating granuloma is the hallmark pathologic finding. Pulmonary, lymph node, ocular, and skin lesions are the most common manifestations of Sarcoidosis [3]. Nervous system can also be involved. The reported prevalence of neurological involvement varies due to case selection and diagnostic criteria, and can be as high as 5% [1,4].

Cranial neuropathies from chronic basal meningitis constitute the most common neurological manifestation of Sarcoidosis; most often the facial nerve is involved, sometimes bilaterally. Encephalopathy, seizures, mass lesions, obstructive hydrocephalus, basilar granulomatous meningitis, aseptic meningitis, myelopathy, and pituitary-hypothalamic lesions are the other manifestations of Cranial Nervous System (CNS) involvement [1,5]. In 62 to 74% of neurosarcoidosis patients, neurological symptoms are the primary manifestation of the disease [6,7]; therefore, many patients with neurosarcoidosis develop systemic symptoms only after presenting with neurological signs of the disease [2]. In this article, we describe a case of neurosarcoidosis who presented with nervous system complaints.

Case Report
A 40 year old woman was presented to Shariati hospital (TUMS), with headache, nausea, and episodic amnesia. The episodes lasted less than five minutes, and resolved after vomiting.

The patient complained of no fever, focal weakness, or sensory symptoms; but she mentioned episodic dyspnea for more than 5 years, leading to cardiac and respiratory evaluations, which failed to show any specific etiology.

She had a history of admission to other hospital, about 9 months earlier, due to headache, blurring of vision and diplopia. At that time she was diagnosed as cerebral venous sinus thrombosis and had been treated with anticoagulants.

Examination disclosed blurred optic disc margins and slight neck rigidity. Brain MRI showed diffuse leptomeningeal enhancement, with scattered white matter changes (figure1).

Cerebrospinal Fluid (CSF) examination showed lymphocytosis (WBC:37,70% lymphocyte), elevated protein (277mg/dl) and low glucose (13mg/dl); but cytological examination, stains for bacteria, mycobacteria, and fungi, VDRL, Wright, and PCR for TB, all were negative in the CSF.

CSF exam was repeated three more times, each time
with complete laboratory evaluation (including cytological examination), with similar results.

Routine hematology and biochemistry tests, liver function tests, thyroid function tests, PPD (purified protein derivative), HIV antibody, C-ANCA, P-ANCA, C3, C4, CH50, anticardiolipin and antiphospholipid antibodies, were normal/negative, as well as tumor markers in search of malignancy. Chest roentgenography, HRCT of thorax, and the result of bronchoscopy with BAL (BronchoAlveolar Lavage) were unrevealing. Serum ACE level was normal, but CSF ACE was raised (24; NL<5).

The patient was treated with IV methyl prednisolone and continued on prednisolone with the impression of probable neurosarcoïdosis.

Two weeks later, she developed another episode of dyspnea. She was re-evaluated by pulmonary specialist, and because of an abnormality in the base of her lung, an open biopsy was performed. The pathology report was consistent with Sarcoïdosis. Dyspnea was ascribed to pulmonary hypertension.

Patient’s neurological symptoms and signs improved after 2 months and after 7 months follow up, she was still free of symptoms.

**Discussion**

We describe clinical and laboratory data of a case of neurosarcoïdosis. In accordance with Zajieck’s diagnostic criteria, diagnosis of probable Sarcoïdosis was made at first; and after open lung biopsy, definite Sarcoïdosis was diagnosed [6].

In most reported series, the incidence of clinically evident neurosarcoïdosis is about 5%. Neurosarcoïdosis is often suspected in patients with known Sarcoïdosis [8]. Stern et al reported that 73% of patients with neurosarcoïdosis developed their symptoms and signs within the first two years of their systemic Sarcoïdosis [8]. In our patient, systemic (pulmonary) Sarcoïdosis was diagnosed after diagnosis of neurosarcoïdosis in the form of chronic meningitis, which presented with headache, nausea, vomiting, and episodic amnesia. Brain MRI showed diffuse meningeal enhancement, without any parenchymal involvement. Meningeal inflammation with concomitant CSF abnormalities is one of the most common manifestations of neurosarcoïdosis [4]. CSF analysis in our patient showed abnormalities compatible with previous studies in the literature [5]. Although ACE level in the CSF is not a reliable marker for CNS Sarcoïdosis; with both false negative and false positive results [6], significant elevation of CSF ACE in our patient, along with exclusion of infections and malignancies, can be in favor of neurosarcoïdosis. Diagnosis and management of neurosarcoïdosis are both challenging; Stern et al recommended biopsy proof of granulomatous inflammation from any involved (including nervous system) tissue specimens before initiation of treatment [1,5]. We started treatment by a probable diagnosis of neurosarcoïdosis, and definite diagnosis was made two weeks later by open lung biopsy.

Patient’s neurological symptoms and signs improved after 2 months and now, after 7 months of follow up, she is still free of neurological symptoms.

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References


