Extradural ganglioneuroma with T1–T2 involvement mimicking spondylodiscitis: a case report and a review of the literature

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Abstract

Ganglioneuroma (GN) is a rare benign neural tumor, usually derived from the ganglia of the sympathetic system. This report describes a 36-year-old man who presented with back pain and local tenderness that closely mimicked the clinical and imaging findings of spondylodiscitis. However, histologic examination made the diagnosis of GN. To our knowledge, this is the first report presenting the pattern of a GN as a differential diagnosis of spondylodiscitis.

KEY words: neural crest tumor, spinal cord neoplasms, ganglioneuroma, radionuclide imaging

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Background

Ganglioneuroma (GN) is a rare benign neural tumor, usually derived from the ganglia of the sympathetic system [1]. The most common anatomical sites are the mediastinum, retroperitoneum, adrenal gland and also head and neck [1]. This report describes a 36-year-old man presenting with back pain and local tenderness, which closely mimicked the clinical and imaging findings of spondylodiscitis. However, histologic examination demonstrated the diagnosis of GN.

Case report

A 36-year-old man presented with a history of progressive back pain for three months. His past medical, surgical and family history were unremarkable. He denied any other symptoms such as fever, night sweats, cough, breathing difficulty and weight loss. In physical examination, except local tenderness over upper thoracic vertebrae, no other abnormal finding was found. Neurologic examination was also normal. Laboratory data revealed hemoglobin 10 g/dL, hematocrit 27.8%, mean corpuscular volume (MCV) 75.54 µm³, erythrocyte sedimentation rate (ESR) 92 mm/h and CRP 2+. The patient subsequently underwent magnetic resonance imaging (MRI) of thorax, which revealed signal changes of T1, T2 bodies, erosion of the endplates with invasion into the intervertebral disc space and a soft tissue mass, bulged anteriorly. The canal was preserved. For three phasic bone scans, immediately following injection of 20 mCi ⁹⁹ᵐTcMDP, bone scan was obtained in perfusion and blood pool phases from thoracic vertebrae. Delayed whole-body bone scan with spot views from that region was also performed. The finding was regional hyperemia and increased radiotracer uptake of T1–T2 vertebrae (Figure 1). Remainder of the skeleton was unremarkable.

Altogether, the clinical history, laboratory data and imaging findings were suggestive of spondylodiscitis, most probably spinal tuberculosis [2], which is not an uncommon disease in our country [3, 4]. The patient underwent open biopsy which histologic analysis of specimen derived from the spinal mass demonstrated mature ganglion cells, with abundant cytoplasm, large nuclei and prominent nucleoli, distributed within an abundant stroma, compatible with ganglioneuroma (Figure 2).

Discussion

Sympathetic trunks and their associated ganglia, located on the side of vertebral bodies, can be the origin of sympathetic ganglion tumors. These include a spectrum from aggressively malignant neuroblastoma (NB), to malignant ganglioneuroblastoma (GNB) and to purely benign GN. Gangliocytoma is a rare and benign tumor arising from ganglia-type cells, which are groups of nerve cells. This tumor arises from ganglia. When neuronal elements make up the majority
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of the mass, the tumor is referred to as a ganglioneuroma. In contrast, gangliocytomas are composed of mature ganglion cells and therefore have no potential for malignant change. Gangliocytoma is a surgically curable disease and surgical resection performs in symptomatic patients. Overall, prognosis is good and recurrence rate is about 25 percent.

Peripheral GN is a well differentiated, benign and slowly growing tumor. It can arise anywhere from the base of the skull to the pelvis, but its most common location is the posterior mediastinum (41.5%), followed by retroperitoneum (37.5%), adrenal gland (21%) and neck (8%) [1, 5]. Spinal GN usually involves the paraspinal region and can grow into the spinal canal through the neural foramina [6]. Depending on the location of the tumor, spinal GN may cause various symptoms, including scoliosis, peripheral radiculopathy, paraparesis or gait disturbance [6, 7]. Although, GN is a benign well- encapsulated tumor and typically does not invade adjacent structures, it has been reported that it can attach to the vertebral bodies [8], or involve the spinal column [6, 9–11]. However, in our patient a unique pattern of bone involvement was noted, mimicking spondylodiscitis. In addition, in most of reported cases, scintigraphic appearance of bone involvement has not been mentioned. In our knowledge, this is the first report presenting the pattern of a GN as a differential diagnosis of spondylodiscitis.

The diagnostic value of metaiodobenzylguanidine (MIBG) scintigraphy in neural crest / neuroblast-derived tumors such as neuroblastoma and pheochromocytoma is established [12, 13].

It should be mentioned that main differential diagnoses of such pattern in the thoracic spine on bone scan are inflammatory process and tumoral involvement, for which radiolabeled WBC’s and gallium scanning in addition to radiological modalities, especially MRI, are helpful to reach the diagnosis of inflammation [14].

References