Primary Extraskeletal Ewing Sarcoma of the Thoracolumbar Epidural Space: Rare Case Report in a Child

Sarcoma de Ewing extraesquelético primário no espaço epidural toracolombar: Raro relato de caso em criança

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Abstract

Introduction Ewing sarcomas are a family of tumors that can be of skeletal or extraskeletal origin. We report a rare case of a child with extraskeletal Ewing sarcoma in the thoracolumbar epidural space.

Case Report The patient was a 1-year-old female child with sphincter alteration, flaccid paraplegia, and areflexia. A magnetic resonance imaging (MRI) scan showed a large extensive epidural lesion with compression of the dural sac in the D6–L2 segment, and a left paravertebral extension through the L1–L2 foramen. Laminotomy was performed, with subtotal resection of the lesion. The histopathological and immunohistochemical analyses indicated Ewing sarcoma. Due to the child’s age, radiotherapy was not performed, only chemotherapy, due to the aggressiveness of the neoplasm. The patient showed rapid tumor recurrence and ended up dying.

Discussion Extraskeletal Ewing sarcoma can appear in different locations in the body. They are aggressive tumors with local recurrence and distant metastases. In our case, a combination of MRI and positron-emission tomography–computed tomography scan presented a clearer result, especially in the presence of metastasis. In the histopathological analysis, small blue cells with a clear cytoplasm and indistinct nucleoli were observed. In the immunohistochemical analysis, CD99 (MIC2) expression is
Introduction

Ewing sarcoma is a rare malignant neoplasm that was first described by James Ewing in 1921. He described it as a small-blue-round-cell tumor. Initially, Ewing sarcoma was believed to originate in the undifferentiated endothelial and mesenchymal cells of the bone marrow, but, with the advent of immunohistochemistry and cytogenetic tests, its neuroectodermal origin was discovered.1-3 It is the second most common bone tumor in children and adolescents.4

The Ewing sarcoma family of tumors (ESFT) includes peripheral primitive neuroectodermal tumors (pPNETs) and Askin tumors. They are morphologically similar in terms of malignancy, and they can be of skeletal or extraskeletal origin. Extraskeletal Ewing sarcomas account for 6% to 47% of all tumors in the ESFT.5,6

The patient in the case herein reported was a child with a rare extraskeletal Ewing sarcoma of the thoracolumbar epidural space.

Case Report

A 1-year-old female child presented to the hospital with difficulty in walking, with motor deficit progressively worsening and dorsalgia for 30 days. Upon physical examination, she presented with intestinal constipation, bladder dysfunction (neurogenic bladder), flaccid paraplegia, and areflexia. The exame de ressonância magnética (RM) revelou volumosa lesão expansiva epidural com compressão do saco dural no segmento de D6-L2, e extensão paravertebral esquerda através do forame L1-L2. Realizou-se laminotomia, com ressecção subtotal da lesão. O exame histopatológico e a imunohistoquímica indicavam sarcoma de Ewing. Por conta da idade da criança, ela não foi submetida a radioterapia, apenas a quimioterapia, e, devido à agressividade da neoplasia, a paciente evoluiu com rápida recidiva tumoral e terminou falecendo.

Discussion

The thoracolumbar epidural Ewing sarcoma can appear in different localities of the body. They are aggressive tumors with local recurrence and distant metastasis. In the investigation, the combination of RM and tomography of positron emission tomography–computed tomography presents better results, especially in the presence of metastasis. In the histopathological exam, we observed small blue cells with clear cytoplasm and indistinct nucleoli. In the immunohistochemical exam, the main expression is CD99 (MIC2). The best treatment outcome would have been surgical resection with chemotherapy and radiotherapy.

Conclusion

We reported a rare case of thoracolumbar epidural Ewing sarcoma in which, despite surgery and chemotherapy, the tumor behaved very aggressively, leading to an unfavorable prognosis.
positive/pan-cytokeratin positive/CD99 positive/friend leukemia integration-1 (FLI-1) positive.

Due to the patient’s age, radiotherapy was not performed, only chemotherapy. The patient then presented with rapid lesion recurrence (Fig. 3), with tumor lysis, septic shock and, after several days in the intensive care unit (ICU), she died.

Discussion

In 1969, Tefft et al. first described four patients with paravertebral soft-tissue tumors that were histologically similar to Ewing sarcoma. In 1975, Angervall and Enzinger reviewed 39 patients with paravertebral malignant soft-tissue tumors that did not originate in the bone, but were morphologically similar to skeletal Ewing sarcoma.

Extraskeletal Ewing sarcoma is part of the ESFT. This group of tumors affects bones and soft tissues. The peak incidence is between 10 and 15 years of age, with a specific translocation t(11; 22)(q24; q12) in > 90% of the cases.

Extraskeletal Ewing sarcoma can appear in different locations of the body, such as the central nervous system, the chest wall, the retroperitoneum, the skin, the kidneys, the small intestine, the pelvis, the rectum, the vagina, the fingers, the arms, the scalp, the lips, the nasal passages, the paravertebral region, and the perineum. They are aggressive tumors with a high incidence of local recurrence and distant metastasis, mainly to the lungs, spine, and brain.
tumors have worse prognosis compared with that of other bone tumors of the ESFT.²,⁶

Extradural tumors account for 30% of all spinal-cord tumors in children. Therefore, the differential diagnoses may be: benign bone tumors (such as osteoid osteoma, osteochondroma, giant-cell tumor, aneurysmal bone cyst, and hemangioma), Langerhans cell histiocytosis, and fibrous dysplasia; and malignant tumors, such as sarcomas, teratomas, chordomas, and metastatic lesions.¹¹

For the diagnosis, imaging techniques like CT be used to look for heterogeneous masses, and the MRI, for hypo- or isointense signals in T1-weighted images and hyperintense signals in T2-weighted images. Fluorodeoxyglucose positron emission tomography has not shown good results in the diagnosis and staging of soft-tissue sarcomas.²,¹²

A combination of MRI and positron-emission tomography–computed tomography (PET-CT) has been shown to yield satisfactory results, mainly in the identification of metastasis.²,¹²

As clinical findings are inaccurate and the diagnosis of extraskeletal Ewing sarcomas by imaging is nonspecific, the histopathological analysis becomes vital, and, on it, uniform proliferation of small blue round cells with a clear cytoplasm and indistinct nuclei is observed. The immunohistochemical markers traditionally used in the differential diagnosis of ESFT are CD99 (MIC2), FLI-1, and human natural killer-1 (HNK-1). Other more specific markers are also used, such as enolase, S-100, chromogranin A, synaptophysin, cytokeratin, and EMA. The expression of Ki67 also represents an indicator of poor prognosis in ESFT.²,⁸,¹³

In terms of treatment, patients with extraskeletal Ewing sarcoma who underwent surgical resection (partial or total), chemotherapy, and radiotherapy had a 1-year survival rate compared with those who underwent surgery, chemotherapy, or radiotherapy.¹⁴ Patients with metastasis had an even worse prognosis.⁶

In the present article, we reported a rare case of thoracolumbar epidural Ewing sarcoma, and due to the atypical age group (1 year of age) for the pathological condition, in which radiotherapy is not recommended, the tumor proved to be extremely aggressive. Despite surgery and chemotherapy, the tumor presented a rapid degree of recurrence, leading to an unfavorable outcome.

Conflict of Interests
The authors have no conflict of interests to declare.

References