Extraskelletal Ewing’s Sarcoma of the Thoracic Nerve Root: A Case Report

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Extraskelletal Ewing’s sarcoma (EES) is a rare malignant soft tissue tumor which is morphologically indistinguishable from skeletal ES. EES usually occurs in young adults and children and there has been only one case reported in a patient aged over 70 years old. We report a case of an EES arising from the first thoracic spinal nerve root in a 73-year-old female, which was misdiagnosed as benign nerve sheath tumor in preoperative imaging evaluation.

Index terms Sarcoma, Ewing; Spinal Nerve Root; Thoracic Nerves; Soft Tissue Neoplasm

INTRODUCTION

Extraskelletal Ewing’s sarcoma (EES), especially involving spine accounts for only 5% of all ES (1). It is known to be found mainly in the paravertebral and epidural space (2). And ES usually occurs in children or young adults (1). Primary EES of spinal nerve has been reported only a few cases and there has been only one case of a senior patient in eighth decade (3).

Herein, we report a case of spinal epidural EES in a 73-year-old female.

CASE REPORT

A 73-year-old female visited to our hospital complaining pain in the posterior neck and left arm. The symptoms started 6 months ago and aggravated 2 days ago. Physical examination revealed hypoesthesia on her left ulnar nerve territory without decrease of muscle strength. She underwent plain radiography and CT of cervical spine. Plain ra-
diographs of the cervical spine showed no significant abnormality other than disc space narrowing in C5–C6 level (not shown).

CT revealed a dumbbell-shaped mass in the left neural foramen of T1–T2. No calcification was seen in the mass. There was mild widening of left neural foramen (Fig. 1A). For further evaluation, MRI of the cervical and thoracic spine was performed. MR images showed that the medial part of the mass is in the intradural and extramedullary region and the lateral part of the mass extends to the paravertebral area through the left neural foramen. The lesion shows intermediate signal intensity on T1-weight images and homogenously slightly high signal intensity on T2-weighted images (Fig. 1B). After administration of intravenous contrast material, the mass is enhanced homogeneously. Size of the lesion is measured approximately 2.6 × 1.3 × 1.5 cm. The mass is branching outside the left neural foramen and extending along the first branch of left T1 nerve (Fig. 1B).

Based on the CT and MRI findings, the benign nerve sheath tumor was suggested. And then, the patient underwent surgery. In the operation, the tumor was revealed after laminectomy of T1 vertebra and also after incision of dura (Fig. 1C). The mass was strongly attached to the nerve root, partly intermingled with the nerve fibers. Subtotal resection of the tumor was done with motor evoked potential monitoring to minimize injury of the nerve.

Microscopic examination revealed that the tumor was composed of alternating highly cel-

![Fig. 1. Extraskeletal Ewing’s Sarcoma in a 73-year-old female.](https://doi.org/10.3348/jksr.2019.80.3.568)

A. Thoracic spine CT with contrast enhancement shows a dumbell-shaped mass in the T1–T2 left neural foramen extending inside the central canal and outside the neural foramen, with mild homogeneous enhancement (arrows). Mild widening of the neural foramen is noted.

B. On MRI with contrast enhancement, the lesion shows intermediate signal intensity on the axial T1WI and slightly high signal intensity on the T2WI. Homogeneous enhancement of the lesion is seen on the FS-T1WI after administration of Gd-CE. The FS-T2WI shows the tumor extending along the first branch of left T2 nerve root (arrowheads).

Gd-CE = gadolinium contrast-enhanced, FS = fat saturation, T1WI = T1-weighted image, T2WI = T2-weighted image
Extraskeletal Ewing’s Sarcoma

lular and paucicellular components with dense collagen bands (× 10). The high-power field view showed highly overlapping blue round cells with oval nuclei and scanty cytoplasm, compatible with blue round cell tumor. No evidence of neuroectodermal differentiation, such as rosette formation was identified (× 400) (Fig. 1D). The result of immunohistochemical staining revealed weak positivity for vimentin and CD56, and strong membranous staining for CD99, compatible with ES (Fig. 1D). Further gene mutation study revealed EWSR1 gene translocation and the final pathological diagnosis was made as EES.

After confirmative diagnosis, the patient transferred to cancer clinic for adjuvant chemotherapy and radiation therapy. After twelve months from the surgery, the tumor was reduced

Fig. 1. Extraskeletal Ewing’s Sarcoma in a 73-year-old female.
C. Intraoperative microscopic photograph. The tumor is located in the intradural portion of T1–T2 (asterisk). The T1 nerve root is seen on the left side (arrowhead).
D. Left photomicrograph of the specimen shows alternating (highly cellular and paucicellular) cellularity with dense collagen bands. The right high-power view shows highly overlapping blue round cells with oval nuclei and scanty cytoplasm. Neuroectodermal differentiation such as rosette formation was not observed (H&E stain × 10; left, H&E stain × 400; right). Immunohistochemical staining result reveals weak positivity for vimentin, CD56, and strongly membranous staining for CD99 (H&E stain × 400).
H&E = hematoxylin and eosin
in size and the patient was in stable condition without local recurrence, only complaining mild tingling sensation on her elbow. However, recently, after post-operative sixteen months, local recurrence and leptomeningeal seeding developed and the patient is receiving chemotherapy.

**DISCUSSION**

EES is a rare form of round cell malignant tumor with extraskeletal origin (1, 2, 4). It accounts for about 10% of ES in children and 5% of ES in adults (3). The primary site of EES involving extradural and intradural extramedullary space is extremely rare (1-10). Preferred anatomic origin of EES is paravertebral and epidural space, so associated symptoms are variable such as local pain, gait disturbance, motor deficit, radicular pain, etc (1, 4, 5).

Imaging features of EES are nonspecific but, usually appears solid, iso signal intensity on T1-weighted image, intermediate to high signal intensity on T2-weighted image and moderate to strong enhancement after gadolinium administration (1-10). Some reported lesions show slightly inhomogeneous signal intensity or low signal intensity on T2-weighted images (1, 3-5) and heterogeneous enhancement (1, 3, 5, 6).

In our case, the mass was iso signal intensity in T1-weighted image and slightly high signal intensity in T2-weighted image with marked enhancement, similar to previous cases. The radiologic differential diagnosis of EES involving epidural space includes benign nerve sheath tumor such as schwannoma, lymphoma and malignant nerve sheath tumor (2-4, 7). Schwannoma has smooth margin and dumbell shaped with intense enhancement. On T2-weighted image, schwannoma appears heterogeneously high signal intensiti with target sign (3, 4, 7). Lymphoma involving epidural space originates from body of vertebra or paravertebral lymph node then extends to epidural space. It is slightly high signal intensity to muscle on T1-weighted image and T2-weighted image involving multi-segment of spine with homogeneous enhancement along epidural infiltration (9). Malignant nerve sheath tumor of spine appears expansile soft tissue tumor destructing adjacent vertebra with paraspinal and epidural extension. On MR, the mass appears low signal intensity on T1-weighted image and high signal intensity on T2-weighted image. It is mostly associated with neurofibromatosis type 1 (10).

This case was initially presumed to be a benign nerve sheath tumor, however the final diagnosis was an EES. There was no definite aggressive feature such as ill-defined margin, adjacent bony destruction, heterogeneous signal intensity or attenuation of the mass, or large size of the mass. However, in the literature, there are some reported cases of EES mimicking benign nerve sheath tumors (4, 6, 7) and it was difficult to differentiate EES from benign nerve sheath tumor only with radiologic imaging, without pathologic confirmation in those cases. In our case, the lesion extended to the 1st branch of thoracic nerve root and this feature may indicate relatively more aggressive kinds of tumor such as lymphoma or malignant peripheral nerve sheath tumor rather than benign nerve sheath tumor, even though ES is very rare in elder patients. However, this finding may be seen in neurofibromas.

Therefore, even there is well-marginated dumbbell-shaped extradural mass which looks benign, if there is subtle ill-defined enhancing or high signal intensity lesion extended to peripheral nerve, possibility of malignancy might be considered.
We reported a case of EES arising from thoracic nerve root in 73-year-old woman. Because of its rarity and varying morphology on imaging modality, it is difficult to diagnosis from radiologic finding before pathologic confirmation. We suggest that in case of nerve root tumor showing atypical radiologic findings of nerve sheath tumors, EES also could be considered as differential diagnoses although these tumors are very rare in elder patients.

Conflicts of Interest
The authors have no potential conflicts of interest to disclose.

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흉부 신경근에서 발생한 골외성 유잉육종: 증례 보고
김정원1 · 이지혜1* · 김재형1 · 정명자1 · 김성희1 · 김지영1 · 김수현1 · 강미진1 · 김태규1 · 배경은1 · 신준재2 · 김현정3 · 김정연3
골외성 유잉육종은 드문 악성 종양으로 육안적으로는 골성 유잉육종과 감별할 수 없다. 골외성 유잉육종은 대부분 젊은 성인이나 20세 이하의 소아청소년에서 발생하며 70세 이상에서 발생한 경우는 현재까지 1예가 보고되었다. 우리는 양성 신경초종양으로 오인되었던 73세 여성의 첫 번째 황부 신경근에서 기원한 골외성 유잉육종의 증례를 보고하고자 한다.
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