Case Report

Thoracic extradural paragangliomas: a case report and review of the literature

P Conti\textsuperscript{1,*}, H Mouchaty\textsuperscript{1}, B Spacca\textsuperscript{1}, AM Buccoliero\textsuperscript{2} and R Conti\textsuperscript{1}

\textsuperscript{1}Department of Neurosurgery, University of Florence, Florence, Italy; \textsuperscript{2}Department of Human Pathology and Oncology, University of Florence, Florence, Italy

Study design: Case report.
Objectives: To report on a case of paraganglioma presenting in an uncommon extradural thoracic localization.
Setting: Department of Neurosurgery, Florence, Italy.
Case report: A 43-year-old woman with a thoracic lesion extending into the extradural space along four levels, T\textsubscript{1}–T\textsubscript{4}, presented with sudden spastic incomplete paraplegia and paresthesia at the lower limbs.
Results: The neoplasm was surgically resected 'en bloc' and histological findings corresponded to paraganglioma. One year after surgery, the patient was walking without assistance, a T\textsubscript{2}–T\textsubscript{4} hypoesthesia was still present and an magnetic resonance imaging (MRI) study showed no signs of focal recurrence.
Conclusions: The imaging features of thoracic paragangliomas may be misleading and an advanced malignant lesion could be primarily suspected; thus, a histological study is always needed. Total resection is the gold standard therapy. Owing to the risk of recurrence or multicentric growth, follow-up must be prolonged and accurate.

Keywords: paraganglioma; spinal neoplasm; extra-adrenal

Introduction

Paragangliomas are tumors originating from the sympathetic or parasympathetic autonomic nervous system associated to the paraganglia, most frequently found in the jugular glomus and the carotid bodies (90\% of all paragangliomas). There is no precise data in the literature about prevalence of spinal paragangliomas. Most cases describe the lesions as extramedullary and intradural, most frequently sited at the level of the cauda equina and the filum terminale.\textsuperscript{1–6} Findings in the extradural space at a cervical or dorsal spinal level are particularly rare.\textsuperscript{1,3,4,7–15}

We describe the case of a thoracic extradural paraganglioma invading the thoracic cavity. A revision of the literature is reported.

Case report

During alopecia treatment with steroids, a 43-year-old woman suffered acute pain involving the interscapular region, the shoulder and the right arm with paresthesia and weakness in the legs and was transferred to our department. At examination, incomplete spastic paraplegia, hypoesthesia in T\textsubscript{3} and T\textsubscript{4} dermatomes bilaterally with a Lhermitte sign, and incomplete ptosis with a slight miosis in the right eye were found. Blood and urine chemical and physical tests were normal, as were blood pressure values.

Computed tomography (CT) scans showed an almost spherical, 3.13 cm mean diameter, mediastinic mass that greatly enlarged the right vertebral foramina at the T\textsubscript{2} and T\textsubscript{3} level (Figure 1). Magnetic resonance imaging (MRI) revealed the lesion as hypo-isointense in T\textsubscript{1} and iso-hyper-intense in T\textsubscript{2} while contrast enhancement was intense and homogeneous. The lesion distorted the medulla and surrounded the right nerve roots at these levels (Figure 2).

Although the imaging characteristics were not of clear infiltration but rather of expansion (dislocated dural theca and vertebral osteolysis with hyperdense margins), the diagnosis of a primitive lung tumor invading the vertebrae was advanced.

Abdomen and thorax contrast-CT scans and bone scintigraphy excluded the presence of other lesions.

*Correspondence: P Conti, Department of Neurosurgery, University of Florence, L.go P. Palagi 1, 50139 Florence, Italy
Owing to the equivocal interpretation of the imaging study and considering the neurological conditions, despite visceral involvement, it was decided to treat the lesion surgically. In the prone position, a median incision centered on T2 and arcuately prolonged under the right scapula was performed. Thoracotomy was achieved and a voluminous extrapleural mass, adherent to the T2-T4 vertebral bodies and to the 2nd, 3rd and 4th right ribs, was visible. The macroscopic aspect was an irregularly ovoid, variegated rosy–reddish lesion with duro-elastic consistency, which surrounded two consecutive nerve roots (right T3 and T4). After having incised the affected ribs, a right T2–T3 lamino-arthro-transversectomy was performed, and the lesion was easily detached. A partial T2 somatectomy allowed 'en bloc' removal of the lesion including its thoracic part.

An expansible cylinder (ADD, Anterior Distraction Device, Ulrich, Germany) was introduced to replace the removed body and one longitudinal bar fixed with hooks on the C7 and T4 right laminae was inserted (Figure 3).

The surgical specimen was fixed in formalin and embedded in paraffin. 5 μm sections were stained with hematoxylin–eosin or mounted on electrostatic slides for immunohistochemical study. Light microscopy revealed...
polygonal cells with a pink-stained granular cytoplasm and rounded, occasionally pleomorphic, nuclei showing fine chromatin and inconspicuous nucleoli in a prominent stromal and vascular hyalinization. Tumoral cells formed lobules and cords often obscured by the hyalinization phenomena. Mitoses were not seen. Immunohistochemically, the lesion was SP, NF, NSE, S-100 protein, and CHROMO A positive. No tumoral labeling was observed for VIM, CK, and EMA. The proliferation index, estimating the percentage of Ki-67-positive cells was low (1%) (Figure 4).

Postsurgically, the patient presented T3–T4 hypoesthesia as well as progressive resolution of the hyperreflexia and the ability to walk without assistance. A slight right Horner Syndrome persisted.

At 1 year from surgery, the area of hypoesthesia shrank and only a slight drooping of the upper eyelid persisted. No residual or recurrent lesion was detected on MRI.

**Discussion**

Extraadrenal (10–15%) and adrenal (85–90%) paragangliomas associated with the sympathetic nervous system (always chromaffine), produce catecholamine in 50% of the cases but rarely release it and thus are rarely symptomatic (weight loss, hypertension, flushing, sweating, tremors, tachycardia, nausea, vomiting, etc). Parasympathetic autonomic nervous system paragangliomas are nonchromaffine and rarely produce catecholamine.5,16

Spinal paragangliomas are rare and there is no agreement on their origin since there are no paragangliomas in this region (sympathetic autonomous medullar
Table 1  Thoracic extradural paraganglioma: review of the literature

<table>
<thead>
<tr>
<th>No</th>
<th>Year</th>
<th>Author</th>
<th>Sex</th>
<th>Age</th>
<th>Site</th>
<th>Preoperative symptoms</th>
<th>Surgical treatment</th>
<th>Follow-up</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1983</td>
<td>Boker et al</td>
<td>F</td>
<td>35</td>
<td>T&lt;sub&gt;e&lt;/sub&gt;-T&lt;sub&gt;5&lt;/sub&gt;</td>
<td>2 months back pain 4 weeks paraparesis and T&lt;sub&gt;4&lt;/sub&gt; sensory loss</td>
<td>Total resection</td>
<td>Incomplete recovery after surgery. 8 months: tumor recurrence, partially removed. 13 months: death</td>
</tr>
<tr>
<td>2</td>
<td>1983</td>
<td>Boker et al</td>
<td>F</td>
<td>36</td>
<td>T&lt;sub&gt;11&lt;/sub&gt;</td>
<td>10 months back pain  Sudden paraparesis and sensory loss below T&lt;sub&gt;11&lt;/sub&gt;</td>
<td>Incomplete resection</td>
<td>3 years: back pain</td>
</tr>
<tr>
<td>3</td>
<td>1991</td>
<td>Cybulski et al</td>
<td>M</td>
<td>34</td>
<td>T&lt;sub&gt;8&lt;/sub&gt;</td>
<td>4 months midthoracic back pain  Sudden paresthesia and stiffness at lower limbs</td>
<td>Stage 1: incomplete resection  Stage 2: embolization  Stage 3: total resection</td>
<td>13 months: neurologically intact  Later: abdominal paraganglioma was detected</td>
</tr>
<tr>
<td>4</td>
<td>1996</td>
<td>Fitzgerald et al</td>
<td>M</td>
<td>30</td>
<td>T&lt;sub&gt;4&lt;/sub&gt;</td>
<td>Back pain  Numbness below T&lt;sub&gt;3&lt;/sub&gt;  Sustained ankle clonus in both legs</td>
<td>Total resection</td>
<td>18 months: neurologically intact  18 months: MRI showed a tumor recurrence at T&lt;sub&gt;5&lt;/sub&gt;</td>
</tr>
<tr>
<td>5</td>
<td>1996</td>
<td>Noorda et al</td>
<td>F</td>
<td>52</td>
<td>T&lt;sub&gt;7&lt;/sub&gt;-T&lt;sub&gt;9&lt;/sub&gt;</td>
<td>2 years neuralgia in the thoracic spine and the right thoracic wall  Lower extremities weakness, T&lt;sub&gt;3&lt;/sub&gt; local tenderness and decreased sensibility</td>
<td>Stage 1: embolization  Stage 2: total resection  Stage 3: anterior stabilization</td>
<td>After surgery: pneumonia, pericarditis and perforation of a rod through the skin 4 months: implants removal. 1 year: neurologically intact</td>
</tr>
<tr>
<td>6</td>
<td>2001</td>
<td>Shin et al</td>
<td>M</td>
<td>43</td>
<td>T&lt;sub&gt;6&lt;/sub&gt;</td>
<td>1 year recurrent back pain  Sudden paresthesia of the lower extremities  Sudden urinary incontinence</td>
<td>Stage 1: embolization  Stage 2: resection</td>
<td>6 months: headache  6 months: on MRI, 2 cerebral metastases  Radio- and chemo-therapy were not satisfactory</td>
</tr>
<tr>
<td>7</td>
<td>2001</td>
<td>Shin et al</td>
<td>F</td>
<td>67</td>
<td>T&lt;sub&gt;11&lt;/sub&gt;</td>
<td>Back pain for several years  Lower abdominal pain for 1 year</td>
<td>Incomplete resection</td>
<td>Not mentioned</td>
</tr>
<tr>
<td>8</td>
<td>2002</td>
<td>Houten et al</td>
<td>M</td>
<td>41</td>
<td>T&lt;sub&gt;9&lt;/sub&gt;</td>
<td>2 weeks gait disturbance and proximal lower limbs extremity weakness  Increasing sensation of numbness  Diminished sensation to pinprick under T&lt;sub&gt;10&lt;/sub&gt;  Low limbs hyperreflexia (small enhancing liver lesions detected on CT scans)</td>
<td>Incomplete resection  Radio- and chemo-therapy</td>
<td>1 year: Neurologically intact  1 year: on MRI no growth of spinal and liver lesions</td>
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</table>
nerves or heterotopic cells). Their most common localization is intradural extramedullar in the lumbosacral tract. We found only eight cases of thoracic primitive nonseccentrum extradural paragangliomas in the literature and only two cases of thoracic extradural seccentrum ones(9-13,15) (Table 1). One of these latter cases, described by Hamilton and Tait, corresponded to a metastasis. The other, described by Graham et al., had anamnesis that included surgical removal of a paraganglioma in a different localization, leading to the hypothesis that the thoracic lesion was probably a metastatic dissemination or a multicentric growth. A similar observation was made in a case reported by Boker et al. For these reasons, these cases of thoracic extradural paraganglioma were excluded from among the primitive ones.

**Differential diagnosis**

The diagnosis of paraganglioma cannot be based on imaging features. On CT scans, in fact, such lesions appear as homogeneous masses, sometimes with calcifications, and they present homogeneous enhancement due to their rich vascularization. Areas of osteolysis or even pathological fractures and/or enlargement of the vertebral foraminae are often observed. Osteoclasts have rarely been described and may also be found in the adjacent costs.

On MRI, paragangliomas appear hypo-isointense on T1 and iso-hyper-intense, often with a 'salt & pepper' appearance on T2-weighted images. They present an intense homogeneous gadolinium-enhancement. They may be distinguished from more malign lesions on MRI when thecal compression and dislocation are present instead of infiltration.

The most specific diagnostic technique is mIBG (m-iodobenzylguanidine) scintigraphy, even if its use has been limited to seccentrum lesions or for the research of synchronous or metachronous metastasis. Further to imaging features leading to paraganglioma, mIBG scintigraphy may be another confirmation. Yet, definitive diagnosis is possible only by means of the histological exam.

The differential diagnosis of paragangliomas includes ganglio-neural tumors (schwannomas, ganglioneuromas), secondary lesions, and all the mediastinal tumors, for extradural thoracic lesions. Compared to mielomas or metastasis, paraganglioma vascularization is not only rich but also homogeneous. Hemangiomas are differentiated due to cortical expansion and a trabecular 'honeycomb' aspect. Angiomielopomas present fat cumulus.

Macroscopically, paragangliomas are characterized by a rosy-red to brown color, seldom hemorrhagic. Microscopically, the principal Type I cells (polygonal, abundant eosinophilic cytoplasm, sometimes presenting granules grouped in nests, so-called 'zellballen') are found surrounded by sustaining Type II cells. Mitosis is rare and the pleomorphism is limited. A narrow and rich vascular net is often found. Degenerative, calcific, cystic, hemorrhagic, and/or necrotic areas are also sometimes found at times. Immunohistochemical colorations are essential for a secure diagnosis. The most important neuroendocrin markers are chromogranine and sinaptofisine that identify Type I cells and S-100 protein that identifies Type II cells.

To reduce the risk of recurrence, radical surgical removal is mandatory. Radiotherapy and chemotherapy have minor efficacy and are often used palliatively in cases of aggressive or multicentric paragangliomas or when the patient conditions cannot tolerate surgery. Best results were obtained with m-131-mIBG chemotherapy: interruption of disease progression, clinical improvement and less captation on scintigraphy.

Post-surgical radiotherapy is also used in cases of recurrence or after resection of metastasis. The most frequently described surgical approaches in the treatment of mediastinal lesions invading the vertebral bodies are anterior or double approaches (anterior and posterior) in one or two sessions.

In the presented case, the surgical treatment was performed in one session, with the patient prone. The posterolateral approach allowed complete removal of the lesion with anterior and posterior monolateral stabilization.

**Conclusions**

Thoracic epidural paragangliomas are rare but have to be considered in the differential diagnosis of thoracic/mediastinal lesions since they present a better postsurgical prognosis with respect to malignant lesions, more frequently encountered in this area.

Imaging features, typical of benign lesions, are good primary elements of differentiation but histological examination is necessary for a definitive diagnosis.

Surgical removal is the gold standard treatment and has to be total, possibly 'en bloc'.

**References**