Extramedullary plasmacytoma presenting as a mediastinal mass

Mediastende kitle ile beliren ilik dışı (ekstramedüller) plazmositom

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Abstract

Extramedullary plasmacytoma (EMP) is a plasma cell neoplasm without bone marrow involvement or other systemic characteristics of multiple myeloma. Few large-scale clinical studies have been conducted because of the rarity of EMP, especially when it arises from the mediastinum. Herein we report a rare case of solitary mediastinal plasmacytoma with reactive pleural effusion. A 58-year-old female presented with grade 4 dyspnea and dysphagia, with a mediastinal mass observed with on PET. CT-guided biopsy results were suggestive of IgG kappa-type EMP arising from the anterior mediastinum. The patient was treated with local radiotherapy to the mediastinum, and had clinical and radiological response s were good. Radiotherapy is an effective treatment for mediastinal EMP, but a complete workup is mandatory, including PET, as the majority of such masses coexist with multiple myeloma.

Keywords: Extramedullary, plasmacytoma, mediastinal, radiotherapy

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Özet

Ekstramedüller plazmasitom (EMP) kemik iliği tutulumu veya multipl miyelomun diğer sistemik özelliklerinin bulunmadığı bir plazma hücresi neoplazisidir. Özellikle mediasten kökenli EMP seyrek görüldüğünden büyük ölçekli klinik çalışma sayısı oldukça azdır. Burada reaktif plevara sıvısı ile birlikte mediasten plazmasitomun görülüğünü nadir bir olgu bildirilmektedir. Derece 4 dispne ve disfajiden yakın 58 yaşındaki bir kadında PET incelemesi mediasten kitlenin ortaya çıkarmıştır. CT-yönlendirmeli biyopsi bulguları ön mediasteniden kaynaklanan IgG kappa-tip EMP tanısını koydurmıştır. Mediastene lokal radyoterapi uygulanarak iyi klinik ve radyolojik yanıt elde edilmiştir. Radyoterapi mediasten kökenli EMP için etkin bir tedavi olmakla birlikte, bu kitelerin büyük bir bölümü multipl miyelomla birlikte gelistigiinden PET’i de içeren tam bir klinik çalışma yapmak zorunludur.

Anahtar kelimeler: Ekstramedüller plazmasitom, plazmasitom, mediastinal, radyoterapi

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Introduction

Extramedullary plasmacytoma (EMP) is a plasma cell neoplasm involving soft tissues without any signs of systemic spread. It occurs in <5% of cases and mediastinal presentation is a rare presentation. Herein we report a rare case of solitary mediastinal plasmacytoma and a review of the literature.

Case presentation

A 58-year-old female presented with upper chest swelling associated with grade 4 dyspnea and difficulty swallowing. Physical examination showed a 4×3-cm supraclavicular mass with diffuse borders and a 5×5-cm infraclavicular bulge fixed to underlying structures. Her breathing sounds were slightly coarse, without rales or rhonchi on auscultation. She was treated in the intensive care unit due to her inability to maintain oxygen saturation. PET CT showed an anterior mediastinal mass (10.3×8.2 cm) extending up to the left supraclavicular region (Figure 1). The mass was abutting the major mediastinal vessels, overlying the manubrium sterni and chest wall, and was associated with bilateral pleural effusion. Cranial, spinal, and pelvic X-ray showed that there weren’t any osteolytic regions. Light microscopic examination showed scattered oval cells with eccentric nuclei and abundant eosinophilic cytoplasm suggestive of plasmacytoma. The plasma cells stained positive with LCA, CD138, and reduced kappa light-chain concentrations (Figure 2). Bone marrow biopsy results showed normal patterns of cell distribution. Pleural fluid cytological examination showed lymphocytes and reactive mesothelial cells. Serum calcium was normal and urine was negative for Bence Jones proteins. Serum electrophoresis showed an M spike in the gamma globulin (IgG kappa) region with a value of 3.07 g/dL. Based on these findings, the patient was diagnosed as mediastinal plasmacytoma. The patient received 3-dimensional conformal external beam radiotherapy (3D-CRT) to the mediastinum (46.2 Gy in 26 fractions). Gross target volume GTV was contoured under FDG-PET guidance with margins based on our institutional protocol. The treatment was
planned using a PLATO 2.7.7 system and was administered under imaging guidance. Clinical improvement was evident after 2 weeks of treatment, as the patient was weaned from oxygen, and able to walk and swallow liquids, and at the end of the treatment an excellent radiological response was noted. At the time this report was written, the patient was receiving adjuvant chemotherapy. We had taken a written informed consent from the patient. We have ethical committee approval.

**Discussion**

EMP is a soft tissue plasma cell tumor occurring in the absence of systemic signs of multiple myeloma, such as a bone osteolytic lesion, plasma cell infiltration in bone marrow, a lytic bone lesion, and serum or urine myeloma proteins [1]. A historical review [2] of 700 patients showed that there was head and neck involvement in 80%-90%, especially in the aerodigestive tract. Case reports of involvement of the liver, pancreas, lungs, skin, and thyroid gland do exist. It is most unusual for this tumor to present in the mediastinum as a primary solitary lesion. To date, only 12 cases of EMP involving the mediastinum have been reported in the English language literature; mean age at presentation was 55 years. Presentation (Table 1) included a mediastinum mass associated with intralobar effusion [3], pulmonary nodules [4], multiple myeloma [5,6], and mimicking hemangioma [7]; however, the presented case is unusual, as it involved a solitary anterior mediastinal mass that presented with grade 4 dyspnea, and responded quickly during the course of treatment. The plasmacytoma in the presented case was aggressive, which was evident by increased contrast uptake observed with PET, in contrast to the lack of increased uptake in indolent plasmacytoma. The presented case also highlights the clinical usefulness of PET/CT in imaging plasmacytomas, as suggested by Masood et al. [5]. Pathologically proven pleural effusion is an occasional finding in patients with multiple myeloma, which occurs in approximately 6% of patients [8]. The cause of pleural effusion in the presented case was not clear and as it was reactive, lymphatic drainage obstruction caused by the mediastinal mass might have been responsible.

Plasma cell neoplasms are relatively sensitive to radiotherapy; however, a review of the literature by Alexiou et al. [2] reported that combined therapy (surgery and radiotherapy) results in better overall and recurrence-free survival (p=0.027). Nonetheless, these results span the greater part of a century (1905-1997) and radiotherapy techniques have changed

<table>
<thead>
<tr>
<th>Serial no</th>
<th>Author</th>
<th>Presentation</th>
<th>Treatment</th>
<th>Follow-up</th>
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<tbody>
<tr>
<td>1</td>
<td>Divis and Sikl (1928)</td>
<td>Anterior mediastinum</td>
<td>Resection</td>
<td>Alive at 3 months</td>
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<tr>
<td>2</td>
<td>Bross (1931)</td>
<td>Posterior mediastinum</td>
<td>Resection</td>
<td>Necropsy report</td>
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<td>3</td>
<td>Snapper (1953)</td>
<td>Mediastinum and lung with MM</td>
<td>Radiotherapy</td>
<td>Necropsy report</td>
</tr>
<tr>
<td>4</td>
<td>Niwa1 (1987)</td>
<td>Superior mediastinum</td>
<td>-</td>
<td>-</td>
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<tr>
<td>5</td>
<td>Miyazaki (1992)</td>
<td>Anterior mediastinum + lobar effusion</td>
<td>No treatment</td>
<td>Alive at 2 years</td>
</tr>
<tr>
<td>6</td>
<td>Moran (1995)</td>
<td>Anterior mediastinum Posterior mediastinum</td>
<td>Chemotherapy Surgery</td>
<td>6 months 2 years (developed multiple myeloma)</td>
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<td>8</td>
<td>Nandekar (2000)</td>
<td>Middle mediastinum mimics hemangioma with MM</td>
<td>Chemothepapy</td>
<td>Alive at 2 years</td>
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<td>9</td>
<td>Yamaguchi (2005)</td>
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<td>Radiotherapy</td>
<td>Alive</td>
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<td>10</td>
<td>Lee (2005)</td>
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<td>Radiotherapy</td>
<td>Alive at 6 months</td>
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<td>11</td>
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<td>12</td>
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<td>Anterior mediastinum + pleural nodules + pleural effusions with MM</td>
<td>Chemotherapy</td>
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</table>
during that time. Currently, radiotherapy is the treatment of choice for solitary plasmacytoma and the dose of 40-50 Gy (depending on tumor size) is delivered over 4-6 weeks. It is associated with a <5% risk of local recurrence and most carefully staged patients with solitary EMP can be cured [9, 10]. There is no published evidence for the role of adjuvant chemotherapy in the treatment of solitary extramedullary plasmacytoma. Tsang et al. [11] and Holland et al. [12] suggest that tumours >5 cm are associated with a high risk of failure, which was a rationale for proposing chemotherapy to the presented patient.

Though 5% of patients have coexistent multiple myeloma [11], our literature review suggests 30% involvement in mediastinal EMP. This indicates that plasmacytoma involving the mediastinum provides an early hint to the diagnosis of occult multiple myeloma and should be included in the differential diagnosis of mediastinal masses. Thusly, after the diagnosis of plasmacytoma, aggressive investigation for multiple myeloma is vital, as such patients can be candidates for adjuvant chemotherapy.

Conflict of interest statement
The authors of this paper have no conflicts of interest, including specific financial interests, relationships, and/or affiliations relevant to the subject matter or materials included.

References