Solitary plasmacytoma of rib in a young adult – a rare case report

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Abstract: Solitary plasmacytoma of bone (SPB) is a rare disease, defined as clonal proliferation of plasma cells identical to those of plasma cell myeloma, which manifest as localised osseous growth with no evidence of disease dissemination. We report a case of solitary plasmacytoma of Right 6th rib in 28-year-old male. The patient underwent complete en-bloc resection of the chest wall harbouring the tumour, ribs, muscles, and parietal pleura with reconstruction of chest wall. Patient is asymptomatic and in regular follow up.

Key Words: Chest wall, plasmacytoma, ribs, solitary tumor

I. Introduction

Solitary plasmacytoma of bone (SPB) is a rare localised lesion that accounts for only 5% of all plasma cell neoplasias[1-3]. It may present as the sole manifestation of plasma cell neoplasm, as a solitary plasmacytoma of the bone or as a consequence of multiple myeloma. Solitary plasmacytoma of rib usually shows destruction of the bone cortex with extension into the surrounding soft tissues. Plasmacytoma may be primary or secondary to the disseminated multiple myeloma and may arise from the osseous (medullary) or nonosseous (extramedullary) sites. Primary extramedullary plasmacytoma can be solitary or multiple[4]. SPB has been considered a genetic abnormality that can often change to multiple myeloma[5]. Solitary plasmacytoma is rare as compared with multiple myeloma.

II. Case Report

A 28 year old nonsmoker male reported with swelling associated with right lower chest pain, gradual in onset, mild-to-moderate in nature, radiating to the back, since last 7 months. On local examination of the chest, there was a tender, solitary swelling of size 6cm x 4cm, on the right anterolateral chest wall arising from 6th rib, firm, non mobile, fixed to chest wall but free from skin, with decreased right basal air entry. Rest of systemic examination was normal. Chest X-ray showed an extra pulmonary opacity around the right 6th rib anteriorly [figure1]. On CECT of the chest a soft tissue density mass causing osteolytic destruction of right 6th rib anteriorly [figure2], indenting the adjacent lung parenchyma with no pleural effusion or pleural thickening [figure3]. CECT of the skeletal spine and head and neck was normal. Bone marrow aspirate, trephine biopsy and flowcytometry revealed increased bone marrow plasma cells (reactive plasmacytosis) with no evidence of acute leukemia/lymphoma/ metastatic disease. Plasma cells in bone-marrow differential count was 5.8%. Serum electrophoresis for M band proteins showed prominent M spike in gamma fraction. Immunofixation showed monoclonal IgA lambda. Urine for Bence Jones proteins was negative. Rest of the haematological investigations were normal. Pulmonary function test was suggestive of moderate restrictive pattern. CECT guided fine needle aspiration cytology was inconclusive.

Right anterolateral thoracotomy was done which revealed a well circumscribed tumor of 7cm × 5cm x 2 cm in size arising from the sixth rib anteriorly. En-bloc resection of the 5th, 6th and 7th rib (harboring the tumour) with involved parietal pleura and muscles was done with 5 cm from the tumor margins. Lung tissue was not adherent to the tumor. Chest wall reconstruction was done with 25 cm x 25 cm prolene mesh in double layer. Histopathology report was suggestive of plasmacytoma of the 6th rib. Margins of the resected specimen were free from the tumor. Patient fulfilled the diagnostic criteria of solitary plasmacytoma of bone[1]:

I. A radiological lytic bone lesion with histological confirmation of plasma cell cytology.
II. Clinical and radiological evidence of a solitary lesion on skeletal survey.
III. Absence of myeloma cells and plasmacytosis of less than 10% in a bone marrow examination.
IV. Absence of anemia, hypercalcemia, or renal impairment.
V. Less than 2.0 g/dl M-protein in serum.

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VI. Discussion

Localised solitary plasmacytoma of bone is a rare disease and is characterised by only one or two isolated bone lesions with no evidence of disease dissemination. Plasmacytomas can be divided into multiple, solitary osseous, and solitary extra-osseous or extramedullary plasmacytomas and rare as compared with multiple myeloma[6]. Localized SPB is a rare disease and has been considered to be curable with radiotherapy/surgical resection. This treatment is sufficient to achieve long term survival[5,7]. The incidence of SPB has been reported to be approximately 3/10 00 000 annually in US[8]. SPB is an uncommon disease that accounts for only 5% of malignant plasma cell tumors. Most commonly tumor develops along spinal column with decreasing frequency in pelvis, ribs, upper extremeties, face, skull, femur and sternum[5]. Men are diagnosed twice as often as women. SPB occurs most commonly in African-Americans and least commonly in Asians and Pacific Islanders. Mean age at diagnosis is 55-65 yrs. According to the current recommendations, the detection of a monoclonal component in the serum or urine does not exclude a diagnosis of solitary plasmacytoma[1,5]. Urine electrophoresis is an important test, because it may show abnormalities in a few patients even when the serum electrophoresis is normal.

In Japanese's literature, 14 cases of solitary plasmacytoma of rib origin have been described[9,10]. The ratio of male to female patient was approximately 1.3:1. The average age on presentation was 59.5 years with a range form 39 to 77 years[5]. Our patient presented at a young age of 28 yrs. In past, radiation therapy was used as the primary treatment for solitary plasmacytoma. Mendenhall et al, reported a 6% local failure rate in patients with solitary plasmacytoma treated with doses of 40 Gy or above in contrast to 31% for doses below 40 Gy[8]. According to Bataille and Sany, the primary methods for treating solitary plasmacytoma were surgery with radiation therapy in 95 cases and surgery alone in 15 cases[1,5]. Plasmacytoma almost always destroys bone. CECT scan and percutaneous needle biopsy are best investigations to diagnose chest lesions. The diagnosis is based on identification of the localized tumor composed of monoclonal plasma cells identical to those observed in multiple myeloma, and absence of the signs in favour of a disseminated form. In our case the needle biopsy was inconclusive. We subjected our patient to radical surgery after discussion with the tumor board of the hospital. Post-operative patient did well, he is asymptomatic and tumor free and in regular follow up with us. In cases were residual lesion persist adjuvant chemotherapy and radiotherapy should be prescribed[11,12].

V. Conclusion

Patients with solitary plasmacytoma originating in the rib have a feasibility of operative resection and radical treatment, and cure can be expected by adequate surgical resection of the tumor.
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Figure 1. Chest x ray showing tumor of 6th rib on right side

Figure 2. Three dimensional reconstructed CT film showing destructing tumor of the 6th rib on right side

Figure 3. CT film showing the rib tumor abuting the pleura but not invading the pulmonary parenchyma

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References