Chondrosarcoma of the Rib: Atypical Presentation and Management

Abstract
Chondrosarcoma is an uncommon malignant tumor of the rib and can have an atypical presentation based on age, gender, and clinical manifestation with differential diagnosis of intrathoracic mass. Management is surgical as the tumor is resistant to chemoradiation. Access to chest wall reconstruction is limited in many low-income countries and forms a barrier to patient compliance. We report an atypical presentation and describe a simple, easy, and cost-effective chest wall reconstruction method for chondrosarcoma of the rib in any resource-constrained setting.

Keywords: Chondrosarcoma rib, intrathoracic mass, polypropylene mesh, surgery

Résumé
Le chondrosarcome est une tumeur maligne peu commune de la côte et peut avoir une présentation atypique en fonction de l’âge, du sexe et de la manifestation clinique avec un diagnostic différentiel de masse intrathoracique. La prise en charge est chirurgicale car la tumeur est résistante à la chimioradiation. L'accès à la reconstruction de la paroi thoracique est limité dans de nombreux pays à faible revenu et constitue un obstacle à l’adhésion des patients. Nous rapportons une présentation atypique et décrivons une méthode de reconstruction de la paroi thoracique simple, facile et rentable pour le chondrosarcome de la côte dans un contexte de ressources limitées.

Mots-clés: Chondrosarcome de la côte, chirurgie, maille de polypropylène, masse intrathoracique

Introduction
Chondrosarcoma is a common malignant tumor that arises in the pelvis or long bones, but its presence in the ribs is uncommon, with an incidence of 0.5 per million person-years.[1] They represent 20% of all chest wall neoplasms.[2] Contrast-enhanced computed tomography is the gold standard investigation of choice for appropriate diagnosis and operative planning.[3] They have significant potential for metastasis and are relatively resistant to chemoradiotherapy. Surgical excision with negative microscopic margins is the treatment of choice. This case report depicts an atypical presentation of primary chest wall mass resembling an intrathoracic tumor diagnosed to be chondrosarcoma managed by surgical excision and reconstruction.

Case Report
A 37-year-old woman who presented with mild chest pain had a chest X-ray done, which showed an opacity in the right lower zone [Figure 1A]. She was referred to the Department of Cardiothoracic Surgery in October 2020. She was hemodynamically stable and afebrile. Her chest and abdominal examinations did not reveal any masses on palpation. Contrast-enhanced computed tomography scan (CECT) [Figure 1B] revealed an exophytic lesion measuring 4.9 cm × 4.9 cm × 8.6 cm, with calcification, arising from the right sixth rib and involving the seventh rib partially. The lesion showed mild post-contrast enhancement with no infiltration of adjacent structures. A CT-guided biopsy of the lesion was inconclusive. Under general anesthesia, she had surgical excision of the lesion, with wide excision margins, and chest wall reconstruction. An oblique incision was placed over the area of the lesion [Figure 2A] after localization with ultrasonography in the operating room. A reddish-white, hard mass measuring about 8 cm × 5 cm was found arising from the sixth rib and infiltrating a small segment of the seventh rib, abutting the diaphragm. The lesion was considered malignant. The lesion was excised, together with adjacent ribs, one above and one below, with excision margins of 4 cm [Figure 2B] without much difficulty.
A single chest drain was placed in a dependent position just above the diaphragm. The chest wall defect, measuring 12 cm × 8 cm was repaired with polypropylene mesh. As the defect was large, two meshes were joined adjacently and folded into two layers. Multiple prolene 2-0 stay sutures were placed in the defect connecting adjacent ribs. The mesh was placed over the defect and sutured with interrupted prolene 3-0 to the intercostal muscles [Figure 2C]. This provided better stability to the chest wall and prevented paradoxical chest wall movement. The wound was closed in layers. The postoperative course was uneventful, with the drain removed on the third postoperative day (POD). She was discharged on the seventh POD and the wound had healed by primary intention at two weeks follow-up [Figure 2C]. Histopathology showed chondroid neoplasm comprising of lobules of pleomorphic chondrocytes in binucleate forms with occasional mitosis and uninvolved bony resection margins, suggestive of grade 1 chondrosarcoma [Figure 3]. Her follow-up visits in the third and sixth months with a CT scan did not detect any recurrence.

**Discussion**

Chest wall chondrosarcoma is a slow-growing tumor presenting as painful swelling, especially in the fifth decade of life with the male gender being predominantly affected.[3,4] They usually have an anterior location arising from the costochondral junction or sternum.[3] Primary chondrosarcomas arise without an underlying etiology, whereas secondary chondrosarcomas arise from pre-existing enchondromas or osteochondromas.[5] The patient in the present case study was a female in the fourth decade of life presenting with mild chest pain and in whom a chest X-ray detected a lesion. This is an unusual presentation seen in 6.5% of individuals who have what looks like an intrathoracic mass without a pre-existing lesion, suggesting a primary etiology in origin.[1,6]
Intratumoral calcification on CT in the form of a “flocculent or popcorn” pattern has been described as the hallmark of chondrosarcoma, similarly was present in our case. They can also have associated bony destruction and soft tissue invasion; however, these were not present in the case presented.

A positron emission tomography (PET) scan can detect extrapulmonary metastases and provide better insight into the tumor grade as well as predict postoperative surgical outcomes. Magnetic resonance imaging is not routinely performed but can be used to assess extraosseous extension. As per the Surveillance, Epidemiology, and End Results (SEER) cancer registry database, tumor grade histologically is classified into four grades: well differentiated (grade 1), moderately differentiated (grade 2), poorly differentiated (grade 3), and undifferentiated (grade 4). Tumor grades have a definite role as a predictor for local recurrence, systemic metastasis, and survival.

Surgical excision with microscopically negative margins is the treatment of choice irrespective of the tumor grade, and chest wall reconstruction is recommended for full-thickness defect of more than 5 cm or anterior resection involving more than three ribs. Various techniques have been used in the reconstruction of the chest wall including meshes, methyl methacrylate cement, flaps, titanium plates, and the creation of neo-rib. However, we preferred using double-layered polypropylene mesh because of its inert property, ease of processing, and versatility for conversion into various shapes; it is also inexpensive, readily available, and produces promising results in extensive chest wall reconstruction. The role of Doxurubin-based adjuvant therapy in grade 2 and 3 mesenchymal tumors has been described, but the available data is non-randomized and small. The role of radiotherapy in high-grade tumors is still unclear and it has failed to achieve better overall survival or cancer-specific survival.

Immune checkpoint blockade therapy is an emerging strategy in the management of chondrosarcoma. However, clinical trial outcomes are still expected, and the role of predictive biomarkers to determine efficacy is an essential step in the future. Post-operative surveillance consists of physical examination and thoracic imaging with chest X-rays or CT scans every six months for the first five years and thereafter annually for a minimum of ten years.

**Conclusion**

Chondrosarcoma of the rib is a rare tumor and can have an unusual presentation. Surgical management with negative margins is necessary to obtain beneficial results.

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**Conflicts of interest**

There are no conflicts of interest.

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