

Revista Portuguesa de Pneumología ISSN: 0873-2159 sppneumologia@mail.telepac.pt

Sociedade Portuguesa de Pneumologia Portugal

Elsayed, H.; Gosney, J.

A massive pleural-based tumour: The challenge of diagnosis

Revista Portuguesa de Pneumología, vol. 17, núm. 6, noviembre-diciembre, 2011, pp. 275-277

Sociedade Portuguesa de Pneumologia

Lisboa, Portugal

Available in: http://www.redalyc.org/articulo.oa?id=169722769008



Complete issue

More information about this article

Journal's homepage in redalyc.org



Rev Port Pneumol. 2011;17(6):275-277



# revista portuguesa de portuguese journal of pulmonology



www.revportpneumol.org

#### CASE REPORT

## A massive pleural-based tumour: The challenge of diagnosis

H. Elsayed a,b,\*, J. Gosney c

<sup>a</sup> Cardiothoracic Department, Liverpool Heart and Chest Hospital, United Kingdom

reserved.

- <sup>b</sup> Thoracic Surgery Department, Ain Shams University, Cairo, Egypt
- <sup>c</sup> Histopathology Department, Royal Liverpool University Hospital Trust, Liverpool, United Kingdom

Received 10 December 2010; accepted 6 June 2011 Available online 21 July 2011

#### **KEYWORDS**

Liposarcoma; Pleural based tumours; Rare tumours

however, primary pleural liposarcoma is extremely rare.

the origin of these tumours to allow accurate diagnosis.

Um tumor gigante de base pleural: o desafio do diagnóstico Resumo Os lipossarcomas do mediastino e da parede torácica foram descritos anteriorment

no entanto, o lipossarcoma pleural primário é extremamente raro. Registamos um caso extremamente raro de um lipossarcoma esclerosante da pleura bem dife enciado num homem de 47 anos que foi reseccionado. Enfatizamos a importância da verificaç

Abstract Liposarcomas of the mediastinum and chest wall have been previously describe

We report an exceedingly rare case of a well-differentiated sclerosing pleural liposarcoma

a 47-year-old male that was resected. We emphasise the importance of careful inspection

© 2010 Sociedade Portuguesa de Pneumologia. Published by Elsevier España, S.L. All righ

cuidada da origem destes tumores, para permitir um diagnóstico preciso.

© 2010 Sociedade Portuguesa de Pneumologia. Publicado por Elsevier España, S.L. Todos direitos reservados.

#### **PALAVRAS-CHAVE**

Lipossarcoma; Tumores na pleura; Tumores raros

### Introduction

Liposarcoma most commonly arises in the retro peritoneum or thigh, but can also involve numerous other anatomical sites, such as the inguinal area, popliteal fossa, buttock, shoulder, arm, neck, back, and scalp, as well as the genitourinary and aerodigestive tracts. 1 Liposarcomas of the

reported once in the English literature. Case report

A 47-year-old gentleman was referred by his general pra tioner. He had been complaining of non-specific chest pa

mediastinum and chest wall have been previously describe however, primary pleural liposarcoma is very rare and or a few isolated cases have been reported. We report

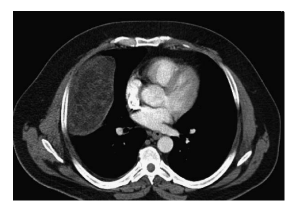
exceedingly rare case of a well-differentiated sclerosi

liposarcoma in a 47-year-old male, which has been or

0873-2159/\$ - see front matter © 2010 Sociedade Portuguesa de Pneumologia. Published by Elsevier España, S.L. All rights reserved. doi:10.1016/j.rppneu.2011.06.001

<sup>\*</sup> Corresponding author. E-mail address: drhany.elsayed@yahoo.co.uk (H. Elsayed).

276 H. Elsayed, J. Go



**Figure 1** A large mass in the right hemi-thorax with low density suggestive of a lipoma/liposarcoma.

for the past 2 years. Over the past 2 months, the pain had become more frequent and sharp in nature. Additionally, he had begun to complain of shortness of breath with a NYHA classification of 2. Apart from that, he was fit with no significant past medical history apart from a fractured elbow and whiplash injury after a road traffic accident in 2002. His current medications included only painkillers on demand. He was a current smoker with a history of 20 packs per year. He drinks no alcohol.

A chest X-ray was done which was commented on as suspicious of diaphragmatic hernia, and hence was referred to the thoracic surgeons.

On presentation, this gentleman was fit and well. Cardiorespiratory revealed decreased air entry on the right basal lung field. The rest of examination was normal. We performed a CT-scan (Fig. 1), which confirmed the presence of a huge mass of an unidentified nature occupying the right hemi thorax. Its measurements were  $17\,\mathrm{cm} \times 13\,\mathrm{cm} \times 10\,\mathrm{cm}$ . A PET FDG confirmed the mass to be of normal activity with no other abnormal uptake elsewhere.

On the basis of the patient's symptoms and the likelihood of resectability, we proceeded with a right thoractomy with single lung ventilation. A huge intrapleural mass measuring  $18\,\text{cm}\times12\,\text{cm}\times10\,\text{cm}$  and weighing 640 g was found attached to the inner surface of the chest wall by a 1 cm stalk. No other masses were found elsewhere. The mass was



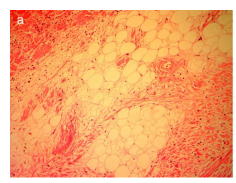
**Figure 2** Gross appearance of the excised mass after rem from the chest.

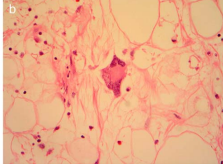
dissected and removed en bloc and was found to be part encapsulated (Fig. 2). Routine haemostasis and closure v performed.

Postoperatively, the patient made an excellent reery and was discharged home three days later. Histopath came back as a huge tumour with sheets of adipocyte variable size with broad fibrous bands and extensive ness. A thick fibrous capsule surrounded the tumour appearance was consistent with a well-differentiated rosing liposarcoma with no areas of high-grade sarc (Fig. 3(a and b)). No immunohistochemistry was needed morphology was diagnostic. On the basis of the pathology our oncologist suggested the need for adjuvant radiother

#### Comment

Liposarcomas of the chest wall and mediastinum been very rarely reported, and distinction from ple liposarcoma with chest wall or mediastinal invasion requesterful radiographic evaluation, surgical evaluation both. In some cases it may be impossible to be certain with tumour is arising but the main differentiating feature whether the tumour is intra or extra-pleural in original part of the control of the con





**Figure 3** (a) Medium power photomicrograph showing mature adipocytes separated by collagenous fibrous tissue (sclero feature). An occasional lipoblast with an enlarged hyper chromatic nucleus is evident within the fat. (b) High power view sho an atypical lipoblast, the presence of which indicates malignant well differentiated sclerosing liposarcoma.

our case, the tumour appeared to be intrapleural, although there was a small stalk attached to the chest wall.

After recognizing the diagnostic histologic features of a liposarcoma, one must exclude the possibility of metastases or spread from either the chest wall or mediastinum.<sup>2</sup> Other sarcomas that may be confused with liposarcoma include malignant fibrous histiocytoma, in which bizarre pleomorphic giant cells may be mistaken for pleomorphic liposarcoma. However, characteristic lipoblasts should be evident in liposarcoma.<sup>3</sup>

McGregor et al.<sup>4</sup> reported the only recorded case of welldifferentiated and pleomorphic liposarcoma of the pleural cavity that recurred 2 years following excision; the histologic pattern of this tumour resembled malignant fibrous histiocytoma.

The rarity of liposarcomas arising in the pleural cavity makes it difficult to identify prognostic factors that correlate with survival. The issue of surgical resectability is of paramount importance for patient survival, especially in low-grade sarcomas.<sup>5</sup>

Based on the small number of pleural liposarcomas reported to date, it is difficult to know if there is clinical importance in separating resectable pleural liposarcomas from chest wall sarcomas. Whether complete resection with adequate surgical margins improves survival for pleural liposarcomas needs further evaluation with a larger number of cases. Our oncologist recommended adjuvant radiotherapy which is supported by other studies in which radiation therapy was instituted, later onset of recurrence and longer disease-free survival were more likely.<sup>6</sup>

Of importance to the pathologist is the histopathologic subtype of liposarcoma. The classification of liposarcoma is based on 4 histologic categories that generally reflect the degree of differentiation, ranging from well-differentiated liposarcoma to myxoid/round cell, undifferentiated, and, finally, pleomorphic liposarcomas. Well-differentiated liposarcomas can be further sub-classified as lipoma-like, inflammatory, and sclerosing, based on features of similarity to lipomas, lymphoplasmacytic infiltrate, or areas of loose to dense fibrous stroma, respectively.<sup>7</sup>

It is very difficult to comment on the prognosis of t subtypes of well differentiated liposarcoma as this is t only second reported case in such an entity and this w require us to gather more cases with longer periods follow up. Our case, as well as those identified in the l erature, demonstrates that the most common histolog subtypes of liposarcoma in the pleura appear to be myxc and well differentiated. All forms of liposarcoma contalipoblasts in varying numbers and morphologic forms; ho ever, the more poorly differentiated liposarcomas behamore aggressively and can produce widespread metastases.

#### Conclusion

Liposarcomas of the pleura are very rare tumours diagnosmainly via CT-scans with lipomas being the main different diagnosis. The challenge of diagnosis remains in identifying its origin and it needs careful inspection from the surge before resection.

#### References

- Enzinger FM, Weiss SW. Soft tissue tumors. St Louis, MO: Most Year Book; 1995.
- Carroll F, Kramer MD, Acinapura AJ. Pleural liposarcoma pi senting with respiratory distress and suspected diaphragma hernia. Ann Thorac Surg. 1992;54:1212-3.
- Evans A, Wolstenholme R, Shettar S, Yogish H. Primary pleu liposarcoma. Thorax. 1985;40:554–5.
- McGregor DH, Dixon AY, Moral L, Kanabe S. Liposarcoma of ple ral cavity with recurrence as malignant fibrous histiocytom Ann Clin Lab Sci. 1987;17:83–92.
- Wong WW, Pluth JR, Grado GL, Schild SE, Sanderson DR. Liposi coma of the pleura. Mayo Clin Proc. 1994;69:882–5.
- Gupta RK, Paolini FA. Liposarcoma of the pleura: report of a ca with a review of the literature and views on histogenesis. A Rev Respir Dis. 1967;95:298–304.
- Snover DC, Sumner HW, Dehner LP. Variability of histologic pitern in recurrent soft tissue sarcomas originally diagnosed liposarcoma. Cancer. 1982;49:1005–15.
- Wouters EFM, Greve LH. Liposarcoma of the pleura. Neth J Su 1983;35:192–3.