Bleeding as a complication of fine needle lung biopsy

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ABSTRACT Two patients in whom bleeding into the pleural space was a major complication of fine needle biopsy of the lung are described. Both had a pleural effusion before biopsy.

In the 20 years since the work of Dahlgren and Nordenstrom showed 87% diagnostic accuracy and no serious complications in 365 cases, fine needle biopsy of the lung has been accepted as a minimally invasive procedure. It is widely and correctly considered to be safe, although there is a well-recognised incidence of minor complications, principally pneumothorax. We report two major complications, one fatal.

Case reports

PATIENT 1
A 32 year old woman was admitted to our hospital with a one week history of pleuritic chest pain, fever, and cough. A plain chest radiograph and a computed tomography scan (fig 1a) showed a 5 cm left lower anterior chest wall mass adjacent to and eroding the anterior end of the left fourth rib. There was also a small left pleural effusion. She had a normal clotting profile and underwent computed tomography guided biopsy with a 20G Surgimed-Rotex screw needle, an anterior approach being used. The biopsy showed syncytia of malignant cells with small nucleoli and the microglandular pattern of adenocarcinoma. Two days after the procedure she complained of left breast pain. Chest radiographs showed a progressively increasing left pleural fluid collection (fig 1b); needle aspiration of this showed only a little bloody fluid, which did not contain any neoplastic cells. The pleural opacity persisted despite intercostal catheter drainage of fluid, which was mostly blood. Because of continued bleeding that required blood transfusion and also because of pain and fever, the patient underwent a left anterior thoracotomy 12 days after biopsy. A large blood clot was evacuated from the pleural space, and a 10 cm diameter mass of friable tumour and blood clot was incompletely removed from behind the 4th rib. No bleeding point was found. The final histopathological diagnosis was of poorly differentiated spindle cell sarcoma. Bleeding subsequently ceased and radiotherapy was given when the wound had healed.

PATIENT 2
A 70 year old hypertensive, cigarette smoking woman was admitted for investigation of haemoptysis, and a right lower lobe lung mass and pleural effusion were seen on the chest radiograph and computed tomogram (fig 2a). She had a normal blood clotting profile and underwent a computed tomography guided biopsy with a 20G Surgimed-Rotex screw needle, together with aspiration of pleural fluid. Examination of both fluid and biopsy material revealed squamous cell carcinoma. Two radiotherapy treatments were given. After the biopsy recurrent right sided pleural haemorrhage (fig 2b) required intercostal tube drainage, transfusion of nine units of blood, and eventually, 19 days after biopsy, thoracotomy. A clotted haemothorax was evacuated and partial decortication was required to re-expand the lung. A small (1-5 cm diameter) mass of adherent blood clot was found on the pleural surface overlying the tumour. This was removed and a possible bleeding point sealed by diathermy. The friable blood clot contained aggregates of tumour cells. An attempt was made to induce pleurodesis with 1% silver nitrate solution, but pleural bleeding continued and drains were reinserted. Despite blood transfusion and bronchial artery embolisation she eventually succumbed to persisting blood and pleural fluid loss and died 46 days after the original biopsy.

Discussion

Diagnostic needle aspiration of the lung was first performed by Leyden in 1883 to obtain pneumonic organisms. This was 12 years before the discovery of x rays. Worldwide experience with the technique is now considerable. Although many needle designs are available, most are 20 to 22G in diameter. Published reports suggest that the complication rate is more dependent on the operator's experience than on the needle used. Sinner has shown that the accuracy rates are fairly similar in different series; it was 90-7% in an overview of 5300 biopsies. The most common complication is pneumothorax, which occurs in 20% to 34% of biopsies, up to 10% requiring treatment. Pneumothorax occurs more frequently when biopsy is performed through the anterior chest wall (22%) than via the posterior approach (12%). Spread of lung tumour along the needle track after aspiration biopsy has been reported but is extremely uncommon.

About 10% of patients experience a short period of haemoptysis immediately after biopsy and minor pleural haematomas occur in 1%. We have found only one case report of death attributable to fine needle biopsy apart from ours, though deaths after lung biopsy with large bore needles have been reported. There have been two reports of potentially fatal complications associated with the fine needle technique. In one tracheobronchial haemorrhage resulted in cardiac arrest and in the other acute cardiac tamponade.
resulted from haemorrhage of a biopsied lesion near the mediastinum, emphasising the need for care in this area.

Pleural space bleeding should be recognised as a potentially serious complication of fine needle aspiration biopsy. The problem is most likely to occur if the lesion is adjacent to a pleural surface. The combination of continuous respiratory movement and the lack of tamponade in the pleural space may lead to major haemorrhage despite a normal clotting profile, as in our cases. In the second case there was a substantial pleural effusion at the time of biopsy and this is likely also to have prevented tamponade of bleeding once the tumour had been punctured. It could reasonably be argued that pleural fluid aspiration alone would have been sufficient to make a cytological diagnosis in this case and that tumour biopsy was unnecessary or even contraindicated. At present, however, a pleural fluid collection is not generally regarded by interventional radiologists as a contraindication to biopsy of an ipsilateral lung mass. Perhaps it should be considered as a potential hazard in the future.

References