Percutaneous biopsy of mediastinal tumours under sonographic guidance

Until the early 1980s biopsies of mediastinal tumours were performed exclusively under fluoroscopic control, following the report by Nordenström in 1967.1-4 Computed tomography now has largely replaced fluoroscopy for this purpose as it allows accurate localisation of the lesion and depicts its anatomical relation to the large mediastinal vessels.5,6 Accidental puncture of the mediastinal vessels, which is not uncommon under fluoroscopic guidance,7 can be largely avoided with exact computed tomography guided planning of the biopsy route. In addition, the exact location of the needle tip in the target lesion can be reviewed before biopsy.

In 1986 and 1988 we described the techniques of supraperasternal and parasternal mediastinal sonography, which allows a considerable part of the mediastinum to be assessed.7-8 We have applied this technique to guidance of percutaneous mediastinal tumour biopsy,7 because sonographic guidance has all the advantages of spatial orientation offered by computed tomography and also provides continuous monitoring of the biopsy under real time conditions.

SELECTION OF PATIENTS FOR SONOGRAPHICALLY GUIDED BIOPSY

The essential criterion for selection of patients is the availability of an avascular approach to the mediastinal tumour at sonography. With the use of both a supraperasternal and a parasternal approach sonography has proved to be highly sensitive in the detection of tumours in most parts of the mediastinum.9 The approach to certain mediastinal regions (paratracheal, subcarinal, and aorticopulmonary) is, however, obstructed by mediastinal vessels and sonographically guided biopsy is not appropriate. Sonographic guidance is mainly suitable for the biopsy of anterior mediastinal and supra-aortic tumours.9

TECHNIQUE OF SONOGRAPHICALLY GUIDED BIOPSY

Preliminary sonographic examination with a 3-5 MHz convex probe serves to locate the tumour accurately and to define its anatomical relations to enable the optimal biopsy approach to be selected. The biopsy is performed by means of a sterilisable biopsy probe with a central perforation for needle guidance (LSC 7000, Picker International, Munich). The needle guide can be adjusted to any angle in the range 0–30°. The trajectory line is displayed electronically on the monitor (figs 1 and 2).

Most sonographically guided biopsies are performed via the right or left parasternal approach with the patient in the supine, right, or left decubitus position. The biopsy route is planned to avoid all mediastinal vessels, even for small calibre needles (20 gauge). To avoid inadvertent puncture of the internal mammary artery, the immediate parasternal (fig 2) or lateral approach (more than 2·5 cm from the lateral margin of the sternum—see figure 2) should be selected.11 When a large cutting needle is used the position of the mammary vessels can be displayed by preliminary sonographic examination with 5-0 or 7-5 MHz transducers (fig 3).12 The large mediastinal vessels should be avoided by selecting an appropriate angle of entry of the biopsy route (fig 1). Mediastinal tumours of the supra-aortic regions are biopsied via the suprasternal approach. The patient is in the supine position with cushions placed under the shoulder girdle. The transducer is placed immediately above the pre sternum in the jugular fossa.

The choice of biopsy needle depends on tumour size, tumour location, and the clinical questions being asked. Mediastinal tumours of unknown cause should be biopsied...
with a 14 gauge cutting biopsy needle (Biopty-cut, Radiplat, Uppsala, Sweden) when tumour size and location permit. As the tip of the Biopty-cut needle is difficult to localise precisely (in millimetres), the large cutting needles should be used only for mediastinal tumours greater than 3.0 cm in diameter. Biopsies of tumours with a diameter of less than 3 cm are performed with a 20 gauge or 18 gauge needle (TSK Supra and Surecut, TSK Laboratories, Tokyo). If mediastinal tumour biopsy is indicated for staging of a malignant tumour of known origin (for example, bronchogenic carcinoma or breast carcinoma), the use of a fine needle (20 gauge) is generally sufficient to obtain an adequate diagnosis.

After administration of local cutaneous anaesthesia the needle is inserted through the biopsy sleeve and pushed as far as the tumour under real time control. Rapid repeated insertion and withdrawal of the fine needle and simultaneous aspiration are required to obtain the tissue specimens. Before biopsy with a cutting needle one pass with a fine needle is generally made to assess the consistency and vascularity of the tumour. Further passes with a fine or cutting needle (or both) are done in different parts of the tumour until adequate tissue cylinders are obtained. We generally perform three to five passes even with a large cutting needle, to obtain representative tissue cylinders from different parts of the tumour.

After the biopsy pneumothorax can be excluded initially by sonography. We also obtain an expiratory postero-anterior chest radiograph four hours after the biopsy.

**RESULTS OF SONOGRAPHICALLY GUIDED BIOPSY**

The results of sonographically guided mediastinal tumour biopsies are comparable to those of computed tomography guided biopsies (table). As the needle can be positioned precisely within anterior and supra-aortic mediastinal lesions under real time control, the results of biopsy depend mainly on the quality of the tissue cylinders and the histological nature of the tumour.

<table>
<thead>
<tr>
<th>Definitive diagnosis</th>
<th>No of patients</th>
<th>Needle diameter (mm)</th>
<th>Correct differentiation (benign/malignant)</th>
<th>Correct histological diagnosis</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Malignant tumours</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hodgkin’s lymphoma</td>
<td>9</td>
<td>2.0 (n = 6)</td>
<td>6</td>
<td>6</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1.2 (n = 2)</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1.0 (n = 1)</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Non-Hodgkin’s lymphoma</td>
<td>3</td>
<td>2.0 (n = 1)</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1.0 (n = 2)</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>Thymoma</td>
<td>2</td>
<td>1.5</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Thymic carcinoma</td>
<td>1</td>
<td>2.0</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Bronchogenic carcinoma</td>
<td>2</td>
<td>2.0 (n = 1)</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Breast carcinoma</td>
<td>2</td>
<td>1.2 (n = 1)</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Adenocarcinoma (uterus)</td>
<td>1</td>
<td>1.0</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Poorly differentiated small cell epithelial tumour</td>
<td>1</td>
<td>1.0</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td><strong>Benign tumours</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Scar tissue after non-Hodgkin’s lymphoma</td>
<td>1</td>
<td>1.0</td>
<td>1</td>
<td>—</td>
</tr>
<tr>
<td>Mediastinal thyroid cyst</td>
<td>1</td>
<td>1.0</td>
<td>1</td>
<td>—</td>
</tr>
<tr>
<td>Mediastinal haematoma</td>
<td>1</td>
<td>1.0</td>
<td>1</td>
<td>—</td>
</tr>
</tbody>
</table>
Percutaneous biopsy is highly accurate in the diagnosis of carcinomas of the mediastinum that have metastasised either from the lung or from extrapulmonary sites. As epithelial metastases are frequently highly cellular, even small tissue specimens are sufficiently representative to obtain an adequate diagnosis.

In general, the results of percutaneous biopsy of mediastinal lymphoma are not as satisfactory as those for carcinoma. In our series the correct histological diagnosis was reached in eight of nine cases of Hodgkin’s lymphoma by identification of Reed-Sternberg giant cells in large tissue cores. In contrast, no definite histological classification was possible in two of three patients with non-Hodgkin’s lymphoma. This may be related in part to the large amount of reactive inflammatory or fibrotic tissue associated with or surrounding the lymphomatous masses. Pathological differentiation of mediastinal lymphomas and thymomas from a fine needle biopsy specimen and even from histological specimens is difficult. As differentiation between a thymoma, which may require surgery, and systemic lymphatic malignancy is of therapeutic relevance, biopsy needles with a larger diameter (14 gauge) should be used for tumours of adequate size to obtain a large volume of intact tissue. It is important to emphasise that sarcomas and other rare primary mediastinal tumours cannot be diagnosed accurately by percutaneous biopsy even on the basis of large tissue cylinders.

CONCLUSIONS

Sonographically guided biopsy of mediastinal tumours is a technically simple, rapid and accurate procedure that has all the advantages of spatial orientation offered by computed tomography and, in addition, provides continuous monitoring of the biopsy under real time conditions. Its application is limited, however, by the location of the tumour. In our experience, only tumours of the anterior mediastinum and the supra-aortic region permit safe biopsy with this technique. The main indication for biopsy of an anterior mediastinal lesion is to differentiate between a primary mediastinal tumour that requires surgery and systemic lymphatic malignancy. As histological differentiation between lymphomas and thymomas is difficult, cutting biopsy needles of larger diameter should be used initially, when tumour size and location permit. With larger cutting needles the biopsy must be monitored more accurately because the potential risk of vascular trauma increases; real time sonography is ideal for this purpose. In our sample of patients (more than 30 sonographically guided mediastinal biopsies) no complications such as haemorrhage or pneumothorax were encountered despite the increasing use of large cutting biopsy needles.

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