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Endobronchial ultrasound-guided transbronchial needle aspiration in lung cancer: the first experience in Hong Kong

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Objective To investigate the diagnostic performance and safety of endobronchial ultrasound-guided transbronchial needle aspiration in patients presenting with radiological features of lung cancer.

Design Prospective case series.

Setting University teaching hospital, Hong Kong.

Patients Consecutive patients with mediastinal or hilar abnormalities suspected of or confirmed as having lung cancer underwent endobronchial ultrasound-guided transbronchial needle aspiration and presented between August 2006 and December 2010.

Main outcome measures Diagnostic performance (including sensitivity, specificity, negative predictive value and accuracy), procedural complications, and tissue adequacy for molecular profiling.

Results A total of 269 procedures were performed in 259 patients, with malignancy confirmed in 210 (81%) of them. In the whole cohort with confirmed or suspected lung cancer, the overall sensitivity, specificity, negative predictive value, and accuracy of endobronchial ultrasound-guided transbronchial needle aspiration were 87%, 100%, 74%, and 91%, respectively. Among 42 patients with tumour samples sent for mutation tests (epidermal growth factor receptor and/or anaplastic lymphoma kinase), 40 (95%) were found to be adequate. No complication or mortality ensued from these procedures.

Conclusion Endobronchial ultrasound-guided transbronchial needle aspiration is highly effective in determining the diagnosis and lymph node staging in patients with lung cancer. In combination with its excellent safety profile, it should be considered a frontline diagnostic test for patients presenting with mediastinal abnormalities suspicious of lung cancer.

Key words Biopsy, fine-needle; Bronchoscopy; Lung neoplasms; Neoplasm staging; Sensitivity and specificity

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New knowledge added by this study
• Endobronchial ultrasound-guided transbronchial needle aspiration (EBUS-TBNA) carries high diagnostic accuracy for lung cancer in Hong Kong.
• The safety of EBUS-TBNA has been well-demonstrated in our local setting.

Implications for clinical practice or policy
• EBUS-TBNA should be considered as one of the upfront investigations-of-choice for suspected lung cancer.

Introduction
When there is an abnormal lung lesion evident on radiological examination, the next step is to establish the diagnosis and accurately determine its stage if lung cancer is suspected. Both computed tomography (CT) and positron emission tomography (PET) are effective in localising the primary lesion and any metastatic foci. However, pathological confirmation remains the gold standard, especially in Hong Kong, where tuberculosis is endemic and not uncommonly masquerades as malignancy.

Conventional flexible fiberoptic bronchoscopy (FOB) is useful in the diagnosis of endobronchial lesions, but many lung cancers are extraluminal and thus invisible...
during bronchoscopy. Because of the extensive branching and angulation of the lower bronchial tree, the diagnostic yield of FOB coupled with fluoroscopy-guided transbronchial lung biopsy (TBLB) is still suboptimal for parenchymal lung lesions, and depends on the lesion's size and location. Conventional transbronchial needle aspiration (TBNA) without image guidance has high diagnostic yield for lesions adjacent to the airway, but has been underutilised in most parts of the world including Hong Kong, partly due to reported serious complications such as bleeding and pneumothorax. Alternatively, CT-guided needle aspiration is useful in the diagnosis of peripheral lung lesions but carries significant risks of a pneumothorax (25%) and having a chest tube inserted (5%). Pathological lymph node staging of the mediastinum remains unattainable with both TBLB and CT-guided needle biopsy.

Recently, endobronchial ultrasound-guided transbronchial needle aspiration (EBUS-TBNA) has been shown to be a highly accurate and safe procedure for the diagnosis and staging of patients with confirmed or suspected lung cancer. The procedure can be incorporated with conventional FOB in a single bronchoscopic session using local anaesthesia. This can provide adequate tissue to assess epidermal growth factor receptor (EGFR) mutation status to guide treatment with modern targeted oral therapeutic agents. Since the first introduction of EBUS-TBNA in Hong Kong in 2005, the technology has now become widely available in the community. As pioneers in adopting this new technology locally, we describe the first experience in Hong Kong and discuss its role in comparison with currently available alternatives.

Thus, the aim of this study was to define the overall diagnostic performance of EBUS-TBNA in the setting of suspected or confirmed lung cancer, as well as according to the final diagnoses including malignancy and infection. The performance of EBUS-TBNA as the first pathological diagnostic test was also defined.

Methods

Recruitment and procedure of endobronchial ultrasound-guided transbronchial needle aspiration

This study was performed prospectively between August 2006 and December 2010, allowing at least 1 year of clinical follow-up at the time of analysis. Queen Mary Hospital was the first centre in Hong Kong that provided EBUS-TBNA with the approval from the Hospital Authority. All patients having confirmed or suspected lung cancer referred for EBUS-TBNA with enlarged (short diameter >1 cm) mediastinal/hilar lesions or lung lesions closely abutting adjacent airways on CT (with or without PET confirmation) were recruited. If the target lesion was subcentimetre on CT and yet 18F-fluorodeoxyglucose avid on PET, EBUS-TBNA was also considered. All the procedures were performed by a chief operator (MKW) who received relevant training in year 2005 and launched the EBUS-TBNA service in Queen Mary Hospital from 2006. The abnormal lymph node stations as detected by CT scan and/or PET were taken as reference. If more than one lymph node station was involved, the procedure was performed in the order N3 to N1 node, according to the seventh edition of the American Joint Committee on Cancer cancer staging manual. Contra-indications and preparations were the same as for patients undergoing FOB with biopsy intent, and included having a medically unstable condition, bleeding diathesis, and lack of consent. The procedures were performed under conscious sedation with midazolam and pethidine used intravenously, in addition to local anaesthesia. The specially designed EBUS scope (XBFUC260F-OL8; Olympus, Tokyo, Japan) had the curvilinear transducer incorporated at the tip with an outer diameter of 6.7 mm. It was administered trans-orally. A saline-inflated balloon kept the probe in contact with the bronchial wall while sampling the lymph node. The system had an integrated colour
Doppler function, with enhanced visualisation of blood vessels to prevent inadvertent puncturing. A dedicated 22-gauge fine needle was deployed to perform TBNA under direct visualisation, so that the excursion of the needle within the lesion could be maximised and large histological cores could be obtained. Rapid-on-site cytological examination (ROSE) was not available in our centre and the procedure was stopped if no satisfactory specimen was obtained after five needle passes per lesion. All specimens were sent for histological (preserved in 10% buffered formalin), cytological (fixed in 50% alcohol), and microbiological examinations (acid-fast bacilli and fungi). Molecular profiling, namely EGFR mutation tests, were analysed for those with confirmed adenocarcinoma from October 2008 when the test became commonly adopted. Conventional FOB was carried out if indicated. Chest radiography was performed after the patients were transferred back to the wards to exclude pneumothorax. Procedure-related complications such as excessive bleeding, infection, pneumothorax, and respiratory failure warranting intervention were to be recorded.

Written informed consent was obtained from all study patients, and the study was approved by the Institutional Review Board of the University of Hong Kong/Hospital Authority Hong Kong West Cluster (HKU/HA HKW IRB: UW 06-236T/1261).

Statistical analyses
In this study, diagnostic performance was procedure-based. A positive result was defined if a specific diagnosis (e.g., carcinoma or tuberculosis) was made, without the need for surgical diagnostic intervention. A negative result referred to proceeding to further investigation (e.g., mediastinoscopy or thoracotomy), which was always undertaken unless the patient was unfit for such a procedure or declined it. A false-negative result was defined as no specific diagnosis until further investigation yielded positive findings from the target area or follow-up eventually confirmed a positive diagnostic result in the area of interest. True negative was defined as a biopsy being confirmed as negative after surgical exploration, or unremarkable follow-up for at least 1 year. The diagnostic yield including sensitivity (true positives/[true positives + false negatives]), negative predictive value (true negatives/[true negatives + false negatives]), and accuracy ([true positives + true negatives]/[true positives + true negatives + false positives + false negatives]) were calculated. Taking the pathological or microbiological diagnosis revealed by EBUS-TBNA as the gold standard, false positives were assumed to be absent.

Skewed continuous and normally distributed variables were presented as medians (ranges), and means and standard deviations (SDs), respectively.

Results
Clinical and demographic characteristics
Over the study period of 4 years and 4 months, there were 269 EBUS-TBNAs performed in 259 patients, which included three who had repeat procedures to confirm the diagnosis. Seven patients who had first EBUS-TBNA–confirmed lung cancer underwent mediastinal restaging by EBUS-TBNA after neoadjuvant therapy. The median age of these 259 patients was 62 (range, 24-86) years; all were Asians and there was a predominance of males (63%). The
The prevalence of confirmed malignancy in this cohort was 81% (210 patients), among whom malignancy was revealed in specimens obtained by EBUS-TBNA in 158 (75%; Table 1). The characteristics of the lesions sampled are shown in Table 2. Regarding the 260 EBUS-TBNA procedures performed, 38 (14%) had lung lesions sampled, of which 29 (76%) were found to be diagnostic. Whereas, 231 (86%) of these procedures entailed mediastinal lymph node sampling and 94 (35%) entailed hilar/interlobar lymph node sampling. The right paratracheal and subcarinal lymph nodes were the most commonly sampled, and accounted for 59% and 61% of all such sampling, respectively (Fig 1). The size of the lymph nodes sampled had a mean diameter of 1.36 cm (SD, 0.61 cm). Before their EBUS-TBNA procedure, results from PET were only available in 122 (45%) of the patients. Among the group with non-diagnostic EBUS-TBNA findings, there were 30 patients who underwent subsequent surgical exploration, and three had a second EBUS-TBNA. The remaining 60 patients were clinically followed up only. Among the latter, there were 33 with no evidence of malignancy during a mean follow-up duration of 1.7 (SD, 1.3) years (Fig 2).
Diagnostic performance

The diagnostic performance of these 269 EBUS-TBNA procedures resulted in a sensitivity, specificity, negative predictive value, and accuracy of 87% (173/[173+25]), 100%, 74% (71/[71+25]), and 91% ([173+71]/269), respectively. In the 220 procedures yielding a final diagnosis of malignancy, the respective sensitivity and negative predictive values were 89% (162/[162+20]) and 66% (38/[38+20]). Regarding the lesions that turned out to be infections (10 cases of tuberculosis and 1 of cryptococcosis), though lung cancer was suspected initially, the sensitivity and negative predictive values were 75% (6/[6+2]) and 60% (3/[3+2]), respectively. Regarding EBUS-TBNA used as the first diagnostic test (ie for tissue sampling), there were 158 procedures that yielded sensitivity and negative predictive values of 90% (100/[100+11]) and 81% (47/[47+11]), respectively (Table 3).

In all, 95 (37%) of the patients had a pre-existing malignancy diagnosed with a median of 3.8 years before the EBUS-TBNA. In 27 (28%) of these individuals, the pathology revealed by the procedure was other than that of the pre-existing one.

Molecular profiling and histological classification

When EBUS-TBNA was introduced locally in 2006, molecular profiling such as for EGFR and anaplastic lymphoma kinase (ALK) was not widely accepted as standard practice. It was not until October 2008 that molecular profiling was included as routine when adenocarcinoma was diagnosed by EBUS-TBNA. Subgroup analysis was therefore performed to determine whether EBUS-TBNA was a reliable diagnostic tool for retrieving adequate tissue for EGFR or ALK mutation tests. Among 40 patients with samples sent for EGFR and/or ALK, 38 (95%) were found to be adequate. Among these, 24 (60%) harboured mutations in the *EGFR* gene exons 18-21 as detected by polymerase chain reaction and DNA sequencing and 13 (33%) were negative (wild-type), and one had the ALK translocation.

Apart from the origin of the tumour, it is now considered imperative to know the cell type within non–small-cell lung cancer (NSCLC) as chemotherapy may then be individualised. Of the 158 patients confirmed to have malignancy by EBUS-TBNA, only 22 (14%) were revealed to be NSCLC without information about the exact cell type, differentiation of the tumour, EGFR status, or primary origin of the tumour.

Complications

Regarding all 269 procedures performed, there were no significant complications (bleeding, pneumothorax, infection, or respiratory distress) warranting reversal of sedation or any other intervention. All the patients, including 14 with superior vena caval obstruction and 1 with severe heart failure (left ventricular ejection fraction 20%), tolerated the procedures well and no procedure was prematurely terminated.

Discussion

Most patients with lung cancer are in locally advanced or metastatic stages upon diagnosis and proper staging is important in classifying patients into operable or inoperable status.\textsuperscript{10,11} In the absence of distant metastases, mediastinal lymph node staging remains the most important factor determining overall staging. Sampling of mediastinal lymph nodes is therefore important in both diagnosis and staging, which should guide subsequent treatment. The term ‘personalised therapy’ is coined in contrast with that in the past, when treatment for lung cancer was mainly classified into small-cell lung cancer or NSCLC only. Nowadays, among advanced stages of NSCLC, oral targeted therapy (EGFR tyrosine kinase inhibitor) is considered the preferred first-line treatment in the presence of EGFR mutations in exon 19 and 21.\textsuperscript{12} Falling short of this, specific anti-cancer therapeutic options (chemotherapy and anti-angiogenic agents) are currently decided

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<th>TABLE 3. Diagnostic yield of endobronchial ultrasound-guided transbronchial needle aspiration (EBUS-TBNA)</th>
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<td>EBUS-TBNA procedure</td>
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<td>All procedures</td>
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<td>Procedures with a final diagnosis of malignancy</td>
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<td>Patients with a final diagnosis of infective cause</td>
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* CI denotes confidence interval
according to the carcinoma cell types, namely: small, large, adeno, or squamous. In patients with pre-existing malignancy, subsequent treatment is largely dictated by differentiating metastatic disease from primary carcinoma of the lung. Based on our 4-year experience since 2006, EBUS-TBNA has demonstrated high diagnostic accuracy in providing clear-cut histological classification and molecular profiling that facilitated state-of-the-art treatment for lung cancer.

Previously, surgical intervention such as mediastinoscopy was the only diagnostic modality with access to the mediastinal lymph nodes for tissue sampling, though the hilar area remained a sanctuary site. However, surgical exploration is not usually considered the upfront diagnostic investigation, due to the need for general anaesthesia for which most patients with advanced lung cancer are not good candidates. Recourse to EBUS-TBNA opens a new diagnostic paradigm for patients having mediastinal or hilar lymphadenopathy. In our study, its diagnostic yield was close to 90%, which is similar to most clinical series reported worldwide. A negative result from EBUS-TBNA (with the negative predictive value of 74% only), however, warrants further investigation and may be best explained by the sampling error associated with a fine needle.

The advantage of using EBUS-TBNA with a 22-G fine needle for sampling under real-time guidance is that it minimises inadvertent puncturing of blood vessels by the Doppler mode screening. This was exemplified by the excellent safety demonstrated in our series. Regrettably, ROSE was not available in our centre, but has the potential to reduce the number of needle passes and sites sampled per patient, and hence the complication rate. The procedural time was not recorded but was estimated to be around 20 to 50 minutes, depending on the number of lesions sampled. Recently, we have shown that EBUS-TBNA was safe even in patients with superior vena caval obstruction. Using the fine needle does not jeopardise securing adequate pathological data required for formulating personalised treatment. Of the 210 patients confirmed as having malignancy as the final diagnosis, EBUS-TBNA was able to detect extrathoracic malignancy in 23 (11%) and primary lung cancer in 135 (64%). Among those 135 patients with confirmed primary lung cancer, 118 (87%) had the exact cell type delineated, although this could have been confounded by concomitant changes in the pathological classification and management of NSCLC during the study period. For the 40 patients who had molecular profiling, adequate tissues for EGFR or ALK mutation tests were obtained in 38 (95%), which was much higher than the 36% reported in a recent landmark clinical trial on targeted therapy (IPASS study) using conventional diagnostic methods. Moreover, EBUS-TBNA is useful in the diagnosis of synchronous lung cancers and other benign conditions affecting the mediastinum such as tuberculosis, fungal infection, or sarcoidosis.

Endoscopic ultrasound-guided fine-needle aspiration (EUS-FNA), via the trans-oesophageal approach, was developed earlier than EBUS-TBNA and is also reported to have high diagnostic performance in mediastinal lymphadenopathy. However, EUS-FNA provides limited access to the hilum and the right side of the mediastinum. Although the inferior mediastinal nodal stations such as para-oesophageal (below the carina) and pulmonary ligament can be approached by EUS-FNA but not EBUS-TBNA, metastatic involvement in these areas is relatively uncommon. In a local study, it only accounted for about 6% of lymph nodes sampled by EUS-FNA. Nevertheless, the combination of EBUS-TBNA and EUS-FNA has even better diagnostic performance than either procedure alone. However, whether this combined approach can replace mediastinoscopy is yet to be elucidated.

In summary, EBUS-TBNA has been shown to have high diagnostic performance and excellent safety. Similar to other technical procedures, it requires a defined period of training. Due to the steep learning curve at the beginning, some professional bodies have recommended a minimum requirement of 25 procedures be performed annually to maintain the technical skill for EBUS proficiency. The diagnostic yield was also associated with annual hospital EBUS-TBNA volume and reflects the combined team experience, and also involves anaesthetic use and cytopathological interpretation.

In Hong Kong, EBUS-TBNA has been introduced for the past 7 years, and has emerged as an important tool in the field of interventional pulmonology, which is rapidly evolving worldwide. The management of lung cancer is greatly advanced with all these new diagnostic and therapeutic modalities in the pipeline, underscoring the importance of personalised therapy, which is the new paradigm.

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References


