Diagnosis of Peripheral Lung Lesions by EBUS-Guided TBB in Routine Practice

Diagnostik peripherer pulmonaler Läsionen mittels EBUS-geführter transbronchialer Biopsie (TBB) in der täglichen Praxis

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Abstract

Objective: To determine the diagnostic yield of EBUS guided TBB performed in routine practice with flexible bronchoscopy and under moderate sedation in ambulatory and hospitalized patients.

Methods: Bronchoscopy was performed under standard conditions in ambulatory and hospitalized patients. Bronchoscopically invisible peripheral pulmonary lesions were located with 20 MHZ-EBUS-probe and transbronchial biopsy was taken using a guiding sheath. Fluoroscopy was additionally performed as required to identify the lesion.

Results: 257 patients with peripheral pulmonary lesions were investigated, with malignancy in 70% of those with a diagnosis established. 175/257 (68.1 %) of lesions were detected with EBUS. In 139/176 (79.4 %) of these lesions, TBB enabled a final diagnosis. The TBB yield depended on lesion size. It was 61.3 % in lesions ≤ 20 mm, 85.5 % > 20 mm/≤ 30 mm, and 81.2 % in ≥ 30 mm (p< 0.0001). This yield was also affected by the position of the probe (centrally 84.5%, tangentially 57.6 %, p= 0.01)). Operator experience did not influence the diagnostic yield but considerably shortened investigation time (4.9 ± 3.5 vs. 6.2 ± 4.2 min, p = 0.042). Relevant complications occurred in only 1.9 % (3 cases of postinterventional pneumothorax).

Conclusions: In an unselected population, EBUS-guided TBB has a high diagnostic yield in peripheral lesions > 20 mm whereas its yield decreases considerably in smaller lesions. Complications are very rare. EBUS-guided TBB can successfully and safely be performed by flexible bronchoscopy.

Zusammenfassung

Fragstellung: Diese Arbeit untersucht die diagnostische Ausbeute der EBUS-orientierten transbronchialen Biopsie (TBB) mittels flexibler Bronchoskopie unter Routinebedingungen bei ambulanten und stationären Patienten.


Ergebnisse: Insgesamt wurden 257 Patienten mit peripheren pulmonalen Rundherden untersucht, davon erwiesen sich 70 % als maligne. 175/257 (68,1 %) der Herde wurden mittels EBUS lokализiert. In 139/176 (79,4 %) wurden diese Herde mittels TBB histologisch diagnostiziert. Die Ausbeute der TBB war abhängig von der Herdgröße: sie betrug 61,3 % in Herden von einer Größe ≤ 20 mm, 85,5 % > 20/≤ 30 mm und 81,2 % bei Herdgrößen ≥ 30 mm (p< 0.0001). Die Ausbeute hing zusätzlich ab von der Lage der Herde (zentral 84,5 %, peripher 57,6 %, p= 0.01). Die Erfahrung des Untersuchers hatte keinen Einfluss auf die Ausbeute, wohl aber ging eine größere Erfahrung mit kürzeren Untersuchungszeiten einher (4,9 ± 3,5 vs. 6,2 ± 4,2 min, p = 0,042). Relevante Komplikationen traten nur in 1,9 % der Fälle auf (3 Fälle eines Pneumothorax).

Introduction
EBUS-guided TBB is an established procedure to diagnose bronchoscopically invisible, peripheral pulmonary lesions for more than 10 years. Nevertheless, available data are quite heterogeneous and poorly comparable. Obviously, the diagnostic success depends on the size of pulmonary lesion. However, there are remarkable differences in diagnostic yield especially in small lesions < 20 mm. While some studies describe high diagnostic yields of about 60–80% independent of lesion-size [1–8], other authors report a dramatic decrease in diagnostic yield in lesions < 20 mm [9–11]. Furthermore, the procedures used in different trials vary widely. In some studies, bronchoscopy is performed in general anaesthesia [1,5,8,10] whereas in others moderate sedation is used. Also support in navigation to the lesion differs (virtual bronchoscopy, fluoroscopy, use of a curette, electromagnetic navigation). Another important issue is the additional analysis of cytology sampled by brush or suction catheter, even when the lesion was not reached with the EBUS-probe.

The aim of this study was to investigate the diagnostic yield of EBUS guided TBB of peripheral pulmonary lesions by flexible bronchoscopy in routine practice applying a moderate sedation protocol. In addition, factors affecting the diagnostic yield and limitations of this method were analyzed.

Methods

Patient population
Overall, 257 patients (179 male, 78 female, mean age 68±12 years) with peripheral pulmonary lesions diagnosed from January 2006 to September 2008 were investigated. Patients were examined either as outpatients (n=86, 33%) or during their hospital stay (n=171, 67%). Data were retrieved retrospectively in a computer database; therefore, informed consent was not mandatory.

Lesions
Most lesions were solid peripheral lesions. In addition, 18 (7%) infiltrative lesions were included. The mean diameter was 31.5±15.3 mm.

Bronchoscopy
All patients were investigated by flexible bronchoscopy. The standardized examination included a) a CT scan of the lung prior to bronchoscopy; b) a sedation protocol with midazolam and/or propofol; c) an EBUS guided TBB; and d) a chest radiograph one hour after bronchoscopy. Flexible bronchoscopy was performed in sedation with a mean dose of 4.5±1.7 mg midazolam. In 14% an additional administration of propofol was necessary (mean dose: 66±33.5 mg). In no case general anaesthesia was applied.

For most examinations a videobronchoscope with 3 mm or 2.8 mm working-channel was used (Olympus BF-1-T-180 and BF-1-T 160). If the lesion was not accessible with this bronchoscope (e.g. because of insufficient angulation of the scope to the apical left bronchus (LB1)), bronchoscopes with smaller diameters were used (Olympus BF-P180 or XBF 4 B40).

After a thorough examination of the bronchial mucosa within view and exclusion of endobronchially visible lesions, endobronchial ultrasonography with 20 MHz radial probe within a 2.6 mm diameter guide sheath was placed into the lesion (a 2.0 mm guide sheath when the small-diameter-bronchoscopes were used). After localisation of the peripheral lesions the ultrasonography-probe was withdrawn and a biopsy forceps (disposable biopsy forceps Olympus: FB 231 D/FB 233 D) was placed in the lesion using the guide sheath. At least four transbronchial biopsies were taken. The transbronchial biopsies were analysed histologically and by imprint-cytology. If the peripheral lesion was not visualized by EBUS, x-ray-fluoroscopy was used to assist correct placement of the EBUS-probe into the lesion. If the lesion was not reached with EBUS and additional fluoroscopy, the examination was terminated. A typical example of an EBUS-guided TBB is shown in Fig. 1.

Patients with a nondiagnostic bronchoscopy were investigated by computertomography-guided transcutaneous biopsy or surgery.

Final diagnoses
Final diagnoses of patients were retrieved from the charts of the patients and classified as malignant or non-malignant. Overall, 25 patients were lost to follow up.

Statistical analysis
Categorical variables were described by frequencies and percentages and continuous variables by means and standard deviations. Categorical variables were compared with the chi-square
Results

Final diagnoses

A final diagnosis could be established in 232 patients (90.3%). In 163 cases (63.4% of all, 70.3% of those with a diagnosis established), the final diagnosis was a malignancy. In this group, non-small cell lung cancer was the most frequent diagnosis. In 69 cases, (26.8% of all, 29.7% of those with a diagnosis established) a non-malignant diagnosis was made (Table 1).

Detection of lesions

In 175 of 257 patients (68.1%), lesions could be visualized by EBUS. Of these 175 lesions, 33 (18.9%) were imaged tangentially only. The overall investigation time in visualized lesions was 5.2 min ± 3.7 min. In 136 (77.3%) cases, x-ray-fluoroscopy was used additionally. The mean fluoroscopy-time was 16.9 ± 24.2 s.

Diagnostic yield

Overall, the diagnostic yield of EBUS-TBB was 139 of 257 patients (54.1%). Of those with EBUS-detected lesions and transbronchial biopsies, the diagnosis could be established in 139 of 175 (79.4%) patients. In 17 of 175 transbronchial biopsies (9.7%), diagnosis was exclusively made by the imprint-cytology.

The diagnostic yield depended on the size of the lesion. Both the detection-rate with EBUS and a diagnostic TBB were related to the size of the lesion (<20 mm, >20 mm/≤30 mm, >30 mm, respectively). The overall diagnostic yield was 24.4%, 64.6% and 69.1% in lesions ≤20 mm, >20 mm/≤30 mm and >30 mm, respectively (p <0.0001).

Factors affecting the diagnostic yield

The EBUS detection rate depended on the size of the lesions (39.7% of lesions ≤20 mm vs. 75.6% of lesions >20 mm/≤30 mm vs. 84.55% of lesions >30 mm) (p <0.0001).

The yield of TBB in patients with a lesion localized with EBUS was 61.3%, 85.5%, and 81.7% in lesions ≤20 mm, >20 mm/≤30 mm and >30 mm, respectively (p <0.0001).

The overall diagnostic yield was 24.4%, 64.6% and 69.1% in lesions ≤20 mm, >20 mm/≤30 mm and >30 mm, respectively (p <0.0001).

Examination times

The mean EBUS-time from first introduction of EBUS-probe up to termination of the EBUS-examination was 5.23 min ± 3.7 min. The EBUS-time for very experienced operators was shorter than EBUS-time for less experienced operators (4.9 ± 3.5 vs. 6.2 ± 4.2 min, p = 0.042).

Complications

Relevant complications were rare. Postinterventional pneumothoraces occurred in 5 cases (1.9%). Three pneumothoraces resolved with oxygen-therapy. In two cases, insertion of a chest drain was necessary. Relevant bleeding requiring further intervention like instillation of noradrenaline occurred in two cases. Bleeding was terminated just with holding the guiding sheath in position after TBB or instillation of arterenol. Coughing affecting the progress of examination occurred in 22 cases (8.5%).

Discussion

The main findings of our study are the following: 1) in an unselected population with malignancy in 70% of those with a diagnosis established, flexible bronchoscopy with EBUS guided TBB had a high diagnostic yield (54.1% overall, 79.4% if detected by EBUS);

<table>
<thead>
<tr>
<th>Lesions detected with EBUS</th>
<th>TBB diagnostic</th>
<th>Diagnostic yield</th>
</tr>
</thead>
<tbody>
<tr>
<td>All</td>
<td>175/257</td>
<td>68.1%</td>
</tr>
<tr>
<td>&gt;30 mm</td>
<td>82/97</td>
<td>84.5%</td>
</tr>
<tr>
<td>&gt;20 mm, ≤30 mm</td>
<td>62/82</td>
<td>75.6%</td>
</tr>
<tr>
<td>≤20 mm</td>
<td>31/78</td>
<td>39.7%</td>
</tr>
<tr>
<td>operator with less EBUS-experience</td>
<td>46/65</td>
<td>70.8%</td>
</tr>
<tr>
<td>tangential lesion detection</td>
<td>19/33</td>
<td>57.6%</td>
</tr>
<tr>
<td>central lesion detection</td>
<td>120/142</td>
<td>84.5%</td>
</tr>
<tr>
<td>infiltrative lesion</td>
<td>17/18</td>
<td>94.4%</td>
</tr>
</tbody>
</table>
2) the diagnostic yield was strongly dependent on the lesion size, with 20 mm as threshold; 3) an additional factor affecting the diagnostic yield was the positioning of the EBUS-probe in the lesion (centrally/tangentially); 4) experience of the operator did not affect the diagnostic yield but determined the duration of the procedure; 5) complications were very rare.

Up to now, 13 studies have been published evaluating EBUS-guided TBB, including 8 prospective studies. Only four of these studies included more than 100 patients, and 8 studies (61.5%) were contributed by two groups from Germany [1, 5, 8, 10] and Japan [2, 4, 7, 11] (Table 3). In addition, a systematic review and meta-analysis has been published recently [12]. The EBUS-detection rate was reported in 11 studies and ranged between 67 and 95% [1–5, 7, 9–11, 13, 14]. The overall diagnostic yield was generally high. A threshold-dependent analysis of the diagnostic yield was provided in 10 studies. Only three report a threshold of 20 mm determining significant differences in the diagnostic yield [3, 9, 11]. Others found no remarkable threshold [1, 5, 8], a higher threshold of 30 mm [2, 7] or a lower 15 mm [14]. These results are only in part comparable to ours due to a wide variation in study protocols, methodology, and patient populations studied. Of note, only few studies were performed using flexible bronchoscopy only [3, 9]. Two studies used additional techniques to detect peripheral lesions such as virtual bronchoscopy and bronchoscopic navigation [8, 11].

Our observational data are characterized by the following: 1) they were generated in routine practice in a large specialized department of pulmonary medicine; 2) only flexible bronchoscopy was performed; 3) they represent the largest series of patients investigated by EBUS-guided TBB. Compared to previous series, the overall diagnostic yield in an unselected population of 54.1% is moderate but generally in line with others reporting overall yields of 53 to 80%. Clearly, the yield of lesions detected by EBUS-guided TBB depends on the size of the lesion. Remarkable variations in diagnostic yields in lesions <20 mm have been reported, some clearly lower than ours (57% compared to 18%–46% [7, 9, 10, 11], some comparable (53–55% [3, 4], some clearly higher (71–76%) [2, 6, 8, 14]. For lesions ≥20 mm, most groups report yields of 57 to 80–90% [2–4, 6, 9, 11, 14]. Our TBB yield of 79.4% is in the upper range of these numbers. Several reasons may account for these differences which apparently are higher in small lesions: 1) two groups identified the position of the probe within or adjacent to the lesion as independent predictors of the diagnostic yield in multivariate analysis [13, 14]. This finding is confirmed by our data showing a difference in the yield of 84.5% for centrally versus 57.6% for tangentially detected versions; 2) one group identified the position of the lesion in CT scan as additional independent predictive factor [13]. This appears plausible in view of several bronchial areas difficult to reach with bronchoscopy, however, we were not able to investigate CT scans retrospectively; 3) the number of biopsies taken was reported to affect the yield, with five biopsies necessary to achieve the optimal yield [14]. We took at least four biopsies, suggesting that the fifth may not be mandatory in every case. However, we cannot provide a formal analysis about the optimal number of biopsies to be retrieved; 4) the lower limit of lesions used to investigate patients with EBUS-guided TBB has not been defined. As such, <20 mm may therefore mean at least 10 mm or even less. Only two studies have formally evaluated the diagnostic yield of EBUS-guided TBB in lesions <15 mm and <10 mm, resulting in yields of 40% and 76%, respectively [2, 14]. Even so, it is largely up to the decision of the investigator to exclude small lesions in areas difficult to reach with the bronchoscope. Other potential factors cannot be related to different yields. Fluoroscopy was used additionally in three quarters of cases, however, fluoroscopy times were generally very short. With increasing experience, we felt fluoroscopy increasingly dispensable. In fact, studies evaluating EBUS-guided TBB without fluoroscopy found comparable yields [3, 10]. An exceptionally EBUS-experienced operator obviously does not necessarily produce better results, but his examination times are shorter [14]. This is plausible, since the method is very easy to apply. In addition the ultrasound image of a lesion is instantly recognizable as opposed to the intricate ultrasound patterns of central peribronchial structures. The learning curve for EBUS-guided TBBS is extremely steep. In our series, all operators had several years experience in bronchoscopy and in all examinations a very experienced EBUS-operator was present for purposes of teaching. The prevalence of malignancy was identified as a possible confounder in a recent meta-analysis [12]; however, underlying disease was not found predictive in a multivariate analysis [13]. Finally, lobar distribution and presence of complications were not predictive [13, 14].

In a recent systematic review and meta-analysis including 16 studies, the reported diagnostic potential for the detection of lung cancer was addressed. EBUS-guided TBB had a point specificity of 1.00 (95% CI 0.99–1.00) and point sensitivity of 0.73 (95% CI 0.70–0.76), with a positive likelihood ratio of 26.84 (12.60–57.20) and a negative likelihood ratio of 0.28 (0.23–0.36) [12].

While the diagnostic yield achieved is already impressive, there may be room for improvement. Two approaches may be useful in this regard. First, to navigate EBUS-probe to the lesion some operators use a ‘curette’ to change direction under fluoroscopy. This method increases examination-time considerably because there is no direct view on the bronchial structures and the navigation results in a kind of “trial and error”. In fluoroscopically invisible lesion this method is not practical. A new navigation-tool is the electromagnetic navigation (EN). This method even allows navigation also to a fluoroscopically invisible lesion. In combination with EBUS to verify location of the probe in the lesion, the diagnostic yield in lesions <20 mm can be increased [8]. However, this approach implies increased costs and prolonged examination time with deeper sedation or even need of general anaesthesia. The same is true for a recently described bronchoscope insertion guidance system [15].

Second, in order to optimize the diagnostic success in several studies cytology sampled by brush or suction catheter was used additional to TBB, or without TBB when lesion was not reached. This method increases the diagnostic yield about 10–20%, although several authors do not report how many diagnoses just rely on cytology without reaching the lesion with EBUS. Especially in lesions <20 mm with lower detection rates the waiver of this additional diagnostic tool may explain lower diagnostic success in this study. However, the diagnostic validity of cytological diagnosis compared with histological diagnosis is not completely clear. Newer studies using EBUS-TBB without another navigation-tool and without cytology show an even lower diagnostic yield over all in lesions <20 mm of 46% [10]. EBUS-guided TBNA was recently shown to increase the diagnostic yield from 60.6% for EBUS-guided TBB versus 78.4% for this procedure plus EBUS-guided TBNA [16].
<table>
<thead>
<tr>
<th>Author/year</th>
<th>Study design</th>
<th>Technique</th>
<th>Patients</th>
<th>Size</th>
<th>EBUS detection-rate</th>
<th>Diagnostic yield over all</th>
<th>Pneumothorax</th>
</tr>
</thead>
<tbody>
<tr>
<td>Eberhardt R, 2009</td>
<td>prospective case series</td>
<td>EBUS</td>
<td>100</td>
<td>≤ 20 mm</td>
<td>67%</td>
<td>46%</td>
<td>3%</td>
</tr>
<tr>
<td>Huang CT, 2009</td>
<td>retrospective</td>
<td>EBUS</td>
<td>83</td>
<td>72%</td>
<td>53%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yamada, 2007</td>
<td>retrospective</td>
<td>EBUS</td>
<td>155</td>
<td>&lt; 15 mm</td>
<td>69%</td>
<td>5%</td>
<td></td>
</tr>
<tr>
<td>Eberhardt R, 2007</td>
<td>prospective rand. cont. trial</td>
<td>EBUS</td>
<td>39</td>
<td>&lt; 20 mm</td>
<td>69%</td>
<td>5%</td>
<td></td>
</tr>
<tr>
<td>EN</td>
<td>39</td>
<td>&lt; 20 mm</td>
<td>59%</td>
<td>5%</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>EBUS + EN</td>
<td>40</td>
<td>&lt; 20 mm</td>
<td>88%</td>
<td>8%</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yoshikawa, 2007</td>
<td>prospective case series</td>
<td>EBUS</td>
<td>76</td>
<td>&gt; 30 mm</td>
<td>62%</td>
<td>90%</td>
<td>83%</td>
</tr>
<tr>
<td>Dooms C, 2007</td>
<td>prospective case series</td>
<td>EBUS</td>
<td>50</td>
<td>&gt; 20 mm</td>
<td>68%</td>
<td>68%</td>
<td></td>
</tr>
<tr>
<td>Herth F, 2006</td>
<td>prospective case series</td>
<td>EBUS</td>
<td>54</td>
<td>&gt; 30 mm</td>
<td>70%</td>
<td>1.9%</td>
<td></td>
</tr>
<tr>
<td>Paone G, 2005</td>
<td>prospective rand. cont. trial</td>
<td>EBUS</td>
<td>87</td>
<td>&gt; 30 mm</td>
<td>79%</td>
<td>75%</td>
<td>71%</td>
</tr>
<tr>
<td>Asahina, 2005</td>
<td>pilot + study</td>
<td>EBUS + virt. bronchosc.</td>
<td>30</td>
<td>&gt; 20 mm</td>
<td>63%</td>
<td>44%</td>
<td>0%</td>
</tr>
<tr>
<td>Kurimoto, 2004</td>
<td>prospective</td>
<td>EBUS</td>
<td>150</td>
<td>&lt; 10 mm</td>
<td>77%</td>
<td>57%</td>
<td></td>
</tr>
<tr>
<td>Yang, 2004</td>
<td>retrospective</td>
<td>EBUS</td>
<td>122</td>
<td>&gt; 20 mm</td>
<td>93%</td>
<td>66%</td>
<td>5%</td>
</tr>
<tr>
<td>Kikuchi, 2004</td>
<td></td>
<td>EBUS</td>
<td>24</td>
<td>&lt; 20 mm</td>
<td>79%</td>
<td>58%</td>
<td>4.2%</td>
</tr>
<tr>
<td>Herth, 2002</td>
<td>prospective case series</td>
<td>EBUS</td>
<td>50</td>
<td>&gt; 30 mm</td>
<td>92%</td>
<td>2%</td>
<td></td>
</tr>
</tbody>
</table>
Although applying EBUS after measuring and applying the distance between the orifice of the bronchus and the lesion has been reported to increase the yield of EBUS-guided TBB, the yield achieved (79.9% versus 57.1%) apparently was not considerably higher than the yield reported by most of groups including that of the present study [17]. The strength of our study is the large number of unselected patients included compared with previous EBUS-studies in a real life setting. Moreover, we used exclusively flexible bronchoscopy obviating the need for general anesthesia and included a considerable amount of outpatients. The complication rate was exceedingly low, comparing favourably to previous reports [4, 8], proving the feasibility of this diagnostic technique in daily practice. Although the study-design was retrospective, data of EBUS-examinations of peripheral pulmonary lesions were collected for internal quality-control. This clearly improves the confidence in these data.

In conclusion, EBUS-guided TBB of peripheral pulmonary lesions is a safe and successful method which can be established in short examination-time within a short examination-time and without X-ray fluoroscopy. Chest 2006; 129: 147 – 150

Conflict of interest

The authors have no conflict of interest.

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