Endobronchial ultrasound in diagnosing and staging of lung cancer by Acquire 22G TBNB versus regular 22G TBNA needles: study protocol of a randomised clinical trial

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ABSTRACT
Introduction Accurate diagnosis and staging of lung cancer is crucial because it directs treatment and prognosis. Endobronchial ultrasound-guided transbronchial needle aspiration (EBUS-TBNA) and endoscopic ultrasound with bronchoscope fine-needle aspiration (EUS-B-FNA) are important in this process by sampling hilar/mediastinal lymph nodes and centrally located lung tumours. With the upcoming of immunotherapy and targeted therapies, assessment of programmed death ligand 1 (PD-L1) expression and molecular profiling has become important but is often impossible in cytological samples obtained through standard 22G TBNA needles. Recently, a three-pronged cutting edge 22G needle was developed that allows for transbronchial needle biopsy (TBNB). Our objective is to determine if EBUS/EUS-B-guided nodal/lung tumour sampling with Acquire 22G TBNB needles results in an improved suitability rate for the assessment of PD-L1 expression in comparison to standard 22G TBNA needles in patients with a final diagnosis of lung cancer.

Methods and analysis This is an investigator-initiated, parallel group randomised clinical trial. Patients are recruited at respiratory medicine outpatient clinics of participating university and general hospitals in the Netherlands, Poland and Italy. In total 158 adult patients with (suspected) lung cancer are included if they have an indication for mediastinal/hilar lymph node or lung tumour sampling by EBUS-TBNA and/or EUS-B-FNA based on current clinical guidelines. Web-based randomisation between the two needles will be performed. Samples obtained from mediastinal/hilar lymph nodes and/or primary tumour will be processed for cytology smears and cell block analysis and reviewed by blinded reference pathologists. An intention-to-treat analysis will be applied. Patients with missing data will be excluded from analysis for that specific variable but included in the analysis of other variables. This study is financially supported by Boston Scientific.

Ethics and dissemination The study was approved by the local Ethics Committee (Medisch Ethische Toetsingscommissie Amsterdam Medical Center (AMC)). Dissemination will involve publication in a peer-reviewed biomedical journal.

Strengths and limitations of this study

The international, multicentre design will improve generalisability of the findings. Two blinded reference pathologists will analyse the tissue samples in order to objectively compare the two needles. The Acquire 22G transbronchial needle biopsy (TBNB) needle is commercially available and therefore, our findings can directly be implemented in clinical practice. Performing bronchoscopists cannot be blinded to needle randomisation which is a potential source of performance bias. This study is not powered to assess if the Acquire 22G TBNB results in an improved diagnostic yield compared with the 22G transbronchial needle aspiration.

Trial registration number NL7701; Pre-results.

INTRODUCTION
Lung cancer is the most commonly diagnosed cancer worldwide with the highest mortality of all cancers.1 Accurate staging is important because it directs treatment and prognosis.2 3 Mediastinal and hilar staging is key in this process. For decades, this was done by (cervical) mediastinoscopy which is a costly and invasive procedure with suboptimal sensitivity for mediastinal metastases.4 Over the past 15 years, clinicians have started to use endobronchial ultrasound-guided transbronchial needle aspiration (EBUS-TBNA) for this purpose which is less invasive and has a similar diagnostic range to cervical mediastinoscopy.5 Bronchial EBUS-TBNA can be combined with endoscopic ultrasound fine-needle aspiration (EUS-FNA) from the
It has been shown that a combined endosonography investigation (EBUS and EUS), followed by mediastinoscopy in the absence of metastases at endosonography, results in greater sensitivity for mediastinal nodal metastases compared with mediastinoscopy alone.\(^5\) Therefore, current guidelines on lung cancer staging recommend endosonography (EBUS and/or EUS) as the initial tissue sampling technique for mediastinal nodal staging.\(^2\) Additionally, EBUS and EUS can be used to sample centrally located lung tumours.\(^7,9\)

Over the past years, the treatment options for patients with advanced lung cancer have been expanding rapidly. Encouragingly, targeted immunotherapies against programmed death ligand 1 (PD-L1) (eg, durvalumab and atezolizumab) and its receptor PD-1 (eg, nivolumab and pembrolizumab) have improved the survival in selected patients with advanced lung cancer. Therefore, to be able to assign a patient to optimal treatment, assessment of PD-L1 expression and molecular profiling of the tumour is crucial. With the advent of EBUS-TBNA and EUS-FNA, the ability to accurately assess the PD-L1 expression and molecular profiling on small, cytological tissue samples has become of great importance. Although the use of cell blocks and improved sequencing techniques have expanded the possibilities for PD-L1 assessment,\(^10\) the success rate of PD-L1 expression and molecular profiling remains suboptimal at cytology compared with histology.\(^15,17\)

Recently, a three-plane symmetric needle with Fransen geometry (Acquire 22G transbronchial needle biopsy (TBNB)) was developed (figure 1). In pancreatic cysts, the Acquire 22G needle allowed improved, true histological core tissue acquisition in pancreatic cysts.\(^18,19\) To date, there is no literature about the Acquire 22G TBNB needle for diagnostic and staging purposes during EBUS/endo- scopic ultrasound with bronchoscope (EUS-B) procedures in patients with (suspected) lung cancer.

In this parallel group randomised clinical trial, we hypothesise that EBUS/EUS-B sampling of mediastinal/hilar lymph nodes and/or primary lung tumours with the Acquire 22G TBNB needle has a higher suitability rate for the assessment of PD-L1 expression in comparison to the regular 22G TBNA needle in patients with lung cancer.

**METHODS AND ANALYSIS**

**Design and study dates**

This is a protocol of an investigator-initiated parallel-group randomised clinical trial. The trial is executed in four centres including both university and general hospitals in The Netherlands (Amsterdam University Medical Centers and Leiden University Medical Center), Italy (Fondazione Pollicino Universitario Agostino Gemelli, Rome) and Poland (Pulmonary Hospital Zakopane). We expect that we will complete inclusion in June 2022.

**Study population**

Patients will be recruited at outpatient clinics of respiratory medicine of the participating hospitals by their treating physicians. If the patient is willing to participate in the study, information about the study will be provided by the local investigator, who will then gain written informed consent. Eligible are patients with proven or suspected lung cancer (either non-small-cell lung cancer or small-cell lung cancer), who have an indication for mediastinal/hilar lymph node or lung tumour sampling by EBUS-TBNA and/or EUS-B-FNA according to current clinical guidelines.\(^2\) Indications for mediastinal/hilar lymph node staging are: (1) suspicion of mediastinal or hilar lymph node metastases based either on size (short axis $>$10 mm on CT) or increased fluorodeoxyglucose (FDG) uptake, (2) a primary tumour without FDG uptake, (3) a primary tumour with a size of $\geq$3 cm or (4) a centrally located primary tumour. Sampling of centrally located lung tumours is attempted if the intrapulmonary mass is located adjacent to the oesophagus or airway wall but not endobronchially visible with conventional bronchoscopy. The following exclusion criteria will be applied: (1) mediastinal restaging after neoadjuvant treatment, (2) contraindication for EBUS or EUS-B (eg, severe respiratory insufficiency), (3) a non-correctable coagulation disorder, (4) pregnancy or (5) inability to consent.

**Study withdrawal**

Subjects can leave the study at any time for any reason if they wish to do so. The investigator can decide to withdraw a subject from the study for unanticipated urgent medical reasons. Such withdrawals will be monitored and reported in the final study report.

**Randomisation and needle allocation**

Randomisation will be performed prior to endosonography. Participants will be randomised to either the Boston Scientific Acquire 22G TBNB needle or the regular Boston Scientific 22G TBNA needle (figure 2). Both needles have a Conformité Européenne (CE) mark. We will use a web-based block-randomisation...
(using Castor Electronic Data Capture (EDC) software, Amsterdam, the Netherlands), stratified by participating hospital to ensure that both groups are of equal size for every hospital. Endoscopists will not be blinded to needle randomisation but the reference pathologists will be blinded for the procurement of the tissue.

Endosonographic procedure

Endosonography will be performed at the department of pulmonary medicine of participating hospitals by experienced chest physicians who are specifically trained in EBUS and EUS-B. Procedures will be performed according to institutional practice, mostly on an outpatient basis. Topical anaesthesia is applied to the pharynx, larynx, trachea and bronchi. Sedation will be administered according to institutional practice, mostly conscious sedation by midazolam 2.5–5 mg intravenous with or without fentanyl or propofol sedation. Vital parameters will be monitored during and after the procedure. Patients will be observed for 1 hour after the endosonography procedure.

Procedures will be performed using an ultrasound bronchoscope with a linear scanning transducer (EBUS scope). A dedicated ultrasound processor is used for imaging processing with Doppler flow imaging for the detection of blood vessels and that enables real-time ultrasound guidance for sampling. All patients will undergo a systematic endosonographic evaluation of all the accessible mediastinal/hilar lymph nodes and the tumour itself.2 20 Following the endosonographic inspection, lymph nodes will be sampled with the randomised needle from N3 to N2 to N1 to the tumour itself. A minimum of two needle passes will be performed for each lymph node and tumour. After insertion of the needle in the target lymph node or tumour, the stylet will be pulled out slowly (slow pull technique). In case of no or limited aspirate after one needle pass, suction using a 10 mL syringe during the following aspirations will be applied.

Endoscopist satisfaction

After the endoscopic procedure, the endoscopists satisfaction score (scale 0–10, where 0 is the lowest score and 10 the highest) will be scored.

Sample processing

Handling of the aspirates will be performed according to institutional practice. Nodal aspirates will be assessed by the local pathologist for clinical decision making. Rapid on-site evaluation (ROSE) is optional. When ROSE is performed, smears of the fine-needle aspirate obtained by EBUS-TBNA or EUS-B-FNA of the intrathoracic lymph nodes are performed in situ. All aspirates (both from lymph nodes as well as the tumour) will be processed for both cytology smears and cell block analysis following local practice. The local pathologist will analyse the nodal aspirates, including assessment of PD-L1 expression and molecular analysis for treatment. In case of N0/N1 disease based on the TBNA/TBNA specimens, surgical pathological staging or clinical radiological follow-up at 6 months serves as reference standard for mediastinal/hilar lymph node metastases (N0/N1 vs N2/N3 disease). Which of the two is performed will be decided by the treating physician, as part of clinical practice. In case no malignancy is detected based on TBNA/TBNA specimens, the final diagnosis after 6 months follow-up will serve as the reference standard.

Independent cytopathology review

The local pathologist will assess the sampling material, including cell blocks and cytological smears for clinical purposes. After inclusion of all patients, all cell blocks will be digitalised and reviewed by two reference pathologists for independent cytopathological review. In case less than two cell blocks of a single patient are available, cytological smears will also be included in the independent cytopathological evaluation. The reference pathologists are blinded to needle randomisation but supplied with the clinical information, endosonography route (EBUS or EUS-B), results of the other tests and are aware of the differential diagnosis prior to endoscopy.

The pathologists will evaluate each cell block for the suitability of PD-L1 expression. Cell block specimens will be considered suitable if more than 100 tumour cells are present in the specimen. Additionally, the suitability for molecular analysis will be evaluated based on the presence/absence of >1000 tumour cells in all the cell block material combined. Also, the cumulative length of tissue core will be measured, the sample adequacy will be evaluated based on the presence of lymphocytes or atypical cells or other pathogenic characteristics (eg, granulomas), the quality of the samples will be evaluated using a Mair’s objective scoring system21 and the bloodiness of the samples will be categorised based on the percentage of blood in the microscopic field. In case the
two independent pathologists do not fully agree on one of the above-mentioned parameters, the pathologists will conduct a consensus review of discrepant cases in order to have a clear final judgement.

Outcomes
The primary outcome is the suitability rate for the assessment of PD-L1 expression on mediastinal/hilar nodal or tumour EBUS/EUS-B aspirates in patients with a final diagnosis of lung cancer.

In the initial study design, we planned to examine the hypothesis that the use of the Acquire 22G TBNA was predicted to result in a higher rate of core tissue samples, as was reported in previous studies. However, this endpoint would primarily assess the specimen quality but would not investigate the clinical impact of improved specimen quality. Therefore, it was decided to define a primary endpoint which would define both specimen quality as part of the clinical impact. For that reason, we found that the suitability rate for PD-L1 assessment would be more informative as primary endpoint. After inclusion of five patients, a protocol amendment was submitted and approved by the local medical ethical committee and changes were updated in the trial register.

The secondary outcomes are:
- Cumulative length tissue core.
- Suitability for molecular analysis/next-generation sequencing in patients with a final diagnosis of lung cancer.
- Sample adequacy (defined as the presence of lymphocytes or atypical cells or other pathogenic characteristics (eg, granulomas)).
- Sample quality using the Mair’s objective scoring system.
- Sample bloodiness.
- Diagnostic sensitivity for mediastinal/hilar nodal staging (defined as the proportion of patients that have N2/N3 disease diagnosed by EBUS/EUS-B, relative to the total number of patients with a final diagnosis of N2/N3 disease as determined by the reference standard).
- Diagnostic sensitivity for malignancy (defined as the proportion of patients that have malignancy diagnosed by EBUS/EUS-B, relative to the total number of patients with a final diagnosis of malignancy as determined by the reference standard).
- Yield for diagnosing malignancy in the subgroup of patients with a centrally located lung tumour (defined as the proportion of patients that have malignancy diagnosed by EBUS/EUS-B, relative to the total number of patients with a final diagnosis of malignancy).
- Complication rate (defined as any complication occurring during or directly after the endosonographic procedure or any procedure-related complication in the follow-up period).
- Procedure duration (defined as start EBUS/EUS-B scope insertion until EBUS/EUS-B scope removal).
- Endoscopists satisfaction of needle use.

Sample size calculation
Based on results on the success rate of PD-L1 assessment in the Amsterdam University Medical Centres and published literature, we expect the acquisition rate of the regular 22G TBNA needle for the assessment of PD-L1 expression in patients with lung cancer is around 64%. We expect that this will be 86% for the Acquire 22G TBNA needle. In total, 120 patients with lung cancer and tumour positive samples (eg, lymph node metastasis/primary tumour sample) are needed to show, with α=0.05 and power=0.80, that the Acquire 22G needle is 22% superior to the regular 22G TBNA needle. Taking into account that 64% of included patients will have a final diagnosis of lung cancer with a tumour positive aspirate and a 5% study drop-out, 158 patients will be included. In our opinion, the expected increase in suitability rate for PD-L1 assessment (from 64% to at least 86%) will result in a clinically relevant improvement of lung cancer diagnosis and staging.

Statistical analysis
An intention-to-treat analysis will be applied, which implies that patients will be analysed according to the needle group they will be randomised to also if endoscopy is eventually performed with another needle due to unforeseen medical decisions or circumstances. All such cases will be clearly reported in the final manuscript. Patients who are randomised but in whom no subsequent endoscopic procedure is performed, will be excluded from analysis. All patients in whom the endoscopic procedure is initiated will be included in the analysis for the primary and secondary outcomes. If bronchoscopists fail to obtain (adequate) tissue samples, this will be considered as a failure on the primary outcome (ie, failure to assess PD-L1 expression) and all relevant secondary outcomes. Because of the prospective and mostly cross-sectional nature of the study, we anticipate minimal numbers of missing data. The number of missing data for each variable will be reported in the final manuscript; patients with missing data will be excluded from analysis only for that specific variable, but be included for the analysis of other variables. Results for continuous variables will be expressed as means and SDs, or as medians with IQRs. Categorical variables will be expressed as frequencies and percentages. $\chi^2$ testing will be used to compare dichotomous outcomes, including the primary outcome. Continuous variables will be compared using Student’s t-tests or Mann-Whitney-U tests. A two-tailed p<0.05 will be considered statistically significant.

Patient and public involvement
Patients or the public will not be involved in the design, conduct, reporting or dissemination plans of this research.

Funding
This is an investigator-initiated study with financial and material support from Boston Scientific.
ETHICS AND DISSEMINATION

The study will be conducted according to the principles of the Declaration of Helsinki (64th WMA General Assembly, Fortaleza, Brazil, October 2013) and in accordance with the Medical Research Involving Human Subjects Act. The study was approved by the local Ethics Committee (Medisch Ethische Toetsingscommissie AMC).

Data handling

Data collection will be performed in the participating centres. Electronic patient record forms will be provided through web-based software (Castor EDC software, Amsterdam, the Netherlands). Data collection and analysis will be monitored according to good clinical practice. Clinical monitoring will be organised in a cross-over mode where CRF files will undergo a quality check. Inclusion rate will be monitored on a monthly basis. The investigators will maintain adequate records, including signed patients informed consent forms and information on adverse events. These documents need to be kept in a secured area with limited access. All records will be signed and dated by the investigators. All records are to be retained for a period of 15 years following the date the entire clinical investigation is completed, terminated or discontinued. The anonymity and confidentiality will be guaranteed and patients identification will be coded. The code will start with the abbreviation of the hospital followed by 100 with the number of inclusion (eg, AMC1001). Patients data will be centralised by the coordinating investigator and kept under strict confidentiality according the General Data Protection Regulation. The data of the patients included in the Netherlands will stay in the Dutch database.

Adverse events

EBUS for staging and diagnosing lung cancer is a routine clinical procedure that is considered safe with a low complication rate of less than 1%.25 Adverse events are defined as any undesirable experience occurring to a subject during the study, whether or not considered related to the endosonographic procedure. All adverse events reported spontaneously by the subject or observed by the endoscopist or his staff will be recorded, as well as those occurring up to 1 week after the procedure. The investigators in the participating centres will report all serious adverse events to the coordinating investigator and principal investigator. They will report the serious adverse event to the sponsor without undue delay after obtaining knowledge of the events, as well as to the accredited medical ethics committee that approved the protocol. Only the serious adverse events with a causal association with study participation will be reported in the final study report following study completion.

Annual progress report

The sponsor/investigator will submit a summary of the progress of the trial to the accredited medical ethics committee once a year. Information will be provided on the date of inclusion of the first subject, numbers of subjects included and numbers of subjects that have completed the trial, serious adverse events/serious adverse reactions, other problems and amendments.

Dissemination

We anticipate to report our findings in an article that will be submitted to a peer-reviewed biomedical journal.

DISCUSSION

In this multicentre, randomised clinical trial, we aim to assess for the first time whether the Acquire 22G TBNB needle results in a higher success rate of PD-L1 assessment compared with the standard 22G TBNA needle.19 This is of great importance as the current success rate of PD-L1 and molecular profiling on cytological samples remains limited compared with histology.13-17 As all ancillary tests (both immunohistochemistry as molecular testing) have to be performed on small cytological specimens, sample exhaustion frequently occurs because of insufficient tumour cells to do all tests,24 resulting in unnecessary repeat procedures and treatment delay.

The novel Acquire 22G TBNB has demonstrated a significant improved diagnostic adequacy with increased tissue acquisition in sixty patients with pancreatic masses.24 Therefore, we anticipate that the Acquire 22G TBNB during EBUS/EUS-B procedures in patients with suspected lung cancer will result in a much higher success rate of assessment of PD-L1 expression as compared with traditional needles.

In order to objectively compare the tissue specimens obtained with the Acquire 22G TBNB needle and TBNA needle, two blinded reference pathologists will analyse the tissue specimens on tissue quality and suitability for PD-L1 assessment and molecular profiling. Combined with the international, multicentre design this will improve the generalisability of our findings. Although the reference pathologist will be masked to needle randomisation, it is not possible to mask the attending endoscopist, which is a potential source of performance bias. After a 6-month follow-up period, the diagnostic sensitivity for malignancy and N2/N3 staging for each needle will be calculated. However, the sample size of this study is not powered to assess whether the use of the Acquire 22G TBNB will actually result in an improved diagnostic sensitivity.

The findings of this study will provide novel insights regarding EBUS/EUS-B-guided tissue acquisition in patients with suspected lung cancer using the Acquire 22G TBNB needle, which could be a significant advantage in terms of treatment selection. As the Acquire 22G TBNB needle is commercially available, study results can directly be implemented in clinical practice.

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