Minimally invasive endoscopic staging for mediastinal lymphadenopathy in lung cancer: a systematic review protocol

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To cite: Liu H, Zhou J, Feng Q-ling, et al. Minimally invasive endoscopic biopsy techniques have been widely available as potential alternatives for mediastinal lesions staging in patients with known or suspected lung cancer. Previous efforts have been made to evaluate the diagnostic performance of specific endoscopic modality alone at the level of the mediastinum for staging lung cancer, however, few studies focus on the accuracy of comparisons between different endoscopic modalities, especially at the level of any individual lymph node station. The objective of our study is to determine the diagnostic yields of different endoscopic modalities for staging mediastinal lymphadenopathy in lung cancer, especially concerning the individual lymph node station.

Methods/design: A systematic electronic search of MEDLINE, EMBASE, SinoMed and ISI Web of Science were performed to identify studies evaluating endoscopic modalities accuracy with restriction of English and Chinese languages from inception to an update until May 2014. Data were extracted with the patient as the unit of analysis with regards to the abilities of different endoscopic modalities at the level of mediastinum and particular lymph node station. The methodological quality was assessed independently according to the Quality Assessment of Diagnostic Accuracy Study (QADAS) criteria. An exact binomial rendition of bivariate mixed-effects regression model was used to estimate the pooled sensitivity and specificity. Also, pre–post probability analysis, publication bias analysis and sensitivity analysis were performed for a synthesis of knowledge of this context.

Dissemination: The findings will advance our better available knowledge of optimal clinical decision-making when dealing with staging of mediastinal metastasis in lung cancer.

Trial registration number: PROSPERO—NIHR Prospective Register of Systematic Reviews (CRD42014009792).

INTRODUCTION

Lung cancer has the highest morbidity among all cancers, with an estimated incidence of over 1.6 million cases/year accounting for 13% of all new cancer diagnoses; it is also the leading cause of cancer-related deaths worldwide, with an estimated mortality of over 1.4 million/year, accounting for 18% of all cancer deaths.1,2

Of crucial importance is accurate diagnosis and precise staging of known or suspected lung cancer for the clinician to better determine treatment, guide prognosis and facilitate continued investigation.3 Central to the diagnostic algorithm is the pathological staging in which the evaluation of mediastinal lymph node is a key step for the management of patients with lung cancer, especially in the absence of distant metastases.4,5

Non-invasive imaging scans involving CT, positron emission tomography (PET) and the integrated PET/CT,6,7 are considered favourable for staging mediastinal lymph nodes due to the morphological and functional characteristics of the lesions.8,9 However, more precise information on staging and typing is required for clinical decision-making.10

Mediastinoscopy and thoracoscopy have been recommended as diagnostic standards...
METHODS
Conception and design
The design of this systematic review was elaborated by the multidisciplinary efforts (eg, cardiothoracic surgery, ultrasonography, radiation oncology, diagnostic medicine and health statistics) using methodological approaches outlined in the Cochrane Handbook for Systematic Review of Diagnostic Test Accuracy. This protocol refers to the Preferred Reporting Items for Systematic reviews and Meta-Analyses (PRISMA) criteria. This systematic review has been registered with No. CRD42014009792 in the Centre of Review and Dissemination (CRD) of York University, PROSPERO (the National Institute for Health Research (NIHR) International Prospective Register of Systematic Reviews).

Eligibility criteria for considering studies
Eligible articles will be identified in accordance with the PICOS criteria. Participants (pretreatment patients with suspected or previously diagnosed lung cancer for staging of mediastinal lymph nodes); Interventions (minimally invasive endoscopic techniques: TBNA, EUS-TBNA and EBUS-TBNA); Comparisons (histopathological validation following mediastinoscopy and surgery or close clinical follow-up for at least 6 months); Outcomes (diagnostic sensitivity, specificity, ORs and likelihood ratio) and Study design (any prospective cohort and case–control study).

Search for identification of studies
In addition to Cochrane Central databases (including the Cochrane Database of Systematic Reviews, the Cochrane Central Register of Controlled Trials, the Cochrane Methodology Register, the Database of Abstracts of Reviews of Effects, the Health Technology Assessment Database and the National Health Service (NHS) Economic Evaluation Database), a systematic electronic literature search of MEDLINE, EMBASE, SinoMed and ISI Web of Science will be performed to identify studies that have evaluated the accuracy of endoscopic techniques compared with reference standards in pretreatment patients with suspected or known lung cancer for staging of mediastinal lymph nodes; these studies will be eligible for inclusion. Search will be limited to articles published in English and Chinese, from inception to an update until May 2014. BIOSIS will be used to identify relevant abstracts and conference proceedings, and abstracts and conference proceedings will be included if appropriate. A combination of subject headings and text words will be used for search on the basis of three sets of terms: index tests (minimally invasive techniques aforementioned), target condition (suspected or known lung cancer for staging of mediastinal lymph nodes) and participants description (pretreatment patients with suspected or known lung cancer). Through the process of search, no filter will be used for diagnostic studies to maximise the sensitivity of the search strategies. Further relevant articles will be retrieved for recall completion by searching the bibliographies of identified trials as well as other related systematic and narrative reviews.

Search strategy
Specific search strategy will be developed for each electronic database, commencing with MEDLINE (table 1). The MEDLINE strategy will be adapted for each subsequent database and search yields reported and compared between databases.

Study screening
The primary, subsequent and conclusive screening for inclusion will be on the basis of the article title, abstract and full text, respectively. The screening at each step will adhere to the same inclusion and exclusion criteria. At
the first step, a single author will exclude studies clearly irrelevant to lung cancer or endoscopic staging for mediastinal lymphadenopathy in terms of the abstracts and titles. In case of selection bias, a 10% random sample of all potential references will be validated by a second author for agreement. A further screening will be conducted by at least two independent reviewers for each article reserved after the first step. Definite inclusion decision will be made on the basis of full text of relevant reports by two independent reviewers. Disagreements will be resolved by discussion and consultation with the aid of a third reviewer, if required.

Inclusion criteria
Population and target condition: The study population was patients either suspected or with known lung cancer who received endoscopic staging for mediastinal lymphadenopathy.

Index tests and reference standard: Patients were staged based on at least one minimally invasive endoscopic technique (eg, TBNA, EUS-TBNA and EBUS-TBNA), which should be compared with at least one of the reference standards (eg, tissue histological confirmation of mediastinoscopy and surgery or close clinical follow-up for at least 6 months), irrespective of availability of imaging-based staging before endoscopic techniques.

Exclusion criteria
Studies focusing especially on mediastinal node staging alone; studies consisting of patients with primary underlying disease other than lung cancer; studies unable to populate two-by-two contingency tables of test performance (absolute numbers of true-positive, false-negative, false-positive and true-negative results) from the text, appendices or despite contacting the authors; studies of sample sizes of less than 15 patients; studies of restaging after induction therapy; studies where the positive results from index tests were scarcely confirmed by any reference standard; and studies where duplicates or subcohorts have already been published. At least two independent reviewers will assess the papers for inclusion and exclusion criteria.

Data extraction
The clinical, demographic and methodological quality characteristics of the reference and index tests, as well as the diagnostic results (eg, true positives, true negatives, false positives and false negatives), will be extracted independently by two reviewers. An in-depth discussion of the variability between studies will be provided where applicable. If possible, data for endoscopic biopsies performed at hilar, subcarinal, paratracheal or other lymph node stations will be extracted separately. Additional data were requested from original study investigators, if needed.

Table 1 Searching strategies of the Medline database (results from 14 May 2014)

<table>
<thead>
<tr>
<th>Specified item</th>
<th>Search Query</th>
<th>Result</th>
</tr>
</thead>
<tbody>
<tr>
<td>Target condition set</td>
<td>&quot;Lung Neoplasms&quot;[Mesh]</td>
<td>170 510</td>
</tr>
<tr>
<td></td>
<td>(&quot;lung&quot;[tw] OR pulmonary[tw])</td>
<td>924 000</td>
</tr>
<tr>
<td></td>
<td>&quot;neoplasm staging&quot;[MeSH Terms]</td>
<td>119 312</td>
</tr>
<tr>
<td></td>
<td>Stage[tw] OR stages[tw] OR staging[tw]</td>
<td>806 735</td>
</tr>
<tr>
<td></td>
<td>(&quot;humans&quot;[MeSH] OR human*[tw]) NOT (&quot;animals&quot;[MeSH] OR animal*[tw])</td>
<td>11 775 100</td>
</tr>
<tr>
<td>Index tests set</td>
<td>&quot;Endoscopic Ultrasound-Guided Fine Needle Aspiration&quot;[MeSH]</td>
<td>385</td>
</tr>
<tr>
<td></td>
<td>biops*[tw] OR aspirat*[tw] OR punct*[tw]</td>
<td>498 201</td>
</tr>
<tr>
<td></td>
<td>endobronch*[tw] OR transbronch*[tw] OR (intervention*[tw] AND bronch*[tw])</td>
<td>12 851</td>
</tr>
<tr>
<td></td>
<td>endosonograph*[tw]</td>
<td>9756</td>
</tr>
<tr>
<td></td>
<td>Endoscop*[tw]</td>
<td>169 758</td>
</tr>
<tr>
<td></td>
<td>#1 OR (#2 AND #3)</td>
<td>224 785</td>
</tr>
<tr>
<td></td>
<td>(#3 AND #5) OR #4</td>
<td>119 312</td>
</tr>
<tr>
<td></td>
<td>#14 AND #15</td>
<td>16 683</td>
</tr>
<tr>
<td></td>
<td>#17 AND #16</td>
<td>16 406</td>
</tr>
<tr>
<td></td>
<td>#18 AND #10 AND #11</td>
<td>405</td>
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<tr>
<td></td>
<td>#19 AND #10 AND #12 AND #13</td>
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<tr>
<td></td>
<td>#20 OR #8 OR #18 OR #19</td>
<td>5133</td>
</tr>
<tr>
<td></td>
<td>#21 AND #20</td>
<td>364</td>
</tr>
</tbody>
</table>

addition, the publication year, the study period, setting (eg, community vs academic hospitals; and primary vs tertiary centres), design (eg, cohort or case–control; and consecutive or random series), centre (eg, single or multiple), potential differences in reference standards, follow-up period and prevalence of mediastinal metastases will be described as covariates that may be the cause of heterogeneity. Two reviewers will independently extract data. Any disagreements will be resolved by consensus.

**Assessment of methodological quality**
In terms of the risk of bias and applicability concerns, the methodological quality of the articles included will be independently assessed by two reviewers using a subset of criteria derived from the Quality Assessment of Diagnostic Accuracy Study 2 (QUADAS-2) tool. A third reviewer will be consulted in case of discrepancies. Blinding implies that the results of minimally invasive endoscopic staging results must be interpreted without the knowledge of those of reference standards. Reviewers’ judgments about risks of bias and applicability concerns will be used in sensitivity analysis to determine the potential effect of methodological quality on diagnostic performance. Two reviewers will independently assess study quality. Any disagreements will be resolved by consensus.

**Data synthesis and analysis**
For each study we will collect the rates of true positives, false positives, true negatives and false negatives and produce from the crude data the sensitivity and specificity. In addition to the diagnostic accuracy of different endoscopic modalities for staging total mediastinal lymphadenopathy, we will shed some light on the accuracy for individual lymph node station and the lesions in different compartments of the mediastinum. A bivariate mixed-effects regression model will be used to generate the pooled estimates of sensitivity and specificity when appropriate. Meanwhile, summary receiver operating characteristic (SROC) curves with the corresponding area under the curve (AUC) will also be graphically generated to determine the diagnostic accuracy, in which a summary operating point (pooled sensitivity and specificity) with the corresponding 95% CI and 95% prediction region, and the point Q* (maximum joint sensitivity and specificity) will also be calculated in addition to the point estimates for each study as well as a symmetrical summary curve. Comparison of diagnostic accuracy between the groups will be achieved by Z test on the basis of AUC of SROC.

**Threshold analysis**
Spearman’s rank correlation will be performed for exploring the effect of threshold effect. To ensure the effect of variability in diagnostic threshold on the shape of SROC curve, threshold effect will be detected by the regression equation D=a+bS, where D is the log of the diagnostic OR and S is a measure of the diagnostic threshold. The variables a and b will then be estimated using a least-squares method weighted by inverse variance.

**Heterogeneity investigation**
The expected heterogeneity across studies will be detected by the meta-regression analysis and subgroup analysis. The pattern of heterogeneity will be detected using $\chi^2$ test and the magnitude using $I^2$ statistic. A study with an $I^2$ greater than 50% will be considered substantially heterogenous.

**Meta-regression analysis**
Meta-regression will be performed to evaluate the importance of potential effect variables and explain variation between studies. Univariate analysis will allow the investigation of the potential source of statistical heterogeneity. Subsequently, multivariate analysis will allow the determination of the important confounders affecting the diagnostic performance by the method of backward elimination. The covariates are consisted with the demographic, clinical and endoscopic characteristics.

**Subgroup analysis**
To explore clinical heterogeneity, we will fit a separate SROC curve for these planned subgroups of patients: (1) different anatomic compartments of the mediastinum (superior vs middle vs inferior mediastinum); (2) different types of endoscopic techniques (TBN vs EUS-TBNA vs EBUS-TBNA vs incorporation of the modalities); (3) different populations (known vs suspected mediastinal lymph node involvement; with vs without the distant metastasis) and (4) sequential imaging (with vs without imaging scans before the implement of endoscopic techniques).

**Probability analysis**
The pretest probabilities of different prevalences of mediastinal metastases among patients suffering from lung cancer and corresponding post-test probabilities will be evaluated, depending on the summary sensitivity and specificity by Fagan’s analysis, which will allow the determination of the relationship between the pretest and post-test probability as well as the likelihood ratio. The difference in the proportions of mediastinal lymphadenopathy in lung cancer (post-test probability minus pretest probability) will gave the overall added value of endoscopic staging.

**Publication bias**
Publication bias will be examined visually by inspecting funnel plots and statistically by using Egger’s regression model. If publication bias was present, the effect of such bias on the summary estimate would be assessed using the trim and fill method. This method will impute the missing studies and recalculate a new summary estimate. The difference between the calculated and
observed value will be then used to determine the effect of bias on the diagnostic performance of the index tests.

**Post-hoc power analysis**

The technique of power analysis allows for the determination of how likely a statistical test of an individual study will be to detect effects of a given sample size in a particular situation. The determination of post-hoc power for testing differences in proportions for the matched pair design with binary response outcome will be performed for each study included, according to the formula derived by Lachin.\(^2^\)

**Sensitivity analysis**

For detecting the robustness of pooled results, sensitivity analysis will be performed to determine the potential impacts of study design on endoscopic staging accuracy.\(^3^\) On the basis of QUADAS-2, the important following sensitivity analyses will be prespecified: risk of selection bias, risk of interpretation bias for the index tests as well as reference standards.\(^4^\)

All the analyses will be conducted using MADIS module of Stata V.10 (Stata Corp, College Station, Texas, USA), except for publication bias using Meta-Analyt \(\beta\) 3.13 (Tufts Medical Center, Boston, Massachusetts, USA), and meta-regression and threshold analysis using Meta-DisC V.1.4 (Clinical Biostatistics Unit, Ramón y Cajal Hospital, Madrid, Spain).

**DISCUSSION**

The problem-oriented research will enable the best available knowledge of clinical practice, on the strength of comprehensively rigorous methodology used in the review. The internal validation and navigation involved in the process will minimise the potential of selection bias and systematic errors. On the basis of a synthesis of methodological quality estimation for each study included, the findings will be available for a further subgroup analysis to explore potential methodological heterogeneity. Considering the high dependence of the findings on the quality of the underlying primary studies, as well as the potential risk of bias from limited quality studies on clinical practice, only the prospective studies will be included in our current review in order to provide more convincing evidence.

The findings of this systematic review will provide important evidence for endoscopic staging for mediastinal lymphadenopathy in lung cancer, as well as enable optimal clinical decision-making when involved in staging for mediastinal lymph node involvement. The findings will also help to better advance our available knowledge of the management strategy in individualised, targeted and comprehensive treatment of, as well as aiding in the determination of prognosis among, patients suffering from lung cancer. Furthermore, the findings may trigger an update or drive the development of related standards, which will lead to refine mediastinal staging in lung cancer.

To our knowledge, the current review will allow the diagnostic accuracy of different endoscopic staging for mediastinal lymphadenopathy in lung cancer, for the first time, from an overall level to the individual node station level, on the basis of evidence-based research. Given that we focus on results from practice and emphasise outcomes evaluation, the results from this review will put forth the advantages and disadvantages in medical practice of using minimally invasive endoscopic biopsy in staging mediastinal lymphadenopathy in lung cancer, and will lay the foundation for further research and the development of establishing optimal staging pathways for patient subgroups.

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**Contributors**

HL conceived and designed the review and completed the PROSPERO registration; HL and JZ conducted the scoping searches and drafted and revised the manuscript. GW and Q-lF were involved in the design of the review and piloted the inclusion and exclusion criteria and the extraction forms. HL, JZ, Q-lF, GW, H-HG and Y-jX provided content expertise and feedback on the design of the review, the protocol and on the manuscript. All authors shared the interpretation of data and critical revision of the manuscript for important intellectual content. All authors critically reviewed the first draft and contributed to the production of the final manuscript and its subsequent revision.

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**Competing interests**

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**Patient consent**

Obtained.

**Provenance and peer review**

Not commissioned; peer reviewed for ethical and funding approval prior to submission.

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**REFERENCES**


