Oesophageal endoscopic ultrasound with fine needle aspiration improves and simplifies the staging of lung cancer


Background: Positron emission tomography (PET) is accurate for mediastinal staging of lung cancer but has a moderate positive predictive value, necessitating pathological verification. Endoscopic ultrasonography with fine needle aspiration (EUS-FNA) is a technique for tissue verification of mediastinal and upper retroperitoneal abnormalities. The use of EUS-FNA may decrease the number of surgical procedures and thereby staging costs.

Methods: EUS-FNA was used prospectively for the cytological assessment of mediastinal and/or upper retroperitoneal PET hot spots in patients with suspected lung cancer. Only if EUS-FNA was positive for malignancy was subsequent mediastinoscopy or exploratory thoracotomy cancelled. The cost effectiveness of EUS-FNA was determined.

Results: Of 488 consecutive patients with suspected lung cancer, 81 were enrolled with mediastinal and/or upper retroperitoneal PET hot spots. EUS-FNA was positive in 50 (62%) patients, negative in six, and inconclusive in 25. Of the 31 negative or inconclusive patients, 26 underwent surgical staging (resulting in 14 patients with and 12 without mediastinal malignancy), while five patients had mediastinal metastases during follow up. No EUS-FNA related morbidity or mortality was encountered. The accuracy of the decision to proceed to surgery (or not) on the basis of EUS-FNA was 77% (95% CI 68 to 86). EUS-FNA detected more mediastinal abnormalities than PET except for the upper retroperitoneal region. Addition of EUS-FNA to conventional lung cancer staging reduced staging costs by 40% per patient, mainly due to a decrease in surgical staging procedures.

Conclusion: EUS-FNA can replace more than half of the surgical staging procedures in lung cancer patients with mediastinal and/or upper retroperitoneal PET hot spots, thereby saving 40% of staging costs.


The prognosis of patients with lung cancer largely depends on the presence of mediastinal and distant metastases. Malignant involvement of mediastinal lymph nodes is a highly significant prognostic factor for survival. Complete surgical resection of lung cancer is often impossible in patients with mediastinal metastases. Better preoperative staging may limit the number of unnecessary surgical interventions in these patients. Positron emission tomography (PET) with 18F-fluorodeoxyglucose (FDG) has already shown interesting properties in this area. Oesophageal endoscopic ultrasonography with fine needle aspiration (EUS-FNA) is a technique for the assessment of lymph nodes and tumours in previously almost inaccessible areas such as the lower mediastinum and upper retroperitoneum. Incorporation of this tumour verification technique in lung cancer patients may improve staging of the disease.

FDG-PET is more accurate than computed tomography (CT) in detecting mediastinal lymph node metastases, with a negative predictive value of 93–95%. However, a positive predictive value of 74–90% makes pathological verification of hot spots necessary to avoid patients being denied a possible cure by surgery. Various staging modalities are used for pathological examination of mediastinal hot spots such as mediastinoscopy, mediastinotomy, and exploratory thoracotomy. However, these techniques are invasive and require general anaesthesia and often hospitalisation, and it is difficult to assess the entire mediastinum—especially the posterior and lower parts. Nevertheless, mediastinoscopy is traditionally considered the “gold standard” for establishing mediastinal metastases.

EUS-FNA combines an excellent view into the mediastinum (EUS) with safe and accurate fine needle aspiration (FNA). It is performed in a short outpatient setting without general anaesthesia, and assessment of at least seven of the 13 right and left mediastinal lymph node levels is possible. The upper retroperitoneum can also be evaluated, especially the area around the celiac trunk and the left adrenal gland. The exact role of EUS-FNA in lung cancer staging is not yet defined, and it is not clear whether it can be used with CT and PET to discriminate accurately between resectable and irresectable lung cancer. Establishing mediastinal or upper retroperitoneal metastases with EUS-FNA makes surgical staging redundant and may result in cost savings for the staging process.

A prospective study was undertaken to determine whether EUS-FNA can reduce the number of surgical staging procedures in lung cancer patients with mediastinal and/or upper retroperitoneal PET hot spots, and whether this approach is cost effective.

Abbreviations: CT, computed tomography; EUS, endoscopic ultrasonography; FNA, fine needle aspiration; PET, positron emission tomography
METHODS

Patients
Consecutive patients with suspected or pathologically established lung cancer were eligible if they had mediastinal and/or upper retroperitoneal PET hot spots without evidence of further distant metastases, if they had a resectable primary tumour as judged by two thoracic surgeons, and if they were medically operable. All patients gave informed consent and the study was approved by the medical ethics committee of the Groningen University Hospital.

Patients were evaluated by history, physical examination, complete blood cell count, renal and liver function tests, chest radiography, bronchoscopy, CT scan of the chest and upper abdomen, and FDG-PET. All test results were presented in a multidisciplinary session where disease stage, tumour resectability, and medical operability were verified before EUS-FNA was performed.

Imaging tests
CT was performed with contrast enhanced single and multislice scanners (Somatom Sensation 16/Somatom AR, Siemens AG, Forchheim, Germany). It was used to assess the resectability of the primary tumour, to evaluate mediastinal lymph nodes, and to exclude distant metastases. Whole body PET was performed with an ECAT 951/31 or ECAT HR+ scanner (Siemens/CTI, Knoxville, TN, USA) 90 minutes after intravenous injection of a median dose of 390 MBq (range 130–670) FDG. Attenuation correction was obtained on the HR+ camera by transmission scanning with a germanium-68/gallium-68 rod source using an interleaved protocol. PET was used for the detection of mediastinal and distant metastases. If mediastinal and/or upper retroperitoneal hot spots were detected, PET was considered positive. Patients with hot spots suggesting metastases outside these areas were excluded from the study. A hot spot was defined as a focal increase in FDG uptake compared with the background not explained by physiological uptake. Both CT and PET were used as guidance for EUS-FNA.

EUS-FNA
EUS was performed with a linear scanning ultrasound endoscope (FG-34UX, Pentax GmbH, Hamburg, Germany) connected to a scanner unit (EUB-525, Hitachi Ultrasound BV, Reeuwijk, the Netherlands) with Doppler flow imaging for the detection of blood vessels. Midazolam 3–5 mg was administered intravenously for conscious sedation with continuous monitoring of pulse rate and peripheral oxygen saturation. The endoscope was introduced into the stomach to scan the left adrenal gland and the area around the celiac trunk. After retraction into the oesophagus, the mediastinum was evaluated by scanning 360° transaxially at 1–2 cm intervals upward up to Naruke level 2. The left adrenal gland was considered enlarged if it had lost its normal elliptical, triangular, or “seagull” shape. Lymph nodes were identified as non-pulsating regions with round, ellipsoid, crescent, or triangular shape. They were separately punctured with a 22 gauge needle (Echotip, Wilson-Cook Medical Inc, Winston-Salem, NC, USA) using a 10 ml syringe for suction. A cytotechnician evaluated the cellular contents of the specimens on site with a modified Giemsa method. If a specimen was inadequate, additional punctures were performed. Specimens were categorised as positive (tumour cells), negative (lymphoid but no tumour cells), or inconclusive (poor cellularity or unable to perform adequate puncture). Subsequent mediastinoscopy and/or exploratory thoracotomy with mediastinal dissection was performed in all patients except those with a positive EUS-FNA. The mediastinal lymph node map of the American Joint Committee on Cancer was used to localise abnormalities at PET, EUS-FNA, and mediastinal dissection.

Because EUS-FNA is a new procedure for lung cancer staging, we were interested in the learning curve of two residents. During each session their ability to identify mediastinal lymph nodes was recorded. One of the residents performed EUS-FNA and an experienced pulmonary physician repeated the procedure. The numbers of observed mediastinal lymph nodes of the residents and the pulmonary physician were compared.

Economic analysis
Economic analysis was performed to assess the cost effectiveness of adding EUS-FNA to the staging algorithm. It was performed from a hospital point of view. All direct medical costs were included until the final lung cancer stage was established. This also included functional tests or additional examinations necessary for the evaluation of the patient’s operability. All cost items of EUS-FNA, mediastinoscopy, and exploratory thoracotomy were estimated by expert opinions of pulmonary physicians, thoracic surgeons, anaesthesiologists, and by observations of two hospital economists. Calculations were based on real costs using the 2002 price level. Costs of hospital admissions were calculated from the actual number of admission days to the general ward and intensive care unit against the Dutch unit price for hospital admission days. Overhead costs were calculated as a 35% surcharge. Because of the short time horizon of the diagnostic process, discounting for future costs was not applied.

Statistical analysis
The study was designed to determine prospectively the proportion of lung cancer patients with mediastinal and/or upper retroperitoneal PET hot spots in whom EUS-FNA can diagnose malignancy. Since EUS-FNA can reach at least seven of the 14 (right and left) mediastinal lymph node levels and the upper retroperitoneum, we estimated that about 50% of the surgical staging procedures could be avoided. Mediastinal PET has a positive predictive value of 67%. We therefore expected EUS-FNA to be able to identify 50% of this 67% as pathologically malignant—that is, 34% of all PET

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>No (%)</th>
</tr>
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<tbody>
<tr>
<td>Male sex</td>
<td>54 (67)</td>
</tr>
<tr>
<td>ECOG performance score</td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>12 (15)</td>
</tr>
<tr>
<td>1</td>
<td>69 (85)</td>
</tr>
<tr>
<td>Pathological diagnosis</td>
<td></td>
</tr>
<tr>
<td>Squamous cell carcinoma</td>
<td>20 (25)</td>
</tr>
<tr>
<td>Adenocarcinoma</td>
<td>17 (21)</td>
</tr>
<tr>
<td>Large cell carcinoma</td>
<td>28 (35)</td>
</tr>
<tr>
<td>Small cell carcinoma</td>
<td>6 (7)</td>
</tr>
<tr>
<td>Malignant lymphoma</td>
<td>3 (4)</td>
</tr>
<tr>
<td>Sarcoidosis</td>
<td>1 (1)</td>
</tr>
<tr>
<td>Aspergillosis</td>
<td>1 (1)</td>
</tr>
<tr>
<td>Unknown</td>
<td>1 (1)</td>
</tr>
<tr>
<td>Other</td>
<td>4 (5)</td>
</tr>
<tr>
<td>CT characteristics</td>
<td></td>
</tr>
<tr>
<td>Mediastinum and retroperitoneum</td>
<td>13 (16)</td>
</tr>
<tr>
<td>normal</td>
<td></td>
</tr>
<tr>
<td>Enlarged mediastinal lymph nodes</td>
<td>56 (69)</td>
</tr>
<tr>
<td>Mediastinal and retroperitoneal</td>
<td>8 (10)</td>
</tr>
<tr>
<td>abnormalities</td>
<td></td>
</tr>
<tr>
<td>No CT performed</td>
<td>4 (5)</td>
</tr>
</tbody>
</table>
positive patients. Eighty patients were needed to estimate this proportion with a precision of 10%.

Statistical analysis was performed with SPSS 10.0 (SPSS Inc, Chicago, IL, USA). Continuous variables are reported as median (range) or mean (SE). Dichotomous variables are reported as percentages with 95% confidence interval (95% CI). The accuracy of EUS-FNA was calculated as the accuracy of the decision to proceed to surgical staging (or not) based on EUS-FNA results. For comparisons between groups of categorical variables, Pearson’s $\chi^2$ test was used. Interobserver agreement for test results was calculated with Cohen’s $k$ value. Multiple logistic regression analysis was used to identify ultrasound features predictive for lymph node malignancy. We performed sensitivity analysis of the cost of the staging procedures by varying the major cost elements.14 15 Reported p values are two sided and p values, <0.05 were considered significant.

RESULTS

Patients

Between January 2001 and March 2003, 488 consecutive patients were screened at the outpatient department, 81 of whom fulfilled all the inclusion criteria and were enrolled in the study (table 1). Their median age was 63 years (range 32–80).

Imaging tests

Images from a typical diagnostic procedure are shown in fig 1. Sixty four patients (79%) patients had mediastinal and/or upper retroperitoneal abnormalities on the CT scan. Interobserver agreement (k value) for the detection of enlarged lymph nodes and left adrenal enlargement on CT scanning was 0.76 (95% CI 0.66 to 0.86). All patients had mediastinal and/or upper retroperitoneal PET hot spots, mostly at Naruke level 4-right, 4-left, and 7 (table 2).

EUS-FNA

EUS-FNA was positive for malignancy in 50 patients, thereby avoiding subsequent surgical staging procedures in 62% of the patients (95% CI 51 to 73). EUS-FNA was negative for malignancy in six (7%) and inconclusive for malignancy in 25 (31%) patients (fig 2). The accuracy of EUS-FNA was 0.77 (95% CI 0.68 to 0.86). Most patients had EUS abnormalities
at Naruke levels 4-left, 5, and 7 (table 2). After PET, EUS detected additional abnormalities at Naruke levels 4-left, 5, 7, 8-right, and 8-left. EUS was not able to visualise accurately Naruke levels 1, 2-right, 2-left, 3, or 4-right.

Twenty six of the 31 patients with a negative or inconclusive EUS-FNA underwent a surgical staging procedure: mediastinoscopy in 12 patients, mediastinotomy in one, exploratory thoracotomy with mediastinal dissection in nine, and mediastinoscopy followed by exploratory thoracotomy with mediastinal dissection in four patients. Of the six patients with a negative EUS-FNA, two had positive and three had negative mediastinal lymph nodes at surgery (fig 2). Further staging was not performed in one patient due to rapid progression of the mediastinal lesions. Of the 25 patients with an inconclusive EUS-FNA, 12 were positive and nine were negative for malignancy at surgery (fig 2). Four patients refused further surgical staging and, after follow up of 6 months, all had progressive disease diagnosed with imaging tests.

After all staging procedures including mediastinal dissection, mediastinal or upper retroperitoneal malignancy was not detected in 12 of the 81 patients (15%). A negative or inconclusive EUS-FNA did not reliably prove the absence of mediastinal metastasis: 19 of the 31 patients (68%) had lymph node metastasis at surgery. Of these 19 patients, seven had lymph node metastasis exclusively at the upper right mediastinum, outside the reach of EUS-FNA.

One mediastinal lymph node (which was localised at the PET hot spot) was randomly selected per patient for multiple logistic regression analysis. Lymph node size, shape (flat, round, irregular), boundary (vague, sharp), core structure (hyperechogenic, hypoechogenic), and visual impression (benign, malignant) were used as factors in the analysis. Lymph node size was the only significant predictor of malignancy ($p = 0.03$). For each additional centimetre the chance of malignancy increased by 0.56 (95% CI 0.04 to 1.00).

The learning process of EUS-FNA of two residents was studied during 54 procedures, 29 by the first resident and 25 by the second. The numbers of lymph nodes detected by the residents failed to reach those identified by the experienced operators. More procedures are therefore required to learn EUS-FNA accurately by those who have not previously performed oesophageal endoscopy.

The median duration of the EUS-FNA procedure (including resident training) was 45 minutes (range 20–60). There was no morbidity or mortality during or after EUS-FNA.

**Economic analysis**

Costs are summarised in table 3. Hospitalisation was a major cost item for mediastinoscopy and exploratory thoracotomy with a mean (SE) duration of 3.0 (0.3) days for mediastinoscopy and 12.0 (1.9) days for exploratory thoracotomy. Average staging costs were $2101 per patient using a staging algorithm with EUS-FNA. For a staging algorithm without

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**Table 3 Real costs per procedure (US$)**

<table>
<thead>
<tr>
<th>Cost item</th>
<th>EUS-FNA</th>
<th>Mediastinoscopy</th>
<th>Exploratory thoracotomy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Personnel</td>
<td>92</td>
<td>308</td>
<td>517</td>
</tr>
<tr>
<td>Equipment and materials</td>
<td>498</td>
<td>143</td>
<td>400</td>
</tr>
<tr>
<td>Medication</td>
<td>2</td>
<td>28</td>
<td>91</td>
</tr>
<tr>
<td>Overheads*</td>
<td>208</td>
<td>168</td>
<td>352</td>
</tr>
<tr>
<td>Other costs</td>
<td>0</td>
<td>43</td>
<td>65</td>
</tr>
<tr>
<td>Mean hospitalisation (days)</td>
<td>(0)</td>
<td>(3)</td>
<td>(12)</td>
</tr>
<tr>
<td>Hospitalisation ($356/day)</td>
<td>0</td>
<td>1068</td>
<td>4272</td>
</tr>
<tr>
<td>ICU stay (days†)</td>
<td>0</td>
<td>0</td>
<td>125</td>
</tr>
<tr>
<td>ICU stay ($1252/day)†</td>
<td>0</td>
<td>0</td>
<td>5822</td>
</tr>
<tr>
<td>Total costs</td>
<td>798</td>
<td>1758</td>
<td>5822</td>
</tr>
</tbody>
</table>

*Calculated as 35% of the costs of personnel, equipment, materials, and medication.  †One overnight stay in the intensive care unit (ICU) was assumed after 10% of the exploratory thoracotomies.
EUS-FNA, average staging costs were $3514 per patient, assuming that the ratio of mediastinoscopies to exploratory thoracotomies remained constant (17:13). Thus, mean cost savings through EUS-FNA were $1413 (40%) per patient. Sensitivity analyses were performed by varying the costs of the staging procedures. Cost savings could be maintained even if the costs of EUS-FNA were 2.5 times the current costs. Likewise, even with a 75% reduction in the costs of mediastinoscopy or a 50% reduction in the costs of exploratory thoracotomy, the outcome still favoured the EUS-FNA algorithm. Increasing the ratio of mediastinoscopies from 17:13 to 10:1 in the staging algorithm without EUS-FNA (that is, a reduction in exploratory thoracotomies of 80%) would result in equal costs of both staging algorithms.

**DISCUSSION**

Mediastinoscopy is still considered the gold standard for mediastinal staging of lung cancer. However, this procedure has no access to the aortopulmonary window and lower mediastinum, and has a complication risk of 2.5%. EUS-FNA, on the other hand, is a minimally invasive short outpatient procedure with hardly any complications. It can reach aortopulmonary, subcarinal, and paraesophageal lymph nodes, as well as the upper retroperitoneum. Due to air in the trachea, EUS cannot image the right upper mediastinum adequately. EUS-FNA is useful and appears more flexible than the transbronchial biopsy with a Wang needle. The nodes can be viewed directly and adjacent structures demonstrated. Mediastinoscopy and EUS-FNA may be considered as complementary techniques since their anatomical views are only partly overlapping. As shown in this study, EUS-FNA can limit the number of mediastinoscopies and exploratory thoracotomies in patients with irresectable tumours by 62%. It is therefore the diagnostic test of first choice for further staging of lung cancer patients with mediastinal and/or upper retroperitoneal PET hot spots.

Our report is the first study in which EUS-FNA has been used for cytological verification of PET hot spots in the mediastinum and upper retroperitoneum. Direct comparison of EUS-FNA with CT and PET is interesting, but it overlooks the actual clinical application of EUS-FNA. With a serial diagnostic approach, performing first CT and PET followed by EUS-FNA for cytological diagnosis, many surgical staging procedures will be prevented. The performance of EUS-FNA in patients with PET negative mediastinum was not part of the study. However, some adenocarcinomas or other low grade tumours which may be PET negative and small lesions not resolving on PET may benefit from EUS-FNA. A randomised study will be needed to determine whether EUS-FNA assesses mediastinal metastases better than PET.

PET is not available in all hospitals. However, EUS-FNA can be guided by CT or even used as the first procedure after (or in conjunction with) bronchoscopy in lung cancer staging. EUS-FNA proved to be of great value in this respect. The decrease in surgical staging procedures in our study was higher than we initially expected. This can partly be explained by a high number of tertiary referrals, resulting in a higher positive predictive value of PET hot spots than expected (85% vs 67%). This figure is similar to the positive predictive values reported in a recent meta-analysis.

Procedures that may be an alternative to EUS-FNA include percutaneous transthoracic needle biopsy (PTNB), transbronchial fine needle aspiration (TBNA), and endobronchial ultrasonography (EBUS). As outlined in a recent review by our study group, PTNB cannot reach as many lymph node stations as EUS-FNA and it has a higher complication rate. The performance of TBNA in mediastinal lymph node staging is variable, and it is only possible for lymph nodes neighbouring large airways. The use of EBUS is promising but there is still little experience with this technique. Diseases other than lung cancer can also be identified with EUS-FNA. Diagnosis and staging are usually performed in concert, since the most efficient way to make a diagnosis is often dictated by the stage of the cancer. We encountered 13% of patients with other diagnoses such as malignant lymphoma, sarcoidosis, and aspergillosis. They mostly presented with atypical signs, symptoms, and CT abnormalities, and without a pathological diagnosis.

The resolution of EUS-FNA within a view field of several centimetres is higher than that of PET, and at least that of CT. Abnormalities of 2–3 mm can easily be located and punctured. EUS-FNA can also localise mediastinal tumour invasion. Moreover, it can be used to search for metastases in areas surrounding mediastinal invasion by tumour, thereby selecting patients who are candidates for more extensive surgery. The number of inconclusive cytological results will increase, since more normal lymph nodes are punctured.

In this study the ability of EUS to identify lymph nodes at the site of PET hot spots was total, but FNA punctures were inconclusive in almost one third of the patients. This can partly be explained by the bloody content of the EUS-FNA specimens and sometimes by attempts to puncture very small lymph nodes. Moreover, in most cases after three punctures at the same site no further punctures were performed. Replacing the on-site cytotechnician by an on-site cytopathologist may perhaps increase the number of adequate specimens obtained.

We analysed the number of procedures required to learn accurate EUS-FNA and found that less than 30 procedures are not sufficient to acquire the necessary skills. We estimate that about 40 procedures will be enough to perform EUS-FNA adequately, including interpretation of ultrasound images of the left adrenal glands and mediastinal lymph node stations.

The cost effectiveness of EUS-FNA in this study was not based on a decision analysis model but on real cost assessments. The addition of EUS-FNA to the staging algorithm avoided surgical staging procedures in 62% of patients, thereby decreasing staging costs by 40%. Our sensitivity analyses show that the cost effectiveness of EUS-FNA is maintained over a wide variation in procedure costs. In conclusion, EUS-FNA is a safe and cost effective procedure for lung cancer staging which detects additional mediastinal abnormalities after PET.

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**Authors’ affiliations**

H Kramer, J W G van Putten, H J M Groen, Department of Pulmonary Diseases, Groningen University Hospital, Groningen, The Netherlands

H M van Dullmen, Department of Gastroenterology, Groningen University Hospital, Groningen, The Netherlands

A H H Bongaerts, Department of Radiology, Groningen University Hospital, Groningen, The Netherlands

A J H Suurmeyer, Department of Pathology and Laboratory Medicine, Groningen University Hospital, Groningen, The Netherlands

T J Klinkenberg, Department of Thoracic Surgery, Groningen University Hospital, Groningen, The Netherlands

W J Post, H Groen, Office for Medical Technology Assessment, Groningen University Hospital, Groningen, The Netherlands

J Pruijn, Positron Emission Tomography Centre, Groningen University Hospital, Groningen, The Netherlands

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REFERENCES


LUNG ALERT

A novel method for the diagnosis of lung transplant rejection


The most common causes of morbidity and mortality following lung transplantation are primary graft failure, infection, and acute and chronic rejection. Clinical criteria alone are inaccurate for the diagnosis of graft rejection. The diagnosis can be made by bronchoscopy or surgical biopsy but this procedure is invasive and cannot be performed repeatedly. Hepatocyte growth factor (HGF) is a glycoprotein highly expressed in the lung parenchyma and produced after acute lung injury. This study was designed to assess whether serum HGF could act as a marker of acute rejection following lung transplantation.

The authors measured serum concentrations of HGF by ELISA in 109 patients undergoing lung transplantation (65 with chronic obstructive pulmonary disease, 23 with cystic fibrosis, and 21 with idiopathic pulmonary fibrosis) and 12 healthy controls. The mean serum HGF concentration before transplantation was higher in the patients than in the controls, and postoperatively rose significantly on the first day before decreasing by day 3. At an episode of acute rejection the serum HGF concentration was significantly higher than in all other groups (controls, patients before transplantation, patients with no rejection, and patients with infection). Logistic regression analysis identified the serum HGF concentration as an independent predictor of lung graft rejection (p = 0.012). After steroid treatment HGF concentrations returned to the preoperative value within 3 days.

The authors conclude that, since serum concentrations of HGF increase with graft rejection but not infection and the values decline with steroid treatment, serum HGF monitoring could be a sensitive predictor of lung graft rejection.
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