Accuracy of staging in early oesophageal cancer using high resolution endoscopy and high resolution endosonography: a comparative, prospective, and blinded trial

A May, E Günter, F Roth, L Gossner, M Stolte, M Vieth, C Ell

Background and aims: The increasing use of endoscopic resection for curative treatment of early oesophageal cancers requires accurate staging before therapy. In a prospective blinded trial, we compared staging of early oesophageal carcinoma using high resolution endoscopy (HR-E) with staging using high resolution endosonography (HR-EUS).

Patients and methods: A total of 100 patients (89 men, 11 women; mean age 63.9 (10.8) years (range 31–91)) with a suspicion of early oesophageal adenocarcinoma (n = 81) or squamous cell carcinoma (n = 19) were enrolled in the study. After endoscopic staging with high resolution video endoscopy by two experienced endoscopists, HR-EUS was performed by an experienced endosonographer who was blinded to the endoscopic assessment. Results of the staging examinations were correlated with the histology of the resected tumours.

Results: Overall rates for accuracy of the endoscopic and endosonographic staging were 83.4% and 79.6%, respectively. Sensitivity for mucosal tumours (n = 68) was more than 90% [EUS 91.2%, endoscopy 94.1%] while sensitivity for submucosal tumours [n = 25] was lower, at 48% for EUS and 56% for endoscopic staging. A combination of the two techniques increased the sensitivity for submucosal tumours to 60%. Submucosal tumours in the tubular oesophagus were significantly better staged with HR-EUS than submucosal tumours close to the oesophagogastric junction (10/11 vs 2/14; p<0.001). Tumours infiltrating the second and third submucosal layers were also more correctly diagnosed than tumours with slight infiltration of the first submucosal layer (sm1).

Conclusions: The overall diagnostic accuracy of both HR-E and HR-EUS with a 20 MHz miniprobe in early oesophageal cancer was high (approximately 80%), with no significant differences between the two techniques. HR-E and HR-EUS provide a high level of diagnostic accuracy for mucosal tumours and submucosal tumours located in the tubular part of the oesophagus. With submucosal tumours located at the oesophagogastric junction or with infiltration of the first third of the submucosa however, the diagnostic accuracy of both techniques is not yet satisfactory.
squamous cell carcinoma (n = 19) were enrolled into this prospective blinded comparison. Staging procedures (video endoscopy with chromoendoscopy, ultrasound of the abdomen, and computed tomography of the chest and upper abdomen) were selected in the same way as in our prospective studies on endoscopic therapy in early Barrett’s adenocarcinoma and intraepithelial high grade neoplasia.9,11 After the staging procedures had been completed, a decision was taken as to whether the patient was suitable for local endoscopic resection with curative intent. Endoscopic resection was carried out using the suck and cut technique in the majority of cases. The resection techniques have been described in detail previously and can be referred to in the literature.9–11

Exclusion criteria for endoscopic resection were any suspicion of a tumour stage more than 1 or suspicious mediastinal lymph nodes (with enlargement >10 mm and/or a hypoechoic pattern). All patients with exclusion criteria were scheduled for radical oesophagectomy if they were generally operable, or for palliative endoscopic treatment if they were inoperable. If there was a suspicion of submucosal tumour infiltration, the patient received extensive information concerning surgical oesophageal resection and endoscopic therapy, including endoscopic resection, photodynamic therapy, or thermal destruction. The decision regarding further therapeutic management was made in consultation with the patient, weighing up personal risks (accompanying diseases, age, etc). All patients provided written consent to endoscopic resection. Further exclusion criteria for the trial comparing endoscopy and were: presence of low grade or high grade intraepithelial neoplasias, as well as clearly advanced tumours; endoscopic evidence of a very large and/or multifocal early cancer, for which photodynamic therapy was planned from the outset; and if blinding of the examiners was not maintained.

Analysis of the data was carried out by two team members who were not involved in collecting the endoscopic or endosonographic data (AM, FR).

### Endoscopic staging

Initially, endoscopic staging was carried out by two experienced endoscopists (CE, LG) using Fujinon EG-450HR and EG-410HR high resolution endoscopes (Fujinon Europe, Inc., Willich, Germany). In addition, chromoendoscopy with a special spraying catheter (Olympus PW-51; Olympus, Inc., Hamburg, Germany) was carried out in all patients, using methylene blue 0.5% in cases of Barrett’s oesophagus and Lugol’s solution 1% in cases of squamous cell cancer. After endoscopy, a standardised questionnaire was completed in which the endoscopist stated his assessment regarding the location of the tumour (oesophagogastric junction, distal, middle, or upper part of the oesophagus), its macroscopic classification according to the Japanese classification10 of early carcinomas (type I, polypoid; type IIa, elevated; IIb, flat; IIIc, depressed, or type III, excavated) and the combination of these types (for example, type IIa+IIb implies that the tumour has elevated and flat parts; type IIa+IIIc that the tumour has elevated and depressed parts; type IIb+IIIc that the tumour has flat and depressed parts; and type I+IIa that the tumour has a polypoid and an elevated part), as well as the depth of tumour infiltration (mucosal or submucosal). Polypoid, depressed, and excavated configurations and ulcerations were regarded as endoscopic signs consistent with a higher rate of submucosal infiltration.

### Endosonographic staging

The next day, HR-EUS using a 20 MHz miniprobe (Fujinon SP-501; Fujinon Europe, Inc.) as a “through the scope” probe with double channel video endoscopes (Fuji ED 410 D double channel; Fujinon Europe, Inc.) was carried out by an experienced endosonographer (EG). Approximately 100–200 ml water volumes were instilled through the second channel of the endoscope using a pump system (Olympus UWS-1; Olympus, Inc.). To avoid aspiration, patients were examined with an elevated position of the upper part of the body. The endosonographer was blinded to the results of the endoscopic evaluation, and his assessment of the depth of tumour invasion (mucosal/submucosal) was also recorded using a standardised questionnaire. In addition, conventional was carried out (Olympus GF-UM-20, 7.5 MHz, Olympus, Inc.) in order to assess lymph node status.
All endoscopies and endosonographies were carried out with the patient under analgesic sedation (midazolam 1–10 mg and/or pethidine 25–100 mg intravenously), with continuous monitoring of heart rate and oxygen saturation.

**Histology**

The gold standard used to establish the accuracy of endoscopic and endosonographic staging examinations was histological examination of the endoscopically or surgically resected tumour. The histological criteria used for early oesophageal carcinoma were the WHO criteria and the Vienna classification. Generally, the resected specimens were fixed in 4% formalin, dehydrated in gradual alcohol series, and embedded in paraffin. Serial step sections of 4 μm were cut and stained with haematoxylin and eosin and/or alcian blue, pH 2.5.

The histological assessment was carried out by experienced pathologists (MS, MV). After resection, specimens were examined with regard to tumour depth (involvement of the mucosa or submucosa)—the principal point of this study. Other histological assessment criteria were: freedom from invasion of the lymphatic vessels and veins; degree of differentiation (grade G1, well differentiated; G2, moderately differentiated; G3, poorly differentiated); and in some cases invasion of the lymphatic vessels and veins.

**Statistical evaluation**

Descriptive statistics were calculated for patient data and clinical parameters: mean, median, SD, and range (minimum–maximum). Paired proportions may also be shown as a contingency table. Although the table closely resembles the usual chi-square test distribution with one degree of freedom. The test of paired proportions is often known as McNemar’s test, especially when the data are shown as a 2×2 contingency table. Generalization for k×k tables is stated as “test of symmetry”.

**Table 3** Correct staging of submucosal carcinomas by high resolution endoscopic ultrasonography (HR-EUS) and HR-endoscopy (HR-E) in relation to the different submucosal types (sm1, sm2, and sm3)

<table>
<thead>
<tr>
<th>Submucosal type</th>
<th>HR-EUS</th>
<th>HR-E</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Correct</td>
<td>Not correct</td>
</tr>
<tr>
<td>sm1</td>
<td>4</td>
<td>8</td>
</tr>
<tr>
<td>sm2</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>sm3</td>
<td>6</td>
<td>3</td>
</tr>
<tr>
<td>Total</td>
<td>12</td>
<td>13</td>
</tr>
</tbody>
</table>

*sm1, first submucosal layer; sm2, second submucosal layer; sm3, third submucosal layer.

In addition, a kappa statistic was performed to test for agreement of the diagnostic methods. Here, a kappa value below 0.4 was regarded as representing poor agreement, a kappa value of 0.41–0.60 fair agreement, a kappa value of 0.61–0.80 good agreement, and a kappa value greater than 0.80 excellent agreement.

Statistical tests were performed using SAS software, version 8.2 (Cary, North Carolina, USA).

**Ethical considerations**

All patients provided written consent to undergo local therapy after receiving extensive information. All patients were informed that surgery was the current treatment of choice for early oesophageal carcinoma, and received a brochure describing endoscopic resection as an experimental procedure.

**Table 4** Accuracy of high resolution endoscopic ultrasonography (HR-EUS) and HR-endoscopy (HR-E) in early oesophageal carcinomas

<table>
<thead>
<tr>
<th>Proportion of correct results</th>
<th>Accuracy</th>
</tr>
</thead>
<tbody>
<tr>
<td>HR-E</td>
<td>64</td>
</tr>
<tr>
<td>HR-EUS</td>
<td>62</td>
</tr>
</tbody>
</table>

*94 patients were enrolled in the study, one patients showed an T2 carcinomas, therefore we have 93 early cancers. m, mucosa; sm, submucosal layer.

**Table 5** Correct staging of submucosal carcinomas by high resolution endoscopic ultrasonography (HR-EUS) and endoscopy (HR-E) in relation to the different locations of the carcinoma (oesophagogastric junction v tubular oesophagus)

<table>
<thead>
<tr>
<th>Histology</th>
<th>Oesophagogastric junction</th>
<th>Tubular oesophagus</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>m</td>
<td>sm</td>
</tr>
<tr>
<td>HR-EUS/HR-E</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>33/34</td>
<td>12/9</td>
</tr>
<tr>
<td></td>
<td>31</td>
<td>11</td>
</tr>
</tbody>
</table>

*One sm carcinoma was assessed as a T2 tumour by HR-E.

m, mucosa; sm, submucosal layer.

**Table 6** Correct staging of submucosal carcinomas by high resolution ultrasonography (HR-EUS) and endoscopy (HR-E) in relation to the different locations of the carcinoma (oesophagogastric junction v tubular oesophagus) and kappa values

<table>
<thead>
<tr>
<th>Kappa value</th>
<th>95% CI</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>HR-EUS/HR-E</td>
<td>0.042/0.346</td>
<td>0.226/0.322</td>
</tr>
<tr>
<td>HR-EUS/HR-E</td>
<td>0.211/0.296/</td>
<td>0.072/0.619</td>
</tr>
<tr>
<td>Other locations</td>
<td>0.836/0.776</td>
<td>0.656/0.106/</td>
</tr>
<tr>
<td>Test for equal kappa coefficients</td>
<td>p&lt;0.001/p=0.012</td>
<td>0.954/1.000</td>
</tr>
</tbody>
</table>

95% CI, 95% confidence interval.
treatment. If patients were interested in obtaining a second opinion, surgeons were regularly asked to discuss all questions once again with the patients.

Approval for the study was received from the ethics committee of the State of Hesse, Germany, in accordance with good clinical practice requirements.

RESULTS

Six of 100 patients had to be excluded from the study. Histological assessment was not possible in five patients with a suspicion of a large submucosal tumour as a decision was made to carry out palliative thermal destruction (APC in one of the five patients and Nd:YAG laser in the other four) due to concomitant diseases and/or age over 70 years. In one patient, carcinoma was not identified in the endoscopically resected specimen although histology of the biopsy in this area had shown a carcinoma. No cancer was found in subsequent check up endoscopies. The neoplasia had probably been so small that extensive biopsies had already removed it.

Endoscopic resection was carried out in 92 patients; five patients underwent surgical oesophageal resection. Three of these five patients were referred to surgery after pretherapeutic staging: submucosal tumour infiltration was seen in two patients, with suspicious lymph nodes in one; in the third patient, a T2 tumour was diagnosed endosonographically. In two of the five patients who underwent surgery, in whom at least one of the diagnostic procedures assessed a submucosal tumour, endoscopic resection was carried out first. Histological classification of the resected specimens not only confirmed a submucosal tumour but also revealed infiltration of the third submucosal layer (sm3). These two patients therefore also underwent surgery for oesophageal resection.

Detailed results with regard to macroscopic classification and histology of the cancers are given in table 1. Comparisons of the endosonographic and endoscopic staging results are given in tables 2 and 3. The overall sensitivity rates were 79.8% for and 82.9% for endoscopy. If the two diagnostic methods were combined and submucosal infiltration was assumed if one of the methods postulated submucosal infiltration, overall sensitivity increased to 84% (79/94); correct staging of submucosal cancers increased from 48% (with alone) and 56% (with endoscopy alone) to 60% if the two methods were combined. Submucosal infiltration was not diagnosed with either method in 40% of cases. The
HR-EUS was 79.6% (table 4).

Of tumour infiltration (the T category).\(^{12,15}\) With regard to the macroscopic classification, type I and type IIa+c lesions were the leading types of submucosal carcinoma: seven of 11 type I lesions (63.6%) showed submucosal infiltration, and submucosal infiltration was found in nine of 22 type IIa+c lesions (40%) (see figs 1–3 for examples). Submucosal infiltration was seen more rarely in type IIa lesions (4/28; 14%). Other types were rarely associated with submucosal infiltration (1/21 type IIb lesions; 5%), or the number of cases was too small for reliable assessment to be made (2/3 type IIc lesions; 1/5 type III lesions). A significant association was not observed between macroscopic type and depth of submucosal infiltration (sm1, sm2, or sm3). There was a trend towards more frequent sm3 tumours with type I lesions while in contrast sm1 lesions more frequently corresponded macroscopically to type IIa+c lesions. In this group of patients, type IIb lesions were only seen in sm1 tumours; a type IIb lesion was never seen in sm2 or sm3 tumours. Ulcerated type III lesions were never found in submucosal cancers of the sm1 type.

DISCUSSION

Conventional is currently the most accurate method for pretherapeutic assessment of lymph node status in oesophageal carcinoma, and is clearly superior to computed tomography and magnetic resonance imaging. In , ultrasound endoscopes with 7.5 MHz ultrasound probes are used either as radial or linear scanners. Conventional EUS is also currently the best diagnostic method for assessing the depth of tumour infiltration (the T category).\(^{12,15}\) However, conventional EUS systems are not ideal for assessing early neoplastic changes, and mucosal and submucosal carcinomas in particular, as the endoscopic view is limited and changes cannot be viewed directly. Early neoplastic lesions may also be pressed into the oesophageal wall by the filled balloon required for water preparation. In addition, conventional EUS systems use 7.5 MHz or maximally 12 MHz ultrasound probes, which are inferior to 20 MHz ultrasound probes with regard to resolution of the tissue layers in the near field.

These disadvantages of conventional EUS are not found with EUS miniprobes, which are therefore suitable for differentiating wall infiltration in T1 carcinomas. Miniprobes are applied under direct vision in a targeted fashion in the area of an endoscopically detected lesion, without pressing the lesion into the oesophageal wall. Using a double channel endoscope, allowing simultaneous working processes, sufficient liquid can be instilled through the second channel for water preparation in spite of active peristalsis in the oesophagus.

Larger studies on HR-EUS published in Western journals have mainly been by Asian research groups, and they have focused on early gastric carcinoma and early neoplasms of the large bowel.\(^{17,19,21,27}\) There have been only a few reports, some with small numbers of patients, on EUS and particularly HR-EUS with the miniprobe in early oesophageal cancer.\(^{16,19,20,26}\) For example, Murata and colleagues\(^{16}\) reported correct differentiation between mucosal and submucosal carcinomas using a 20 MHz probe in 94% of cases (46/49 lesions). The overall accuracy in the 54 patients investigated was 75%. Yanai and colleagues\(^{16}\) found an accuracy of 72.3% in staging 47 mucosal or submucosal lesions; the overstaging rate was 19% and the understaging rate was 2%.

With 100 patients, the present study is currently the largest investigation, providing a prospective assessment of the value of HR-EUS with miniprobes in early oesophageal carcinoma and comparing the results with histological findings after endoscopic resection. The endosonographic accuracy of early
oesophageal carcinomas showed good results at approximately 80%, a value comparable with that in a Japanese study of early gastric carcinoma and in a study of early oesophageal carcinoma where Yanai et al reported an accuracy of 72.3% and 64.7% (80% for mucosal carcinoma). Analysis of submucosal carcinomas, representing approximately one quarter of early oesophageal carcinomas, is all the more disappointing. For these lesions, sensitivity fell to 48% (with a specificity of 91%). Two factors appear to have played a part here: location of the carcinomas and depth of infiltration. Only two of 14 carcinomas located directly at the oesophagogastric junction were correctly assessed, while nine of 10 carcinomas in the rest of the oesophagus were correctly staged. Technical difficulties due to the anatomical conditions in the area of the cardia, particularly in the presence of an axial hiatal hernia, certainly played a major role. Data from the literature confirm this: de Manzoni and colleagues reported an overall accuracy of 55.2% for EUS assessment of adenocarcinoma of the cardia, with T2 tumours showing particularly poor results. Kelly and colleagues, on the basis of a literature search conducted for their review article, similarly stated that the accuracy of EUS is lower for carcinomas of the cardia. It is often not possible to achieve adequate water preparation by instilling fluid through the second channel. In addition, there is substantial motility that prevents dilation of the distal oesophagus from being maintained for a longer period. With regard to water preparation, a condom-like sheath into which water could be introduced from outside for preparation might be helpful. Initial prototypes with a total diameter of 2.6 mm, capable of being introduced through a 3.2 mm instrument channel, have already been successfully tested in clinical practice.

The second factor is the superficial—often focal—infiltration of the uppermost layer of the submucosa, known as type sm1. Eight of 12 sm1-type lesions were not correctly diagnosed; all were located in the distal oesophagus (n = 1) or directly at the oesophagogastric junction (n = 7), again underlining the importance of tumour location. The fact that the sm1 type represents an independent “risk factor” has already been noted by Shimizu and colleagues. In addition, concomitant inflammatory changes in the oesophageal mucosa—which are often seen in reflux induced Barrett’s oesophagus despite administration of proton pump inhibitors—can make the assessment very much more difficult. As described by Shimizu and colleagues, overall sensitivity improves if sm1 carcinomas are included in the mucosal group. If it is assumed on the basis of the data presented by Gotoda and colleagues that sm1-type cancers are endoscopically resectable with no risk, or a very low risk, of lymph node metastasis under certain conditions, patients with histologically confirmed tumour infiltration of the first submucosal layer (sm1-type) could be assigned on clinical grounds to the group of patients with mucosal cancers. On this assumption, the overall correct staging of endoscopic and endographic staging would increase to 89% and 86%, respectively. In the subgroup of submucosal carcinomas, the correct assessment would increase from 48% to 80%. The best diagnostic yield was achieved when the two diagnostic methods were combined, as has already been postulated by Ohashi and colleagues. The overall rate of correct staging was then 91.5% (86/94), with correct assessment of submucosal carcinomas at 88% (22/25), provided that submucosal infiltration was assumed to be present if one of the two diagnostic procedures identified it.

On the basis of the data presented by Gotoda and colleagues on early gastric carcinoma, this regrouping of sm1 carcinomas using clinical criteria appears to be worth considering. A follow up study of more than 5200 patients with early gastric carcinoma who had undergone surgery showed that no lymph node metastases were found when the lesion was type sm1 without evidence of infiltration of lymphatic vessels and a maximum size of up to 30 mm. This means that gastric sm1 carcinomas can receive the same type of endoscopic treatment as mucosal carcinomas under certain conditions. Shimizu and colleagues recently presented long term results on this topic regarding endoscopic treatment of early squamous cell carcinomas that had infiltrated the muscularis mucosae (m4) or had already penetrated the submucosa (sm1). Five year survival rates (YSR) and tumour specific five YSR for patients who had received endoscopic treatment did not differ significantly from those in the control group who had undergone surgical resection. The extent to which these data might be generally applicable to early oesophageal carcinoma of Barrett’s type remains an open question.

The results of endoscopic staging, which with an overall sensitivity of 82.9% and an accuracy of 83.4% are even slightly superior to those of EUS (79.8% and 79.6%), are impressive. However, an absolute prerequisite is that the endoscopist should have a substantial amount of experience with early carcinomas and high resolution endoscopes. Endoscopy is also more sensitive than EUS in the assessment of submucosal carcinomas (56% vs 48%, respectively). These results are not entirely in agreement with Japanese data from a comparison of gastric carcinoma in which endoscopy, with an accuracy of 63%, was inferior to EUS at 71%. No tendency towards overstaging with EUS was found in the present study; on the contrary, submucosal carcinomas tended to be rather underestimated.

In conclusion, overall correct diagnostic assessment of both HR-E and HR-EUS with a 20 MHz miniprobe in early oesophageal cancer is high, with no significant differences between the two techniques. Combining HR-E and HR-EUS can increase sensitivity. HR-E and HR-EUS provide a high level of diagnostic accuracy for mucosal and submucosal tumours located in the tubular part of the oesophagus. With submucosal tumours located at the oesophagogastric junction or with infiltration of the first third of the submucosa however, the diagnostic meaningfulness of both techniques is not yet satisfactory.

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