Diagnosis and treatment of superficial esophageal cancer

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Abstract
Endoscopy allows for the screening, early diagnosis, treatment and follow up of superficial esophageal cancer. Endoscopic submucosal dissection has become the gold standard for the resection of superficial squamous cell neoplasia. Combinations of endoscopic mucosal resection and radiofrequency ablation are the mainstay of the management of Barrett’s associated neoplasia. However, protruded, non-lifting or large lesions may be better managed by endoscopic submucosal dissection. Novel ablation tools, such as argon plasma coagulation with submucosal lifting and cryoablation balloons, are being developed for the treatment of residual Barrett’s esophagus, since iatrogenic strictures still hamper the development of extensive circumferential resections in the esophagus. Optimal surveillance modalities after endoscopic resection are still to be determined. The assessment of the risk of lymph-node metastases, as well as of the need for additional treatments based on qualitative and quantitative histological criteria, balanced to the patient’s condition, requires a dedicated multidisciplinary team decision process. The need for trained endoscopists, expert pathologists and surgeons, and specialized multidisciplinary meetings underlines the role of expert centers in the management of superficial esophageal cancer.

Keywords
Superficial esophageal neoplasm, endoscopic resection, endoscopic submucosal dissection, radiofrequency ablation, Barrett’s esophagus, squamous cell carcinoma, early adenocarcinoma

Introduction
Esophageal cancer is the eighth most common cancer and the sixth most common cause of cancer death, with estimations of 456,000 new cases and 400,000 deaths in 2012 [1]. Squamous cell carcinoma (SCC) is the most frequent histological subtype because of its high prevalence in Asia, with up to 95% of cases of esophageal neoplasia in China [2], and persistent alcohol and tobacco consumption in Western countries [3]. In America and Northern Europe, however, esophageal adenocarcinoma (EAC) arising on Barrett’s esophagus has recently become the most prevalent type of esophageal cancer [4,5]. Upper gastrointestinal endoscopy screening programs in Asia and surveillance protocols for Barrett’s esophagus have allowed a greater proportion of early stage, T1—so-called “superficial” or “early” esophageal cancers—to be diagnosed. Mucosal (T1a) cancers include high-grade dysplasia (HGD), or in situ carcinoma (T1am1), and tumors infiltrating the lamina propria (T1am2) or the muscularis mucosae (T1am3). Submucosal cancers (T1b) can be classified as T1bsm 1, 2 or 3, depending on the invasion of the upper, middle, or deepest third of the submucosa on surgical specimens [6,7], or as superficial submucosal invasion (less than 200 µm from the muscularis mucosae for SCC and less than 500 µm for EAC) [8]. All T1 lesions are technically amenable to endoscopic resection (ER), but two caveats should immediately be raised: first, the risk of severe, refractory esophageal stricture following resections exceeding three fourths of the circumference [9], requiring repeated dilatations with a significant risk of perforation [10], has to be taken into account before performing extensive ER, given the lack of efficient options to prevent esophageal stricture [11]. Second, the risk of lymph-node metastases has to be assessed based on the resected specimen to determine whether an ER has been curative or not. Esophagectomy, which remains a standard treatment for esophageal neoplasms, should now be restricted to lesions extending to the muscularis propria (T2 and over) and to superficial lesions bearing the highest risk of lymph-node metastases. Indeed, ER has proven equally effective in terms of oncological outcomes, with no procedure-related mortality, a reduced morbidity, and a preserved quality of life [12,13]. Therefore, endoscopy is currently the cornerstone of the management of superficial esophageal
cancer, allowing for screening, diagnosis, treatment- and follow up in most cases.

**Diagnostic workup**

**Endoscopy**

The vast majority of esophageal neoplasms are diagnosed at an advanced stage, in which the endoscopist encounters an irregular esophageal stenosis or tissular bud with an ulcerated surface and takes biopsies for histological documentation and therapeutic decision-making. The detection and characterization of superficial neoplasia can be challenging, since lesions typically appear as subtle and flat, Paris 0-1 Ib type. High-definition white-light endoscopy is a minimal requirement [8], and is usually associated with ancillary techniques such as dyes or so called virtual chromoendoscopy. Other prerequisites are the involvement of an experienced endoscopist and enough time for inspection of the mucosa. Noticeably, Gupta et al showed that a 1-min inspection time per longitudinal centimeter of Barrett's esophagus was associated with a higher rate of neoplasia detection [14]. The description of the lesions should include their size, Paris classification type, extension over the esophageal circumference and precise location (e.g., 5 o'clock) and distance from the dental arcade. The extent of the Barrett's esophagus should be reported using the Prague classification [15], which reports the circumferential (C) and the maximal extent (M) of the Barrett's segment, measured from the upper end of the gastric folds.

For the detection and delineation of SCC or squamous dysplasia, Lugol staining has long been the gold standard, with a per-lesion sensitivity of 88%. Narrow-band imaging (NBI; Olympus, Tokyo) is a virtual chromoendoscopy technique that has shown sensitivity comparable to that of Lugol, and a superior specificity of 82% vs. 37% [16]. Indeed, cicatricial or parakeratotic lesions may appear as Lugol-negative and therefore suspicious, while NBI will show a benign pattern of the intrapapillary capillary loops (IPCL). Since there is no such feature as a pit pattern in the squamous epithelium, the microvascular pattern of the IPCL can be used to predict the degree of malignancy and invasiveness of the lesion [17]. A classification developed by Inoue et al subdivides the IPCL into 5 types, with type V (neoplastic) being subdivided into 4 subtypes: the last one, with neovessel formation, suggests deep submucosal infiltration, proscribing ER. However, discrepancies between endoscopic biopsies and resection specimens occur in about a third of cases, and en bloc ER should remain the gold standard for the diagnosis [8].

**Endoscopic ultrasound (EUS)**

The role of EUS in the initial staging of esophageal neoplasia remains controversial. A recent meta-analysis found high sensitivity and specificity of EUS for the diagnosis of T1a lesions of 0.85 (95% confidence interval [CI] 0.82–0.88) and 0.87 (95% CI 0.84–0.90), respectively [24], but about 80% of the patients were investigated with an EUS miniprobe, unavailable in most centers. Conventional EUS probes do allow for an accurate differentiation between T1 and T2 stages, but this is generally not clinically relevant, since T2 lesions can usually be recognized endoscopically as a protruding type, with ulcerated features, large neovessels, and limited motility with esophageal peristalsis. However, EUS is able to detect regional lymph nodes with a sensitivity of 76% and a specificity of 72% [25], significantly better than computed tomography (CT) scan [26]. EUS-guided fine-needle aspiration can then increase the diagnostic accuracy of the N-staging to 90% in advanced carcinomas [27]. However, in a retrospective analysis of 131 patients with early Barrett’s neoplasia, Pouw et al showed that EUS alone never impacted treatment policy [28]. Finally, EUS is not routinely recommended in the workup of all early esophageal cancers [8]: the guidelines of the European Society of Gastrointestinal Endoscopy (ESGE) suggest limiting its use to high-risk lesions, in addition to CT scanning and positron emission tomography-CT.

**Advanced endoscopic techniques**

Advanced diagnostic endoscopic techniques, such as autofluorescence imaging, confocal laser endomicroscopy or optical coherence tomography, have been widely studied,
mainly for the detection of Barrett’s neoplasia. Autofluorescence imaging has failed to prove a significant improvement in the detection of early neoplasia [29]. The development of probe-based confocal laser endomicroscopy is limited by its high cost and low sensitivity [30]. Optical coherence tomography is a promising tool for scanning the entire Barrett’s mucosa; however, it is not yet ready for implementation in clinical practice [31]. Currently, no advanced endoscopic imaging technique is recommended, and targeted biopsies or resection of any visible lesion are advisable, in addition to random quadrantic biopsies in the case of Barrett’s esophagus, according to the Seattle protocol [32].

Endoscopic management of SCC

ER

Endoscopic mucosal resection (EMR) has been widely studied since the 1990s for the treatment of early SCC. EMR can be conducted using submucosal injection, an oblique cap and an asymmetric resection snare [33], or a modified multiband ligator device allowing the passage of an electrosurgical snare [34]. The latter technique has proven as effective as the first, but is cheaper and faster, with a complication rate lower than the 1-2% perforation rate usually observed with cap-EMR [35]. All studies report consistent data, with en bloc resection rates ranging from 20-53%, and local recurrence from 10-26% [9,36,37]. Indeed, the largest lesion amenable to en bloc resection with the EMR cap or ligator device is about 15 mm. Conversely, endoscopic submucosal dissection (ESD) allows for en bloc resection of neoplastic lesions irrespective of their size [38] Figure 1. Since 2005, over 15 studies have reported outcomes of ESD for esophageal SCC in more than 700 patients, with en bloc resection rates of 95.1% and histologically complete resection rates of 89.4%. These results were reproduced in several expert Western centers [39-41]. Complications after endoscopic ESD for SCC include bleeding within the first 24 h in 2.1% of patients and perforations in 5%, both treated conservatively in the vast majority of cases, while strictures occur in 11.2%, mainly after treatment of lesions exceeding 75% of the esophageal circumference or extending beyond m2 [41,42]. No procedure-related mortality was reported after ESD or EMR.

No randomized study has compared EMR and ESD. Takahashi et al compared the outcomes of 184 lesions treated by EMR to 116 lesions treated by ESD and found a significantly higher R0 resection rate with ESD (97.4% vs. 78.3%, P=0.0002) and a lower local recurrence rate (0.9% vs. 9.8%, P=0.0065) [37]. In addition, Cao et al reported that ESD provided a significantly higher en bloc resection rate than EMR, also for lesions smaller than 10 mm [43]. These data, along with the optimal histological assessment of the risk of lymph-node involvement allowed by a single large resection specimen, have made ESD the current standard for ER of esophageal SCC [8] (Table 1).

Endoscopic ablation

Radiofrequency ablation (RFA) probes deliver a 465-kHz energy waveform via a bipolar arrangement of electrodes over a wire-guided balloon or on an articulated over-the-scope distal attachment (HALO system, Medtronics). The safety and efficacy of RFA have been established for the eradication of early neoplasia in Barrett’s esophagus in numerous studies, including three randomized controlled trials [47-49]. Thus, RFA has been proposed for flat early squamous neoplasia, including early SCC, with encouraging results, including 84% complete eradication at 12 months, 2% neoplastic progression managed endoscopically, and a 20% stricture rate after a median of 2 RFA sessions [50]. However, endoscopic recognition of T1a m1-2 lesions amenable to RFA, as opposed to deeper lesions requiring ER or surgery because of potential lymph-node metastases, has proven to be difficult, even for expert endoscopists [51]. Furthermore, the costs associated with RFA remain high. Therefore, the use of RFA in early squamous neoplasia is currently not recommended.

Canto et al recently reported the results of balloon-based cryoablation of early squamous neoplasia, including non-neoplastic Lugol-unstained lesions, with a high efficacy at 1 year and a good safety profile [52]. However, as with RFA, the risk of mistaking an invasive SCC for a dysplastic lesion and ablating instead of resecting it is likely to limit the indications of cryoablation or any ablative technique in the squamous epithelium.

Surveillance after resection

Since ESD has become the standard treatment for superficial SCC, the risk of local recurrence has become virtual in the case of R0 resection. However, the risk of metachronous cancer in the esophagus reaches 20% at 5 years [53] and the risk of distant metastases justifies endoscopic surveillance after resection. No prospective, large-scale study has yet assessed the risk of lymph-node metastases in early SCC. In one of the largest reports, which included over 500 early SCCs treated endoscopically and followed over 5 years, the cumulative risk of lymph-node metastases ranged from 0.4% for T1a m1-2 SCC, 8.7% for T1am3, 7.7% for T1bsm1, to 36.2% for T1bsm2 lesions [53]. These numbers are in line with those reported by other teams, including surgical series, with a 0-2% risk of lymph-node metastases for T1am1-2 lesions, 8-18% for lesions invading the muscularis mucosae (m3), and 7.7 to 19% for lesions invading the submucosa to less than 200 μm (sm1), and 30-54% for sm2 or sm3 lesions [6-8,53]. These figures have to be interpreted with caution for m3 and sm1 lesions, since most of these lesions also have poor qualitative histoprognostic factors, such as poor differentiation or lymphovascular involvement, while well differentiated m3 or sm1 tumors without these poor histoprognostic factors rarely cause lymph-node metastases [54]. As a result, yearly endoscopic surveillance is recommended after R0 resection of a T1am1-2 SCC. For m3 or sm1 tumors, the decision is
made during a multidisciplinary meeting, weighing the risk of lymph-node metastases against the individual assessment of esophagectomy-associated morbidity and mortality. Clear-cut follow-up modalities have not been defined, but should probably include an esophagscopy after 3 months and yearly thereafter, combined with endoscopic ultrasonography and six-monthly CT scan, as for surgically resected esophageal cancer. For sm2 or sm3 tumors, ER is not curative and esophagectomy with lymph-node resection is recommended [8].

In the case of R1 resection on the horizontal (lateral) margins, recurrences can usually be managed endoscopically and close endoscopic surveillance is recommended [8]. A surveillance protocol with an esophagscopy every 3 months for 1 year, then yearly, can be considered. Conversely, positive vertical margins suggest deep mural invasion and prompt surgical resection if the patient's clinical status is compatible.

### Endoscopic management of adenocarcinoma

#### ER

ER is the first step in the treatment of early adenocarcinoma and Barrett's neoplasia but can also be the last step of the diagnostic workup. The standard EMR technique requires a large straight or oblique cap of 12 mm inner diameter with a distal ridge, allowing for the placement of an asymmetrical snare after submucosal lifting. Another widely used technique is the multiband EMR technique, which uses a modified variceal band ligator and a 5-7F polypectomy snare. A randomized controlled study comparing both techniques showed that multiband EMR was cheaper and quicker than cap-EMR, but similarly safe, despite the absence of submucosal lifting [55]. In accordance with these data, Alvarez Herrero et al, in a large prospective study including 1060 ER using multiband EMR, recorded no perforations [56], and even larger studies have confirmed the safety and efficacy of EMR for intramucosal EAC [57]. Finally, diagnostic and therapeutic ER using multiband EMR for all visible abnormalities arising on a Barrett's mucosa can be preferred to biopsy sampling in certain cases, especially since biopsies can downgrade EAC to HGD in up to 40% of cases [8].

In contrast to squamous neoplasia, ESD has not been accepted as the gold standard for Barrett's neoplasia. Indeed, the principle of ESD—allowing for en bloc resection of lesions irrespective of their size, at the cost of longer procedures, and a longer learning curve—is hardly compatible with the extensive and often multifocal nature of Barrett's neoplasia. As a result, prospective studies of ESD for Barrett's neoplasia have shown disappointing results, with 39-74% histologically complete resection rates for HGD or EAC and 48-96% curative resection rates for EAC (Table 2). No mortality was associated with ESD for EAC, while 1.1-2.7% intraprocedural bleeding and 2.3-4% perforation rates were reported [8]. However, the vast majority of the perforations are diagnosed and closed during the endoscopic procedure and treated conservatively. Although the latest ESGE guidelines recommended that piecemeal ER should be preferred over ESD in most cases [8], the recent development of ESD in western centers that specialize in Barrett's esophagus may change therapeutic paradigms in the near future. Until multicenter prospective studies have further demonstrated its efficacy and superiority over EMR, ESD in Barrett's neoplasia should probably be limited to expert centers and selected cases, such as protruding lesions not amenable to cap or multiband EMR, lesions larger than 15 mm in size, and poorly lifting lesions or those with a high suspicion of submucosal invasion.

#### Endoscopic ablation

##### RFA

The first step in the management of a patient with Barrett's neoplasia is an ER of all visible lesions, since they have a probability of harboring HGD or EAC close to 80% [68]. Complete eradication of the remaining Barrett's mucosa is indicated, because 30% of the patients will experience a recurrence of neoplasia during follow up [69]. However, circumferential resection of the Barrett's mucosa results in an esophageal stricture in up to 88% of patients, with a 4% recurrence rate of neoplasia at 24 months [48]. A randomized study has shown that focal ER, followed by ablation of the remaining Barrett's mucosa, was equally effective and safer compared to stepwise ER [48]. The efficacy of this treatment
approach has been confirmed by several larger prospective studies [70,71] that achieved 90% complete eradication of intestinal metaplasia and neoplasia at 5 years [71]. Finally, EMR of visible lesions and RFA of the remaining Barrett’s esophagus is currently the gold standard in the treatment of Barrett’s neoplasia Figure 2. This is particularly true for long circumferential Barrett’s esophagus, in which other ablative techniques can be time-consuming and less effective.

When HGD is diagnosed on random biopsy samples without any visible lesion to resect, and after confirmation of HGD by an expert pathologist, ablation of the complete Barrett’s segment is indicated [32]. Recent international guidelines have recommended offering endoscopic therapy, including ablative therapies, for Barrett’s esophagus containing confirmed LGD [32,72]. However, because evidence is supported by only one randomized trial, in which LGD in the surveillance group had an unexpectedly high rate of progression to EAC [49], this recommendation must be treated with caution: as rightfully stated by the Bobcat consortium and ESGE, RFA can be used in Barrett’s esophagus with LGD containing high risk features, such as multifocality of LGD, confirmation of LGD by repeat, expert-analyzed biopsies, focal visible lesion or long-segment Barrett’s esophagus.

**Other techniques**

Argon plasma coagulation (APC) is a widely available and relatively cheap ablative endoscopic therapy for residual Barrett’s esophagus. Initial studies demonstrated a reduction in the recurrence rate of neoplasia from 37% to 3% at 2 years after ER [73]. However, APC is operator-dependent, time-consuming (particularly for larger Barrett’s segments) and may result in a 9% stricture rate. A recently developed APC probe allowing for submucosal injection of 0.9% saline prior to a high-power ablation, called Hybrid-APC™, could improve the safety and efficacy of APC ablation [74].

Finally, balloon based cryoablation has recently been reported as an effective modality for ablating residual Barrett’s islands after ER [75]. Studies reporting the outcomes of cryoablation performed on larger Barrett’s segments are expected.

**Surveillance after resection**

As mentioned above, the risk of local or metachronous recurrence of EAC is largely reduced by the systematic ablation of the residual Barrett’s esophagus. However, close endoscopic surveillance and prolonged follow up is still recommended by current guidelines [32].

The risk of lymph-node metastases associated with EAC can be estimated according to quantitative factors, such as the depth of tumor infiltration, and qualitative factors, such as the differentiation grade and the presence of lymphovascular involvement. ER can be considered curative for EAC invading the mucosa or the superficial submucosal up to 500 µm, provided they are not poorly differentiated and do not show lymphovascular involvement [8]. For these lesions, the risk of lymph-node metastases is below 2%. In the case of deeper infiltration in the submucosa, positive vertical margins, or the presence of poor qualitative histoprognostic features, the risk of lymph-node metastases exceeds 10% and additional therapy by esophagectomy with lymph-node dissection should be discussed. In the case of positive horizontal margins, additional ER or ablative therapy can be offered.

**Surgery**

Surgery has long been the gold standard for the treatment of localized esophageal cancer, including early esophageal cancer. However, esophagectomy results in 2-5% mortality and 30-40% severe morbidity, and significantly alters the patients’

Table 2 Outcomes of the main studies assessing endoscopic submucosal dissection for Barrett’s associated neoplasia

<table>
<thead>
<tr>
<th>First author</th>
<th>Number of patients</th>
<th>Mean diameter of the lesion, mm</th>
<th>Mean procedure time, min</th>
<th>R0 resection rate for EAC/HGD</th>
<th>Curative resection rate for EAC/HGD</th>
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<th>Stricture rate</th>
<th>Recurrent or metachronous neoplasia</th>
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<td>Nagami</td>
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<td>Kagemoto</td>
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<td>Probst</td>
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<td>Cheaux</td>
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<td>Höbel</td>
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R0, histologically complete resection; HGD, high-grade dysplasia; EAC, early adenocarcinoma; NA, not available
quality of life for at least one year [76-78]. However, surgical resection of a T1 esophageal cancer is associated with an overall 5-year survival of 77-88%, with a 19-43% 5-year risk of tumor recurrence [79-81]. Comparison of the outcomes of surgical and ER treatment lacks well designed prospective studies. Most data pertain to EAC, and in the only large cohort including SCC patients both treatment groups were hardly comparable, with over three times more T1b lesions in the surgery group [81]. All studies resulted in comparable oncological outcomes, with greater postprocedural mortality and morbidity induced by surgery [13,81,82].

Chemoradiotherapy (CRT)

CRT and surgery have comparable efficacy in the treatment of locally advanced esophageal cancer [83]. Therefore, several teams have assessed CRT alone for early esophageal neoplasia. Protocols are usually the same as for advanced esophageal cancer and typically involve 45 to 50 Gy external beam radiotherapy over a 5-week period and 2 cycles of chemotherapy based on cisplatin and 5-fluorouracil.

CRT alone was also studied for early esophageal cancers: in a study involving 37 stage I esophageal cancer, of which 30 were T1b lesions, Jingu et al reported a poor cancer-specific survival rate of 87.6% at 5 years [84]. In a larger study that included 72 patients with T1 esophageal cancer treated by CRT alone, Kato et al found 68.1% 4-year recurrence-free survival, excluding local 15% lesions recurring locally and treated by ER [85]. Nemoto et al, in a large multicenter study that included 141 patients with T1 esophageal cancers treated by CRT or radiotherapy alone, found a 3-year overall survival of 90% and 70% for patients with T1a and T1b lesions, respectively, and better outcomes with CRT compared with radiotherapy alone [86]. However, it has to be noted that the treatment regimens were extremely heterogeneous in terms of radiotherapy dose and type, and the administration of chemotherapy concurrent with radiotherapy. The combination of external-beam radiotherapy and high dose brachytherapy for superficial esophageal cancer has also been reported outside Japan, in 66 patients with T1 esophageal cancers, with 5-year relapse-free survival and cause-specific survival of 54.6% and 76.9% [87]. Finally, the quality of the data on CRT for the treatment of superficial esophageal cancer is currently insufficient to draw definitive conclusions. Available data suggest that CRT, possibly including brachytherapy as an adjunct to external-beam radiotherapy, is feasible and safe in inoperable patients. However, the relapse-free and cause-specific survival rates are far below those of surgery or ER. Furthermore, although ER remains feasible after CRT, the curative resection rates barely exceed 50% and the complication rate is unusually high, mainly because of a higher occurrence of esophageal strictures [88].

A promising role for CRT could be to replace surgical resection after non-curative ER. This treatment strategy, however, already recommended in the ESGE guidelines and performed in expert centers, relies only on small retrospective case series [89-91].

Perspectives

Diagnosis

Currently, the indication for ER relies on a fine endoscopic assessment of mucosal anomalies, and the prediction of the malignant potential is based on architectural features and the presence of dysplasia on histology. The detection of molecular alterations of the Barrett's epithelium could be a useful tool for risk stratification and prediction of response to therapy. For example, p53 aberrant expression (either absent or overexpression) detected by immunohistochemistry has been extensively studied, and could be as discriminative as the presence of LGD in the prediction of neoplastic progression [92,93]. Furthermore, analysis of the p53 expression pattern could help in improving the diagnostic accuracy of esophageal biopsies and lowering the rate of “indefinite for dysplasia” [94]. Specific genetic alterations of the MYC, p16, Her-2/neu, or ZNF217 genes, detected by fluorescence in situ hybridization on cytological brush samples, are associated with a lower rate of response to endoscopic therapy [95]. The recent development of a transoral brush cytology device that allows for RNA extraction and diagnosis by microarray, and quantitative polymerase chain reaction testing for specific genetic biomarkers of Barrett's esophagus and early neoplastic changes, might lead clinicians in the future to switch from endoscopic examination and quadrantic biopsies to transoral unsedated cytological brushing of the entire Barrett's segment. This might help first to overcome the sampling bias; second to assess the genetic diversity among Barrett's cells, better predict their malignant potential and determine tailored surveillance intervals; and finally to detect molecular changes prompting ER or ablation, before any macroscopic or microscopic change can be seen [96,97].

Treatment

Although ablative therapies offer a safe treatment modality for residual Barrett's esophagus, their application for early squamous cell neoplasia is limited, their long-term efficacy is unknown, and their costs are high. Therefore, epithelium-regenerating, anti-inflammatory or antifibrotic drugs or devices are under development in order to prevent post-endoscopic esophageal stricture, and possibly allow for a single-step ER.
Superficial esophageal cancer

Figure 1 Endoscopic submucosal dissection of an early squamous cell carcinoma (T1am2). (A) High-definition white-light endoscopy visualization of a Paris 0-IIb lesion of the mid esophagus. (B) Narrow-band imaging and (C) Lugol coloration showing the limits of the lesion. (D) Narrow-band imaging with magnification showing the type V-2 intrapapillary capillary loops suggesting m2, resectable lesion. (E) Circular markings before endoscopic submucosal dissection. (F) Distal incision. (G) Submucosal dissection using the tunnel technique under the lesion. (H) Resection wound after en bloc endoscopic submucosal dissection. (I, J, K) Three-month follow-up endoscopy showing a clean esophageal, Lugol-negative scar, without evidence for recurrence or residual neoplasia.

Figure 2 Endoscopic treatment of early Barrett’s neoplasia (T1am1). (A) High-definition white-light endoscopy showing a visible abnormality with nodularity and irregular nodularity and irregular pit pattern on a short Barrett’s tongue. (B and C) Narrow-band imaging of the lesion in direct and retroflex view. (D) Band ligation of the lesion without submucosal lifting, before (E) placement of the snare below the band, and (F) resection wound after multiband mucosectomy. (G) Radiofrequency ablation using a focal probe to ablate residual Barrett's esophagus, 3 months after endoscopic mucosal resection. (H and I) Follow-up endoscopy 3 months later, showing a normal-appearing neo-Z line under white-light endoscopy (H) and narrow-band imaging (I).
of large esophageal neoplasms or of the complete Barrett’s esophagus [11].

Finally, the surgical risk of esophagectomy in the case of non-curative ER for a superficial esophageal cancer may be excessive for frail or elderly patients, or patients with a low risk of lymph-node metastases: therefore, alternatives to surgical resection, such as esophagus-preserving lymphadenectomy [98] or “adjuvant” chemoradiotherapy, are under study [89-91].

Concluding remarks

Endoscopy allows for the screening, early diagnosis, treatment and follow-up of superficial esophageal cancer. In squamous cell neoplasia, endoscopic therapy consists in ESD, while EMR followed by the ablation of the remaining Barrett’s esophagus remains the recommended treatment approach for early adenocarcinoma. Optimal surveillance modalities after ER are still to be determined. The need for trained endoscopists, expert pathologists, and dedicated multidisciplinary meetings underlines the role of expert centers in this pathology.

References


