Barrett’s esophagus: lessons from recent clinical trials

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Abstract
Data from recent studies cast doubt on former recommendations on diagnosis and management of Barrett's esophagus. Based on latest research findings several Gastroenterological Associations actualized their guidelines and international experts compiled consensus statements as practical help for clinicians. In this review we discuss recent trials and their impact on clinical practice, current recommendations and persisting controversies in Barrett's esophagus.

Keywords Barrett’s esophagus, esophageal adenocarcinoma, endoscopic eradication, endoscopic mucosal resection, endoscopic submucosal dissection

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Introduction
Barrett’s esophagus (BE) and its underlying condition, gastroesophageal reflux disease (GERD), predispose to esophageal adenocarcinoma (EAC), a tumor whose incidence has risen dramatically in Western countries during the past decades (in the United States more than 6-fold in 40 years from 0.4 cases per 100000 in 1975 to 2.6 cases per 100000 in 2009 [1]). The prognosis of advanced tumor is poor with a 5-year survival for distant staged disease of only 2.8% [1]. If early carcinoma is detected the patient may be offered a potentially curative endoscopic resection (ER), or, if dysplasia is detected, endoscopic ablation to prevent progression to cancer. Hence, screening and surveillance for BE seem rational. Several studies showed that endoscopic surveillance leads to carcinoma detection at earlier stages and to more favorable survival [2]. However, recent studies also showed that the incidence of cancer and the risk of malignant progression among patients with non-dysplastic BE is considerably lower than previously thought [3-5]. Low-grade dysplasia (LGD) on the other hand seems to be an overdiagnosed but underestimated entity [6]. In the past years, tremendous advances evolved as well in ER and ablation techniques as in endoscopic imaging. But is there enough evidence to change practice and what are the lessons learned from recent studies to reconsider diagnostic and therapeutic strategies?

Epidemiology and cancer risk: should we perform screening?
Endoscopic screening is a controversial issue. The primary goal of screening is to identify patients with BE who will benefit from surveillance or therapy to prevent EAC. But first of all who actually should be screened? Known risk factors for BE and EAC are GERD, male sex, white race, older age, obesity, metabolic syndrome, tobacco use, hiatal hernia and a family history of GERD, BE or EAC [7]. The American Gastroenterological Association (AGA) recommends screening for BE in individuals older than 50 years with symptomatic GERD and at least 1 additional risk factor for EAC [8]. There is no definitive study that supports the assumed benefit of this strategy. But the major dilemma is that a significant proportion of patients with BE and EAC lack reflux symptoms. Approximately 50% of patients with short-segment BE deny GERD symptoms and 40% of patients with EAC reported no history of prior GERD [9,10]. Also there are different opinions about the clinical importance of short BE. Another consideration that diminishes the usefulness of screening is the very low risk of malignant progression in non-dysplastic BE. Recent population based studies and large meta-analysis showed an annual cancer incidence of only 0.1-0.3% in these patients and the risk even seems to further decrease over time with follow-up endoscopies showing no progression to dysplasia [3-5,11]. All in all, it is currently difficult to clearly identify the population at risk and more accurate methods for risk stratification are needed. Molecular biomarkers and non-endoscopic technologies for cell collection may help us in the future [12-14]. Promising results have been obtained with the Cytosponge, a cell collection device composed of reticulated foam compressed within a gelatin capsule attached to a string. The capsule is swallowed by the patient and, after 5 min, allowing the dissolution of the gelatin and expansion of the foam, the sponge is retrieved by the operator. During the passage of the sponge cells are absorbed for immunohistochemical analysis.
In a feasibility study the Cytosponge detected BE >1 cm with 73% and BE >2 cm with 90% sensitivity and a specificity of >90% [15]. The results of a large multicenter study (BEST2) will provide further information on the diagnostic accuracy. A simulation model, screening a hypothetical cohort of 50-year-old men with GERD symptoms, showed that Cytosponge screening followed by endotherapy reduces EAC mortality and is cost effective [16]. Accurate, minimal invasive and cost-effective screening tools may soon be available for clinicians. Up to now, the effectiveness of endoscopic screening is debatable and there are variable recommendations on it amongst different Medical Societies.

Definition of BE: do we require goblet cells?

In BE, as a consequence of GERD, the squamous epithelium that normally lines the distal esophagus is replaced by a metaplastic columnar epithelium. Endoscopically this is characterized by the typical salmon color and coarse texture. Histologically it is characterized by specialized intestinal metaplasia with goblet cells. It is a subject of controversy whether or not goblet cells are required as diagnostic criterion for BE. On the one hand, missing goblet cells in a biopsy specimen may represent a sampling error. On the other hand, there is evidence that esophageal cardiac epithelium, although lacking goblet cells, may also predispose to malignancy [17,18]. Two retrospective studies evaluated the risk of neoplasia in patients with columnar metaplasia of the esophagus either with or without goblet cells and found non-goblet cell columnar metaplasia to have the same malignant potential [19,20]. But the magnitude of this risk is unknown and so is the benefit of endoscopic surveillance. The British Society of Gastroenterology considers esophageal cardiac epithelium as a form of BE. The British guidelines point out that the distinction between columnar-lined esophagus and intestinal metaplasia at the gastric cardia can only be made definitively histologically when columnar mucosa is seen juxtaposed with native anatomical esophageal structures such as submucosal glands and/or gland ducts. But native structures are seen in only 10-15% of biopsy samples, which implies that in the great majority it is not possible to distinguish between intestinal metaplasia of the cardia and the esophagus. Biopsies of the normal cardia are not recommended routinely but if there is concern about the appearance at the site and after ablation therapy. The presence of intestinal metaplasia is considered as highly corroborative but not specific for a diagnosis of BE, as cardiac intestinal metaplasia cannot be ruled out. However, the guidelines recommend that this information should be recorded and that the diagnosis of BE should take into account the degree of confidence based on a combined analysis of endoscopic and histopathological criteria [21]. Other societies, including the AGA and the German Society of Gastroenterology, require esophageal biopsies showing intestinal metaplasia with goblet cells to establish the diagnosis [8,22]. After all, intestinal metaplasia is the only type of esophageal columnar epithelium that clearly predisposes to malignancy [8,22].

Diagnosis: can we drop the Seattle protocol with advanced endoscopic imaging?

To evaluate patients with BE high resolution endoscopy is recommended in order to detect subtle abnormalities of early neoplasia [23]. Endoscopic evidence of BE should be recorded using the Prague criteria [circumferential (C) and maximum (M)] extent of endoscopically visible columnar-lined esophagus in centimeters and any separate island above the main columnar-lined segment [24,25]). Current practice standards require the collection of targeted biopsies of every suspicious lesion followed by 4-quadrant biopsies specimens every 1 to 2 cm of BE (Seattle protocol). This approach is labor-intensive, so there has been a great deal of research in image-enhanced technologies.

Chromoendoscopy with contrast enhancing agents such as indigo carmine or acetic acid, virtual chromoendoscopy [Narrow band imaging (NBI, Olympus), Fuji Intelligent Chromo Endoscopy (FICE), and I-scan, Pentax] and confocal laser endomicroscopy; in addition to high-definition standard endoscopy, might increase the diagnostic yield for the detection of dysplastic lesions.

Acetic acid showed a sensitivity of 96% for the diagnosis of high-grade intraepithelial neoplasia or cancer and a 15-fold increase in neoplasia detection compared to the standardized random biopsy protocol [26,27]. NBI, which highlights surface patterns and vessels, was found to have a sensitivity and specificity of 96% and 94% for the diagnosis of HGD in a meta-analysis [28]. In a recent trial, NBI-targeted biopsies showed the same detection rate as high-definition white light examination with the Seattle protocol while requiring fewer biopsies [29]. The Barrett's international NBI Group (BING) developed and validated a NBI classification system to identify dysplasia and EAC in BE. Based on the simple classification of mucosal and vascular patterns as regular (non-dysplastic) and irregular (dysplastic) the BING Criteria could classify BE with >90% accuracy and a high level of inter-observer agreement [30].

Overall, advanced imaging techniques increased the diagnostic yield for detection of dysplasia or cancer by 34% in a recent meta-analysis [31]. In fact they may be very helpful to detect and delineate lesions but their diagnostic power is dependent on the expertise of the individual endoscopist. However, they have not been found to be superior to the standard 4-quadrant random biopsy protocol. Hence, current evidence seems insufficient to change practice. Careful examination using high-resolution endoscopes combined with targeted and 4-quadrant biopsies remains the gold standard [23,24].

Management of BE

Cancer in BE is thought to evolve through dysplasia. Dysplasia may be an imperfect marker to predict malignant progression as it can be patchy and therefore missed during routine biopsy sampling. Also, there may be significant interobserver disagreement about its grading [6]. However, dysplasia remains the basis for clinical decision making.
Endoscopic therapy aims to treat dysplastic precursor tissue to reduce cancer risk. In expert hands, endoscopic therapy of BE-related dysplasia and early neoplasia has shown to be effective and safe. In inexperienced hands, it may be associated with significant complications [32]. Therefore, endoscopic treatment should only be performed in centers with expertise [23]. Before treatment, a lesion should be assessed by an experienced endoscopist, using at least a high-resolution endoscope and one of the advanced endoscopic imaging modalities (NBI and/or chromoendoscopy) to determine whether the lesion is suitable for endoscopic treatment, to choose the appropriate resection technique (endoscopic mucosal resection (EMR) or endoscopic submucosal dissection (ESD)), delineate precise margins and to detect other possible lesions [33].

The management of neoplastic BE has changed considerably over the past decade. Today it consists of a multimodal approach combining tissue-acquiring and ablative techniques. Tissue acquiring techniques, which provide tissue specimens for histological examination, include EMR and ESD. Ablative techniques include radiofrequency ablation (RFA), argon plasma coagulation (APC), cryotherapy and photodynamic therapy (PDT). The common goal of ER, ablation or both is to completely eradicate all of the Barrett’s metaplasia, dysplastic and non-dysplastic. This concept has shown high rates of disease reversal. A brief algorithm for the endoscopic management of BE is provided in Fig. 1; details are discussed below.

**High-grade dysplasia (HGD) and early cancer**

In BE with HGD or intramucosal cancer endoscopic therapy is well established [34]. In contrast to the low risk of malignant progression in patients with non-dysplastic BE, the risk with HGD is considered high enough to warrant intervention. In a meta-analysis, the annual rate of cancer progression with HGD was calculated approximately 6% per year, but in endoscopic intervention studies the risk was found to be considerably higher [35,36]. Also, in HGD a risk of occult adenocarcinoma has been reported as high as 40% [37]. The risk to harbor carcinoma is particularly high in high-grade intraepithelial neoplasia areas that are endoscopically visible. ER of a visible lesion is essential for proper diagnosis and staging. EMR of visible lumps diagnosed with HGD on previous biopsy led in 25-40% of cases to a histological upgrading to cancer [38,39]. Hence, visible lesions should undergo ER. After ER all residual Barrett’s mucosa should be eradicated. This two-step concept can significantly reduce the risk of metachronous neoplasia. It remains the question what would be the best ablation technique. Well-studied alternatives are thermal ablation techniques as RFA and APC, PDT, and cryotherapy. Widespread EMR
can cause strictures, especially when more than two thirds of the circumference is removed. PDT plays no significant role any more due to its side effect of photosensitivity of the skin. Although there are no head to head comparative trials RFA seems to be superior and has become the preferred procedure for endoscopic ablation [40,41]. In a meta-analysis including more than 3800 patients RFA achieved complete eradication of dysplasia in 91% and complete eradication of intestinal metaplasia in 78% of patients. The most common adverse event was esophageal stricture, which was reported in 5% of patients [42].

Today the recommended standard of care in patients with HGD or intramucosal cancer is ER of visible lesions followed by RFA of residual Barrett’s mucosa [22-43]. This combined approach has shown high rates of disease reversal [44]. However, recurrences have been reported after successful endoscopic therapy. Hence, endoscopic follow up is mandatory. The ESGE recommends regular endoscopic follow up after excision/ablation of BE-associated HGD or mucosal cancer, but more research is necessary to determine the appropriate interval. Empirically, endoscopic follow up is recommended 3-monthly for 1 year and yearly thereafter [33].

If adenocarcinoma is found in the EMR specimen the risk of lymph node metastasis has been shown to correlate with the depth of invasion. In patients with mucosal neoplasm lymph node metastases are present in less than 2% but in patients with tumor infiltration into the deep submucosa in more than 20% [45]. In contrast to surgery, endoscopic therapy does not have the potential to cure neoplasm that has metastasized to regional lymph nodes. Therefore, ER is considered curative for intramucosal carcinomas that are well or moderately differentiated (G1-G2) without lymphatic or vascular invasion (L0, V0) [33]. Endoscopic en bloc R0 resection of a sm1 (<500 μm) low risk tumor (G1-2, L0 and V0, size <3 cm) is considered potentially curative [33]. Manner et al reported for these lesions a low risk of lymph node metastasis (1.4%), but only few patients were included in that study [46]. So the risk of lymph node metastasis should be balanced against the risk of surgery for the individual patient in a multidisciplinary discussion [33].

Surgery is recommended in the presence of [33]:
- Lymph vascular invasion (L1,V1)
- Deeper infiltration of the submucosa than sm1 (≥500 μm)
- Poorly differentiated tumor
- Positive vertical margins (R1 vertical)

If the horizontal margin is positive (R1 horizontal) or the tumor was resected piece meal and no other high-risk criteria are met, close endoscopic surveillance/treatment is recommended rather than surgery [33].

The standard for ER of Barrett’s neoplasia in current clinical practice is EMR. But in lesions >15 mm EMR entails piece meal resection, associated with higher recurrence rates and hampers histopathological assessment of free margins [47]. ESD allows the en bloc resection of a lesion regardless of its size (Fig.2). With en bloc resection the histopathological evaluation is improved and an adequate assessment of R0-status and curative resection as defined to oncological standards is possible. The efficiency and safety of ESD with high success rates have been demonstrated by Asian studies [48-50]. But in Asian countries BE and adenocarcinoma are still rare. Therefore, available data on ESD in BE are scarce. Nevertheless, it has been shown that also in western countries ESD of Barrett’s neoplasia is feasible with en bloc resection rates >95%, R0-resection rates >80%, and complication rates comparable to EMR [51-53].

In the presence of HGD or intramucosal cancer without visible lesions (flat HGD/intramucosal cancer) RFA is recommended [21-23,33]. Widespread EMR can cause strictures, especially when more than two thirds of the circumference is removed. RFA has been compared with stepwise ER for complete eradication of BE containing HGD/mucosal cancer. Both methods showed equivalent efficacy but radical ER was associated with higher stricture rates and therapeutic sessions [54]. If HGD or intramucosal cancer are confirmed and there are no visible lesions after expert high resolution endoscopy review, then ablative therapy is the treatment of choice [21]. But to reemphasize it: RFA should only be used as a primary treatment modality in the case of flat HGD/intramucosal cancer. All visible lesions should be resected to provide adequate histological assessment, even if this demands circumferential EMR or ESD. Resulting strictures do respond well to dilatation and there exist strategies for stricture prevention, such as steroid administration.

**LGD**

The management of LGD is confounded by uncertainty of its natural history and difficulties in making the diagnosis. The diagnosis of LGD in BE is a subject of high interobserver variability among pathologists and can be challenging in the presence of inflammation. As demonstrated in a recent Dutch
study, LGD in BE seems to be an over diagnosed and yet underestimated entity [6]. In this study 85% of patients who were initially diagnosed with LGD were down staged to either non-dysplastic or to indefinite for dysplasia (IND) after review by two expert GI pathologists. So it seems essential that the diagnosis is confirmed by at least two GI expert pathologists. The trial also showed that for patients with a consensus diagnosis of LGD, the cumulative risk of progression to HGD or carcinoma was alarming 85% in 109 months and the incidence rate for HGD or carcinoma 13.4% per patient per year. For down staged patients the corresponding incidence rate was 0.49%. Faced with this data gastroenterology societies propose that the diagnosis of dysplasia in BE should be confirmed by at least one additional pathologist, preferably one who is an expert in esophageal/gastrointestinal (GI) histopathology [21,22]. This recommendation takes in account the great medical importance of a “true” diagnosis of LGD but implicates challenges in its practical implementation (definition/qualification of an expert pathologist, independent evaluation, down-staging of diagnoses, financial aspects etc.).

The finding of an endoscopically visible lesion in the setting of biopsy-detected LGD is of special importance as it may contain HGD or invasive cancer. Hence, visible lesions in confirmed LGD should be resected endoscopically to enable accurate histological assessment [55]. ER may result in a change of histological diagnosis, as shown in a multicenter study, where ER in patients diagnosed with LGD on biopsy led to upstaging in 33.3% and downstaging in 13.3% [56]. If HGD or mucosal cancer is detected ER should be followed by ablation [55].

Ablation of BE with only LGD remains controversial because there is no clarity on cancer risk. As mentioned above, LGD generally seems to be overcalled but in those patients with LGD confirmed by at least two expert GI pathologists the risk of neoplastic progressions is considerably high. There are several studies that also indicate some clinical risk features such as multifocality of LGD and the length of BE-segment [57]. RFA can significantly reduce the risk of neoplastic progression to HGD/EAC. In the “SURF” trial it decreased the progression rate from 26.5% (control) to 1.5% (RFA) [58].

Non-dysplastic BE

Noting the success of RFA in eradicating Barrett's metaplasia some physicians have proposed that RFA should be offered to all BE patients rather than to restrict it to patients with dysplasia. But patients with no dysplastic BE have a very low risk to develop HGD or EAC. Recent studies show that the risk may be as low as 0.12-0.33% per year [3]. This low risk does not weigh out potential therapy associated risks and does not justify therapeutic intervention. Visible lesions in non-dysplastic BE (as well as visible lesions in BE with LGD or IND) should undergo ER to enable accurate histological assessment [55].

- We recommend that in the case of BE visible lesions in diagnosed LGD (or IND), ER should be followed by ablation if HGD or intramucosal cancer is detected, rather than continued surveillance [55]

Endoscopic surveillance

Endoscopic surveillance for BE is recommended by all Medical Societies. It is based on the assumption that the transition from BE to EAC progresses through LGD and HGD, thus justifying endoscopic surveillance for these premalignant stages. To date, the only evidence supporting this practice comes from observational studies reporting that patients whose Barrett’s carcinoma was diagnosed during surveillance endoscopy have earlier stage tumor and better survival. The recommendations vary amongst different Medical Societies and further data on optimal intervals and protocols for biopsy collection is needed.

Broadly speaking, in non-dysplastic BE surveillance endoscopy is recommended every 3-5 years. In the presence of LGD, confirmed by at least two expert GI pathologists, without visible lesions surveillance endoscopy every 6-12 months or eradication therapy is recommended [7].

Since recurrences of Barrett's metaplasia after apparently successful eradication are possible and recurrence rates up to 33% have been reported [59], patients should continue to undergo endoscopic surveillance even after therapy. Empirically, in patients treated for HGD, endoscopic follow up is recommended 3-monthly for 1 year and yearly thereafter.

Practical impact

- BE is a combined endoscopic and pathological diagnosis
- The Seattle protocol (4-quadrant biopsies every 1 to 2 cm of BE and of every suspicious lesion) remains the standard; advanced imaging techniques may increase the diagnostic yield
- For any degree of dysplasia, at least two expert GI pathologists are required to confirm the diagnosis
- Visible lesions should be endoscopically resected to enable accurate histological assessment
- In HGD/mucosal cancer ER of visible lesions followed by field ablation of the whole Barrett's segment with RFA is now the standard of care
- In LGD (confirmed by at least two expert GI pathologists) with visible lesions ER should be performed. Without visible lesions surveillance endoscopy every 6-12 months or eradication therapy is recommended
- In non-dysplastic BE the risk of progression is low. Surveillance endoscopies are recommended every 3-5 years
- Recurrences after apparently successful eradication of
Barrett's metaplasia are possible. Further endoscopic surveillance is indispensable.

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