



# Barrett's esophagus: current standards in advanced imaging

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*Contributions:* (I) Conception and design: All authors; (II) Administrative support: None; (III) Provision of study materials or patients: None; (IV) Collection and assembly of data: None; (V) Data analysis and interpretation: None; (VI) Manuscript writing: All authors; (VII) Final approval of manuscript: All authors.

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**Abstract:** Esophageal adenocarcinoma (EAC) continues to be one of the fastest rising incident cancers in the Western population with the majority of patients presenting with late stage disease and associated with a dismal 5-year survival rate. Barrett's esophagus (BE) is the only identifiable precursor lesion to EAC. Strategies to screen for and survey BE are critical to detect earlier cancers and reduce morbidity and mortality related to EAC. A high-quality endoscopic examination with careful inspection of the Barrett's segment and adherence to the Seattle protocol for tissue sampling are critical. Advanced imaging modalities offer the potential to improve dysplasia detection, predict histopathology in real time and guide endoscopic eradication therapy (EET). Several technologies have been studied and although most are not yet recommended for routine clinical practice, high definition white light endoscopy (HD-WLE) as well as chromoendoscopy (including virtual chromoendoscopy) improved dysplasia detection in numerous studies supporting their use. Future studies should evaluate the role of artificial intelligence in optimizing detection of dysplasia in BE patients.

**Keywords:** Esophageal adenocarcinoma (EAC); Barrett's esophagus (BE); advanced imaging; chromoendoscopy

Received: 21 August 2019. Accepted: 21 January 2020; Published: 05 January 2021.

doi: 10.21037/tgh.2020.02.10

**View this article at:** <http://dx.doi.org/10.21037/tgh.2020.02.10>

## Introduction

Esophageal adenocarcinoma (EAC) is a growing threat worldwide with a global incidence of 0.7 per 100,000 person years (1.1 in men, 0.3 in women) (1). EAC has become the predominant subtype in many Western Countries where it is now more common than esophageal squamous cell carcinoma (2). In the US, the incidence of EAC is on the rise with an estimated 18,440 new cases and 16,170 EAC deaths projected in 2020 (3-5). EAC typically affects older adults with identifiable risk factors including male gender, Caucasian, gastroesophageal reflux disease, smoking, and central obesity. The prognosis for patients with EAC is strongly related to stage at diagnosis, but unfortunately the majority of patients with EAC present with late stage disease and have limited options. Despite implementation of screening programs and greater recognition of this

deadly cancer, we have unfortunately still not impacted outcomes related to this cancer on a population level (4). Advancements in endoscopic and minimally invasive surgical interventions have led to improved outcomes for those patients who present with earlier stage disease; thus, prompt identification of patients with risk factors for BE/EAC is critical.

### *Rationale for Barrett's esophagus (BE) screening and surveillance endoscopy*

BE is the only identifiable precursor lesion of EAC and affects up to 5% of the general population (6). It results from chronic acid exposure of the stratified squamous epithelium in the distal esophagus leading to metaplasia and replacement with intestinal type columnar epithelium. BE is believed to progress from non-dysplastic (NDBE) → low

grade dysplasia (LGD) → high grade dysplasia (HGD) → intramucosal carcinoma (IMC) and finally to invasive EAC in a stepwise fashion. Robust screening programs for those at high risk, coupled with guideline supported surveillance strategies for BE patients, offer an opportunity to intervene on this process and change the patient's trajectory. At the present time, there is no level 1 evidence that screening and surveillance reduces incidence, morbidity, or mortality related to BE/EAC. However, there are important case control and cohort studies that address this question. A recent systematic review and meta-analysis demonstrated lower EAC-related mortality associated with regular BE surveillance (RR 0.60, 95% CI, 0.50–0.71) as well as with surveillance detected EAC *vs.* symptom detected EAC (RR 0.73, 95% CI, 0.57–0.94) (7). Barrett's esophagus surveillance allows for detection of EAC at earlier stages, which is associated with better outcomes. Randomized trials are needed to assess the impact of surveillance and potential to decrease the burden and mortality of EAC.

A high-quality endoscopic exam, detection of neoplastic lesions and endoscopic eradication therapy (EET) or minimally invasive surgical procedures for dysplastic lesions and early cancers are critical steps for effective surveillance programs. Several advanced imaging modalities have been investigated with numerous studies on chromoendoscopy (dye-based and virtual), confocal laser endomicroscopy (CLE), volumetric laser endoscopy (VLE), and more recently artificial intelligence. Adjunct use of these technologies during endoscopy allows for real-time diagnosis and prediction of histology which can effectively guide EET (8).

The objectives of this review include to (I) review best practices in Barrett's Esophagus surveillance; (II) examine the role of advanced imaging techniques; (III) highlight guidelines and quality indicators.

### ***Principles for a high-quality endoscopic examination of BE***

The first step to a high-quality examination requires documentation of esophageal landmarks including the presence of a hiatal hernia (9). Use of a distal attachment cap can facilitate improved visualization. The mucosa should be carefully cleaned with the water jet and gently suctioned to remove any debris or mucus without damaging the mucosa. Precise manipulation of insufflation and desufflation can adjust the luminal view to detect subtle abnormalities. The endoscopist should spend enough time (consider 1 minute per cm) carefully inspecting the Barrett's

segment (10). Findings should be reported in a systematic fashion. The Barrett's segment should be carefully measured and reported using the Prague classification (Circumferential and Maximal extent of Barrett's) (11). Any superficial lesions should be described using the Paris classification (12,13) (*Figure 1*). A ten-step approach for a high-quality examination of BE is provided (*Table 1*).

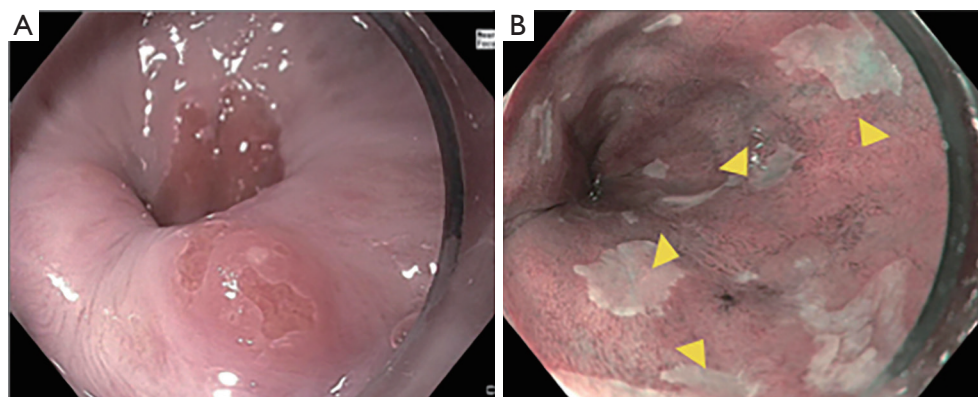
### ***Sampling BE***

The Seattle protocol (four-quadrant biopsies using the "turn and suction technique" at 1–2 cm intervals along the entire length of the Barrett's segment) remains the standard of care for tissue sampling (14). Any suspicious areas (nodularity, erythema, erosion, ulceration) should be biopsied and samples placed in separate jars as these mucosal abnormalities can be associated with dysplasia (15). All visible lesions, no matter how subtle, should be completely removed endoscopically (16).

The major limitation in current BE surveillance practice is the need for random biopsies. Despite international guideline recommendations to follow a systematic approach (the Seattle protocol), non-adherence is common, and increases by 31% for every 1cm increase in length of the Barrett's segment (17). The rate of guideline adherence in the community setting is only about 50%, which significantly decreases dysplasia detection (OR 0.53, 95% CI, 0.35–0.82) (18). This may contribute to high miss rates for EAC. Indeed, a systematic review and meta-analysis of 820 patients (24 studies) with index endoscopy with NDBE or BE-LGD demonstrated up to 25% (95% CI, 16.4–36.8%) of EAC were missed (19). Most neoplasia in BE is nonpolypoid making it difficult to detect, and random biopsy is prone to sampling error as the distribution of neoplasia is highly variable and focal so can be easily missed (20). These limitations inherent in use of white light endoscopy (WLE) with random biopsies may explain why earlier studies underestimated the mortality benefit of endoscopic surveillance (21).

### ***Imaging modalities in Barrett's esophagus***

Imaging modalities have the potential to overcome these challenges. They can help identify suspicious areas for additional biopsies on top of the routine, guideline supported approach. Use of additional technologies does not replace the Seattle protocol. We emphasize their value to facilitate high quality examinations with closer inspection



**Figure 1** Description of visible lesions in Barrett's Esophagus using the Paris Classification. (A) Nodule (Paris 0-IIa) in an area of previous ablation; (B) nodular area (Paris 0-IIa) with abnormal mucosal and vascular pattern under NBI. NBI, narrow band imaging.

**Table 1** Ten step approach to high quality endoscopic examination of Barrett's esophagus

Approach	Rationale
1. Consider use of a distal attachment cap	1. Facilitate visualization
2. Utilize CO <sub>2</sub> insufflation and desufflation	2. Fine adjustments to luminal insufflation can help with detection of subtle abnormalities
3. Clean mucosa well using water jet channel and carefully suction the fluid	3. Remove any distracting mucus or debris and minimize mucosal trauma
4. Identify esophageal landmarks, including the location of the diaphragmatic hiatus, gastroesophageal junction, and squamocolumnar junction	4. Critical for future exams
5. Examine the Barrett's segment using high definition white light endoscopy	5. Standard of care
6. Examine the Barrett's segment using chromoendoscopy (including virtual chromoendoscopy)	6. Enhances mucosa pattern and surface vasculature
7. Spend adequate time inspecting (consider 1 minute per cm)	7. Careful examination increases dysplasia detection
8. Use the Prague classification to describe the circumferential and maximal Barrett's segment length	8. Standardized reporting system
9. Use the Paris Classification to describe superficial neoplasia	9. Standardized reporting system
10. Use the Seattle Protocol (in conjunction with advanced imaging modalities) to sample the Barrett's segment	10. Increases dysplasia detection

and increased identification of suspicious areas, a strategy of "Look more, biopsy appropriately." Furthermore, advanced imaging offers the chance to detect and diagnose dysplasia in real-time and guide EET (*Figure 2*).

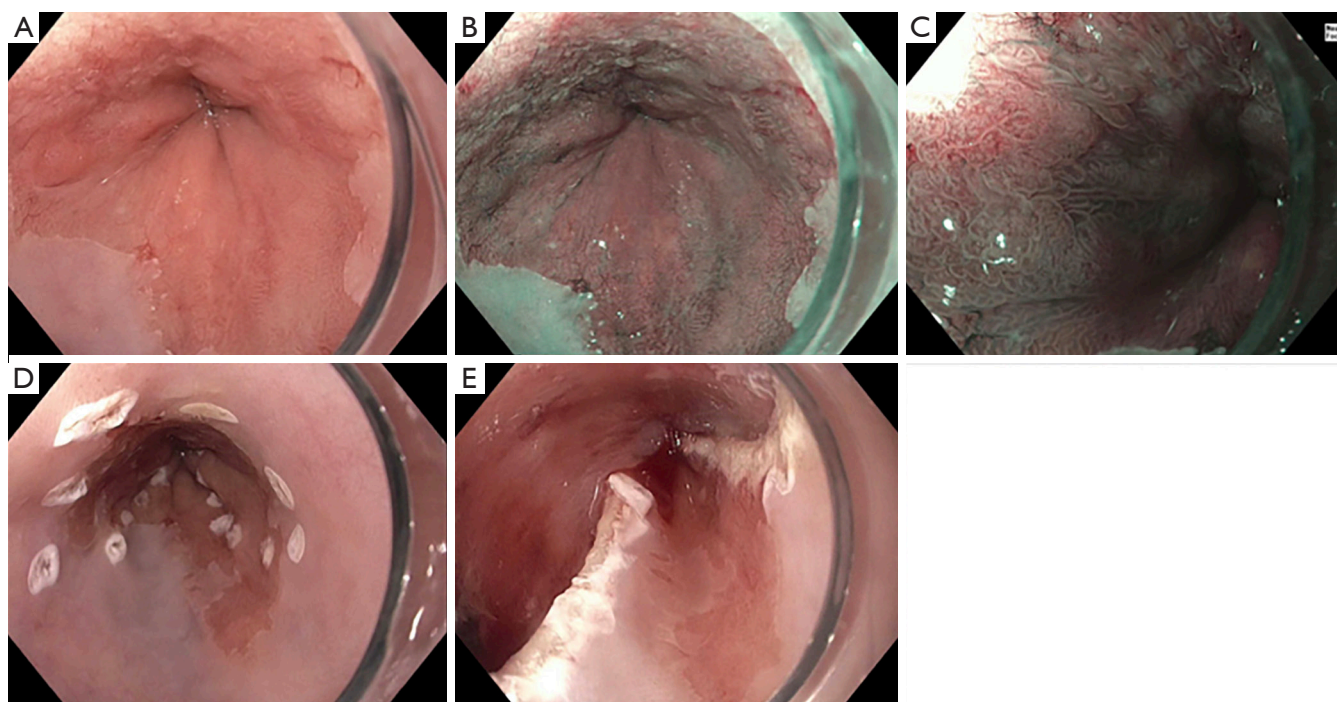
### **High definition white light endoscopy (HD-WLE)**

At a minimum, HD-WLE should be used. Cohort studies have demonstrated clear superiority of HD-WLE compared to standard WLE for dysplasia detection (22). Although there

are no head to head randomized controlled trials, HD-WLE is now considered standard of care and endorsed by societal guidelines and consensus recommendations (23-25).

### **Conventional and virtual chromoendoscopy**

Chromoendoscopy, either dye-based or virtual, is the most extensively studied advanced imaging modality for improving neoplasia detection in BE. A comprehensive review of 14 studies (843 patients) demonstrated a 34%



**Figure 2** Use of advanced imaging to guide endoscopic eradication therapy. (A) high definition white light endoscopy of Barrett's Segment; (B) subtle area of nodularity (Paris 0-II) extending from the 9 to 3 o'clock position; (C) closer inspection of mucosal and vascular pattern using NBI and near focus; (D) marking prior to endoscopic mucosal resection; (E) status post endoscopic mucosal resection.

(95% CI, 20–56%) increase in the diagnostic yield of dysplasia or cancer using chromoendoscopy compared to WLE (26). Importantly, there was no difference between dye based or virtual chromoendoscopy ( $P=0.45$ ).

### Dye-based chromoendoscopy

The most commonly used dyes for BE surveillance are acetic acid, methylene blue, and indigo carmine. Application of these agents to the surface mucosa within the esophagus can enhance surface vasculature and the appearance of neoplasia. They are applied via a spray catheter through the working channel of the endoscope to lightly coat the surface mucosa in an even fashion. General limitations include cost, need for special dye spraying equipment, additional procedure time, and difficulty in adequate application. Acetic acid is a colorless chemical that effectively stains BE a whitish color. It works by disrupting cells in the surface epithelium leading to changes in the mucosal patterns. A clue to dysplasia is an area that loses the whitening effect earlier than the surrounding mucosa. In the hands of expert endoscopists, acetic acid chromoendoscopy performs well and meets the ASGE Preservation and Incorporation of

Valuable Innovations (PIVI) thresholds (sensitivity 96.6%, negative predictive value 98.3%, specificity 84.6%) (27). Methylene blue is a vital dye that is quickly absorbed by intestinal and colonic epithelium with effects lasting up to 20 minutes. In BE, areas of intestinal metaplasia appear dark blue (stained) compared to unstained esophageal squamous epithelium. Areas with irregular uptake of blue stain (dark and light areas) are suspicious for neoplasia. A meta-analysis of 9 studies (450 patients) showed no increased benefit with methylene blue chromoendoscopy compared to WLE with random biopsy for BE (28). The performance characteristics also do not meet PIVI thresholds (sensitivity 64%, negative predictive value 70%, specificity 96%) (27). In light of inadequate metrics and a potential risk of causing DNA damage, methylene blue chromoendoscopy is not currently recommended for use (29). Indigo carmine is a non-vital dye that collects in the pits and grooves of the surface mucosa. It has been studied mostly in conjunction with high resolution magnification endoscopy.

### Virtual chromoendoscopy

Virtual chromoendoscopy uses technology built into the

endoscope to achieve a similar result as dye application with the ease of only pressing a button. The most widely used system is narrow band imaging (NBI, Olympus) which applies a red-green-blue light filter to the target mucosa. NBI is based on the optical phenomenon that the depth of light penetration into tissue depends on the wavelength; the shorter the wavelength, the more superficial the penetration. Whereas standard WLE uses light at wavelengths of 400–700 nm, NBI applies shorter wavelengths (400–540 nm) to maximally highlight the surface mucosa and vascular pattern (30). Additionally, this narrower spectrum is matched to the maximum absorption of hemoglobin, so that structures with high hemoglobin content will appear dark (surface capillaries appear brown, submucosal vessels appear cyan) compared to the brighter surrounding mucosa. Other platforms for virtual chromoendoscopy are The Fujinon Intelligent Color Enhancement (FICE, Fujinon Inc., Japan) and iScan (Pentax, Japan) systems which capture the full spectrum of white light then apply digital processing programs to enhance the surface mucosa and vascular patterns (31). The newer iScan Optical Enhancement system (OE) uses specific wavelengths of light matched to hemoglobin absorption to highlight the microvasculature in combination with magnification endoscopy for closer inspection, and offers potential to improve dysplasia detection in BE (32). Another more recent system is ELUXEO 7000 (Fujifilm, Japan) which uses 4 LED multi light technology that facilitates visualization using blue light imaging (BLI) and can aid in visualization of BE neoplasia (33).

Virtual chromoendoscopy adds no cost (already built into the endoscope), negligible time (pressing a button), and no additional risk to the patient. Results from an international crossover randomized controlled trial showed higher detection of dysplasia (30% *vs.* 21%,  $P=0.01$ ) and fewer biopsies required per patient (3.6 *vs.* 7.6,  $P<0.0001$ ) with NBI compared to HD-WLE (34). The ASGE recently updated previous meta-analyses and provided strong data supporting the use of chromoendoscopy in BE. They noted a 9% absolute increase in dysplasia detection (95% CI, 4.1–14%) and 30.3% relative increase (95% CI, 16.2–44.3%), with no difference between virtual or dye based chromoendoscopy (35).

### Additional advanced imaging modalities

Several other imaging modalities have been investigated for use in BE but most are not ready for clinical application at this time (14).

### CLE

CLE is a technology based on principles of light microscopy that process reflected light into a high-resolution gray scale image for mucosal visualization at the cellular level (36). Endoscopy based (eCLE) is no longer commercially available but probe based (pCLE) can be used. It has the potential for *in vivo* histology, which could facilitate same-session endoscopic therapy and resection. There is also potential for machine learning in endomicroscopy (37). However, it is time consuming, costly, requires IV contrast agents, and evaluation is limited to the mucosa. A recent systematic review and meta-analysis by the ASGE noted a marginal increase in diagnostic yield (absolute increase in dysplasia detection 10.2%, 95% CI, 1.4–19.1%) and a non-significant relative increase in dysplasia detection (36%, 95% CI, –5.4–77.5%), informing their recommendation against routine use of CLE (conditional recommendation, low quality of evidence (35)).

### VLE

Optical coherence tomography (OCT) is a probe-based platform that investigates areas of interest with infrared light and creates high resolution cross-sectional images. VLE employs a technology similar to OCT where the probe is passed through a balloon designed for use in the esophagus and images the esophageal wall layers. VLE scans 6 cm of the esophagus over 90 seconds and provides a resolution of 10 mm and an imaging depth of 3 mm (35). Continuous adaptations to the platform make it difficult to study outcomes, but certain new features such as a laser to tag areas of suspected dysplasia and target them for biopsies, offer great potential to strengthen this technology for BE (38). VLE is time consuming and learning how to interpret images is complex, but computer-aided detection (CAD) may enhance application of this technology. European studies demonstrated strong ability to detect BE neoplasia using a computer-assisted algorithm for VLE (Area under the receiver operating characteristic curve AUC 0.95), and greater AUC with multi-frame analysis (39,40). As the technology continues to evolve, additional studies on diagnostic accuracy and cost effectiveness will be important before it can be recommended (41).

### Other imaging modalities

Autofluorescence imaging (AFI) is a technology that depends

on endogenous fluorophores within the GI mucosa that absorb light of varying wavelengths depending on their metabolic activity, blood flow, and biochemical characteristics. This provides an opportunity to distinguish between normal tissue and tissue which is inflamed or neoplastic. AFI is limited by a high false positive rate and minimal incremental diagnostic yield over Seattle protocol (42). Molecular endoscopy uses targeted probes directed to specific molecules in the GI tract and has the potential for highly specific *in vivo* diagnosis (43). It remains limited by cost, time, and special equipment that is not available for routine clinical use.

### **Artificial intelligence**

CAD systems have arrived and offer promising technology that could transform our ability to endoscopically detect BE related neoplasia (44,45). The ARGOS project was developed in collaboration by expert endoscopists at BE referral centers in the Netherlands, an imaging analysis group, and commercial enterprises with the goal to create the first CAD for BE neoplasia detection. Their initial work focused on training their algorithm using endoscopic still images in WLE and results showed high accuracy (92%), sensitivity (95%), and specificity (85%) for detecting and localizing dysplasia in BE (46). They used over 494,364 still images to train their system, further trained and refined their system on datasets of retrospectively and then prospectively collected images of BE neoplasia (with internal validation), and then a fourth dataset was used for external validation (47). Their CAD system currently shows high accuracy for detecting BE neoplasia and next steps include deeper learning with video recordings to refine an algorithm for real-time analysis during endoscopy. A US group has also developed an artificial intelligence algorithm using convolutional neural networks that detected BE neoplasia with accuracy 93.7%, sensitivity 95.6%, specificity 91.8% for an AUC 0.94 (48). Artificial intelligence is also being developed for VLE and BE (49).

### **Incorporating imaging modalities into practice: guidelines & quality indicators**

Gastroenterology societies worldwide provide guidelines on use of imaging modalities (Table 2) as well as quality indicators in BE (Table 3).

The ASGE set minimum PIVI thresholds for when advanced imaging modalities should be recommended in clinical practice. To eliminate random biopsies in BE

surveillance, an imaging technology with targeted biopsies should meet the following performance threshold: (I) per-patient sensitivity of 90% or greater and a negative predictive value (NPV) of 98% or greater for detecting HGD/early EAC, compared with the current standard protocol (WLE with targeted and random 4-quadrant biopsies every 2 cm); (II) specificity should be sufficiently high (80%) to allow a reduction in the number of biopsies (compared with random biopsies) (53). Currently, only acetic acid chromoendoscopy, virtual chromoendoscopy with NBI, and endoscope-based confocal laser endomicroscopy meet these thresholds, but only when performed by expert endoscopists (27). Additionally, an international survey of BE experts sought to determine the minimum incremental diagnostic yield for detection of dysplasia or cancer of an advanced imaging modality compared to the standard Seattle Protocol. Results showed the incremental diagnostic yield for virtual chromoendoscopy to be fairly low (15%, IQR 10–29%) whereas for VLE it was much higher (30%, IQR 20–50%). These results provide benchmarks that need to be met for guidelines to recommend routine use of these technologies (54).

The need for standardized grading systems has also limited uptake of advanced imaging modalities. Several classification systems have been proposed using NBI, but the criteria were complex and validation studies were poor. To address this gap, The Barrett's International NBI Group (BING) brought together experts from the US, Europe, and Japan to develop a simple, validated system for identification of dysplasia and EAC in patients with BE (55). The BING criteria distinguishes regular mucosal patterns (circular, ridged/villous, or tubular patterns) from irregular mucosal patterns (absent or irregular surface patterns), and regular vascular patterns (regularly positioned blood vessels along or between mucosal ridges and/or characterized by normal, long branching patterns) from irregular vascular patterns (focally or diffusely distributed vessels in discordance with the normal architecture). The BING criteria demonstrated 85% overall accuracy for identification of dysplasia (sensitivity 80%, specificity 88%, 81% positive predictive value, 88% negative predictive value) and high interobserver agreement ( $k=0.681$ ) (55).

### **Training in advanced imaging modalities**

Significant advances in science and technology have tremendously improved our ability to recognize early neoplasia in the gastrointestinal tract. Image enhanced endoscopy is clearly a valuable and efficacious tool to increase detection of dysplasia in BE as demonstrated by

**Table 2** Guideline Recommendations for imaging in BE

Society	Recommendation for Imaging in BE
ASGE 2019 (35)	<p>In patients with BE undergoing surveillance, we recommend using chromoendoscopy, including virtual chromoendoscopy and Seattle protocol biopsy sampling, compared with white-light endoscopy with Seattle protocol biopsy sampling (strong recommendation, moderate level of evidence)</p> <p>In patients with BE undergoing surveillance, we suggest against routine use of confocal laser endomicroscopy compared with white-light endoscopy with Seattle protocol biopsy sampling (conditional recommendation, low quality of evidence)</p>
ESGE 2017 (50)	High definition endoscopy (endoscope, processor, and screen) is recommended for endoscopic surveillance of BE. Routine use of chromoendoscopy, optical chromoendoscopy, autofluorescence endoscopy, or confocal laser endomicroscopy is not advised
ACG 2016 (51)	<p>Surveillance should be performed with high definition/high-resolution white light endoscopy (strong recommendation, low level of evidence).</p> <p>Routine use of advanced imaging techniques other than electronic chromoendoscopy is not recommended for endoscopic surveillance at this time (conditional recommendation, very low level of evidence).</p>
AGA 2011 (25)	<p>We recommend endoscopic evaluation be performed using white light endoscopy (strong recommendation, moderate-quality evidence)</p> <p>We suggest against requiring chromoendoscopy or advanced imaging techniques for the routine surveillance of patients with Barrett's esophagus at this time (weak recommendation, low-quality evidence). (These technologies may be helpful in guiding the performance of biopsies in patients who are known to have dysplasia and in patients who have mucosal irregularities detected by white light endoscopy)</p>
BSG 2014 (24)	<p>High-resolution endoscopy should be used in Barrett's oesophagus surveillance (grade C recommendation)</p> <p>Advanced imaging modalities, such as chromoendoscopy or 'virtual chromoendoscopy', are not superior to standard white light endoscopy in Barrett's oesophagus surveillance and are therefore not recommended for routine use (grade A recommendation)</p> <p>Expert high-resolution endoscopy (HRE) should be carried out in all Barrett's patients with biopsy detected HGD in order to detect visible abnormalities suitable for ER (grade B recommendation)</p>

BE, Barrett's esophagus; ASGE, American Society for Gastrointestinal Endoscopy; ESGE, European Society of Gastrointestinal Endoscopy; ACG, American College of Gastroenterology; AGA, American Gastroenterological Association; BSG, British Society of Gastroenterology; ER, endoscopic resection.

strong clinical trial data. So why is it that uptake remains limited and competency so difficult to achieve? Certainly, resources may be limited in particular practice settings. Time and cost may be a factor. The major driver is likely the lack of training available in these advanced imaging modalities. Technology has evolved quicker than our ability as a society and community to create robust, high-quality educational programs for endoscopists of all levels. To overcome challenges with advanced imaging in BE we propose two major goals: (I) to train non-expert endoscopists to achieve PIVI thresholds; (II) to refine our techniques to exceed the PIVI thresholds at the expert level. The second goal is somewhat longer term as it may rely on technological developments and clinical trial data.

The first task can be addressed in the short term and

requires training the larger community of gastroenterologists around the world. For these technologies to be incorporated into routine clinical practice, providers need to feel comfortable and competent in recognizing BE related neoplasia (54). This may require leveraging societal efforts and linking performance to quality metrics. The need for validated educational programs is now being recognized. A group in the UK developed and validated a training module that was effective at improving endoscopist performance in use of acetic acid chromoendoscopy for Barrett's detection (56). More recently, an international working group developed an interactive, web based tool for detection and delineation of Barrett's esophagus-related neoplasia (BORN) (57). The BORN training module is freely available on the internet

**Table 3** Quality Indicators for Imaging BE

Society	Quality Indicators for Imaging BE
ASGE/ACG 2017 (16)	<p>The rate at which landmarks and length of BE is documented (e.g. Prague Grading System) in patients with BE before EET (Threshold 90%, 75–100).</p> <p>The rate at which the presence or absence of visible lesions is reported in patients with BE referred for EET (Threshold 90%, 60-100).</p> <p>The rate at which the BE segment is inspected by using HD-WLE (Threshold 95%, 0-100).</p> <p>The rate at which biopsies of any visible mucosal abnormalities are performed during endoscopic surveillance after EET (Threshold 95%, 50-100).</p>
ESGE 2017(50)	<p>Endoscopy reports of patients with BE should include:</p> <p>The extent of BE using the Prague criteria [circumferential extent (C), maximum extent (M)], and any separate islands proximal to the maximal extent;</p> <p>A description of location (in cm from the incisors and clockwise orientation) of any visible abnormality within the Barrett's epithelium, in addition to lesion size (mm) and macroscopic appearance using the Paris classification;</p> <p>The presence or absence of erosive esophagitis using the Los Angeles classification;</p> <p>The location of biopsies taken from the Barrett's segment (number of biopsies and location in cm from the incisors);</p> <p>Appropriate photo documentation of the landmarks and of all visible Barrett's epithelium, as well as any visible lesions</p>
AGA 2015 (52)	<p>If BE is suspected on an endoscopic screening examination, the endoscopist should obtain multiple systematic biopsies from the suspected segment (consensus agreement 92%, low quality evidence).</p>

BE, Barrett's esophagus; EET, endoscopic eradication therapy; ASGE, American Society for Gastrointestinal Endoscopy; ESGE, European Society of Gastrointestinal Endoscopy; ACG, American College of Gastroenterology; AGA, American Gastroenterological Association.

to endoscopists of all levels and is Continuing Medical Education-accredited. The program delivers high quality endoscopy videos through an interactive platform that provides scoring and feedback to participants.

Finally, the future of endoscopy lies in our current trainees. The traditional structure of gastroenterology (GI) fellowship has been an apprenticeship-based model much like the rest of medicine, where trainees learn clinical medicine and endoscopy on the job. With implementation of the Next Accreditation System through the ACGME, there has been a shift towards Competency Based Medical Education (CBME), an approach that recognizes trainee progression through a series of clinical milestones. This system relies on defined quality and competency metrics in the cognitive and technical aspects of endoscopy, many of which have been established in the GI core curriculum. As technology evolves and the endoscopic armamentarium grows, it will be critical to incorporate training in advanced imaging modalities into fellowship programs (58). At the present time, there is no data on trainee learning curves or the impact of training on achieving competency in advanced imaging modalities. Training the next generation can be achieved through use of online modules or cloud-based

curriculum, flipped classroom learning, and a case-based approach (59).

## Conclusions

Improving our ability to detect and diagnose early BE related neoplasia can guide EET and may help decrease the burden of EAC. This starts with a high-quality endoscopic examination and careful inspection of the Barrett's segment. HD-WLE is considered minimum standard of care. Additionally, we recommend use of chromoendoscopy (including virtual chromoendoscopy) and the Seattle protocol for biopsy sampling. Practicing endoscopists and trainees should follow guideline recommendations and understand key quality indicators for the management of BE related neoplasia. With evolving technology and integration of artificial intelligence into endoscopy, it will be critical to design educational programs to graduate competent fellows and train practicing endoscopists.

## Acknowledgments

*Funding:* None.



## Footnote

*Provenance and Peer Review:* This article was commissioned by the Guest Editors (Krish Ragnath, Philip WY Chiu) for the series “Advanced Endoscopic Imaging of the GI Tract” published in *Translational Gastroenterology and Hepatology*. The article was sent for external peer review organized by the Guest Editors and the editorial office.

*Conflicts of Interest:* Both authors have completed the ICMJE uniform disclosure form (available at <http://dx.doi.org/10.21037/tgh.2020.02.10>). The series “Advanced Endoscopic Imaging of the GI Tract” was commissioned by the editorial office without any funding or sponsorship. JM Kolb received funding from the National Institutes of Health (NIH) T32-DK007038. S Wani is supported in part by the Department of Medicine Outstanding Early Scholars Program, and is a consultant for Medtronic, Boston Scientific, and Interspace.

*Ethical Statement:* The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved.

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doi: 10.21037/tgh.2020.02.10

**Cite this article as:** Kolb JM, Wani S. Barrett's esophagus: current standards in advanced imaging. *Transl Gastroenterol Hepatol* 2021;6:14.