Recent advances in endoscopic imaging of the esophagus have revolutionized the diagnostic capability for detecting premalignant changes and early esophageal malignancy. In this article, we review the practical application of narrow-band imaging focusing on diseases of the esophagus, including Barrett’s esophagus, adenocarcinoma, and squamous cell carcinoma. (Gut Liver 2021;15:492-499)

Key Words: Barrett’s esophagus; Esophageal adenocarcinoma; Esophageal squamous cell carcinoma; Narrow band imaging; Mucosal imaging

INTRODUCTION

Endoscopic imaging has evolved over time from white light endoscopy to the adoption of novel electronic imaging techniques utilizing the technology of altering white-light with a push of a button. With the addition of magnification technology, the endoscopist now has the capability to assess mucosal surface architecture in greater detail.1 Narrow-band imaging (NBI) is an imaging technique that utilizes specific wavelength of blue and green light to penetrate into the superficial layers of the mucosa to highlight abnormal mucosal and vascular pattern.2 A white-light source filter located in front of a xenon arc lamp in the endoscope produces two selective narrow bands of wavelength lights measuring 415 nm and 540 nm. The length of the wavelength is directly proportional to the depth of its penetration. The 415 nm wavelength highlights only the superficial mucosa where the capillaries appear brown whereas the 540 nm wavelength penetrates deeper into the lower parts of the mucosa and submucosa giving them a blue-green hue.2,3 The end result is the ability to better visualize the microsurface and microvascular patterns on the mucosal surface.4 This allows advanced evaluation of abnormal lesions such as dysplasia and cancer.5

BARRETT’S ESOPHAGUS AND ESOPHAGEAL ADENOCARCINOMA

Barrett’s esophagus (BE) carries the risk of progression to esophageal adenocarcinoma (EAC). BE remains the sole proven premalignant condition for EAC.6 Treatment options are limited and prognosis remains bleak when EAC is diagnosed at a late stage. In order to achieve early diagnosis and treatment, national and international gastroenterological societies have endeavored to draft guidelines for BE surveillance.7 In general, the prevalence of BE has been estimated to be 1%–2% in patients receiving endoscopy for any indication and increases to 5%–15% among patients with symptoms of gastroesophageal reflux disease.8 Risk factors for developing BE include chronic gastroesophageal reflux disease,9,10 obesity (central adiposity),11,12 cigarette smoking,13 male sex,14 older age15 and a positive family history of BE or EAC.16

To diagnose BE, two components need to be present: first, the endoscopic appearance of a salmon-pink colored columnar epithelium extending above the gastro-esophageal junction, replacing the normal tubular esophageal squamous epithelium and second, histological demonstration of esophageal columnar epithelium with intestinal metaplasia showing presence of mucin-containing goblet...
To diagnose dysplasia, current guidelines recommend endoscopic surveillance by performing random four quadrant biopsies every 1 to 2 cm of the BE mucosa. However random sampling can lead to random errors where an area harboring dysplasia may be missed. Furthermore, once worrisome histology is found on random biopsies, it can be technically challenging to re-locate the lesion. Endoscopists adherence to biopsy guidelines is also low likely due to its cumbersome nature. Studies have shown that adherence to biopsy protocols worsen with increasing BE length and this becomes dangerous as the risk of EAC is greater with longer BE segments.

The rationale behind the advent of NBI is to complement and improve observation methods and possibly reduce the need for random histological sampling as suspicious areas may be identified and interrogated further. There have been various attempts to describe pit patterns seen in BE. A formal consensus is required to limit the varying classifications put out by different groups of experts as it only confuses gastroenterologists adopting the utility of NBI.

**NBI OF BARRETT’S ESOPHAGUS**

It has been shown that targeted biopsies with NBI assistance can achieve the equivalent of random biopsies using high definition white light endoscopy (WLE) in the detection of intestinal metaplasia with lesser number of biopsies and better diagnostic yield. The additional employment of high magnification settings significantly increased the accuracy of cancer detection compared to NBI alone. This combination allows the operator the capability to zoom in and focus on an area of interest, thus allowing confirmation of findings when evaluating dysplastic areas. The rapid evolution of NBI with magnification endoscopy assessment of BE is as a result of its excellent correlation with histological outcome. This obviates the need for physical biopsy when optical biopsy with NBI and magnification can confidently confirm non-dysplastic areas.

Sharma et al. from Kansas and Kara et al. from Amsterdam were amongst the first to describe and characterize certain mucosal and vascular patterns observed on NBI magnification images that correlated with non-dysplastic and dysplastic Barrett’s mucosa. The basis of these classifications work on the observations that non-dysplastic Barrett’s mucosa has regular mucosal and vascular pattern while advance dysplastic lesions has irregular, distorted or absent mucosal and vascular pattern.

Sharma’s group was able to identify that irregular and distorted mucosal pattern with abnormal branching, tortuous or non-uniform vascular pattern had 100% sensitivity and 95% positive predictive value of identifying high grade dysplasia. Of note, NBI magnification imaging was not reliable in detecting low grade dysplasia. Kara et al. were first to compare NBI’s practical superiority over chromoendoscopy. They further proposed a classification system based on NBI morphological appearance of the mucosal and vascular pattern. The emphasis was again on regularity of different mucosal patterns such as villous or gyrus pattern, regularity of vascular pattern and absence of any abnormal appearing vessels which was most consistent with non-dysplastic tissue. Presence of any one criteria such as irregularity in mucosal pattern, vascular pattern or abnormal vessel would significantly increase the probability of higher grade dysplasia; up to 85% probability if two features were present. Absence of any of these criteria had high negative predictive probability for dysplasia.

Subsequently, a more refined but complex combination of five different fine mucosal patterns (FMP) and capillary patterns (CP) classification was put forth by Goda et al. from Tokyo to describe various appearances that is predictive for detecting specialized intestinal mucosa and superficial Barrett’s adenocarcinoma. Of all patterns, a cerebriform FMP and ivy-like or DNA-spiral like appearance of CP most predicted specialized intestinal mucosa. The cerebriform FMP pattern consists of complicated branching and unions. As for the ivy-like or DNA-spiral like appearance of CP, it involves spirals and complicated branching with adjacent capillaries showing interconnections resembling a chain or net. In their cohort, all cases of superficial or intramucosal adenocarcinoma (which in Western countries may have been termed high-grade dysplasia) demonstrated both irregular FMP and CP with 100% sensitivity and specificity. Irregular FMP is characterized by irregularity in shape and branching in mucosal pattern and irregular CP is characterized as irregular micro-vessels with irregular course and uneven forms.

In order to provide a practical NBI method to identify dysplastic lesions, a more simplified classification by Singh et al. was described by combining the pit patterns of the mucosal surface and the regularity of the microvasculature in four easily distinguishable types (Table 1).

Type A mucosa was indicative of columnar mucosa without intestinal metaplasia with positive predictive value of 100%. Type B and C were typical of intestinal metaplasia with positive predictive value of 90%. Type D was indicated of high grade dysplasia with 79% positive predictive value and more importantly 100% negative predictive value. It is also important to note that all of the above classifications with NBI patterns were unable to predict low grade dysplasia.
sia with confidence.

Singh et al.\(^{32}\) further simplified the classification system in 2014 where the microstructural pattern and microvascular architecture were classified into three subtypes (Table 2). Type 1 was characterized by regular pits and/or vasculature (Figs 1-3) including absent/round/oval/linear/cerebriform/villous pits with regular vasculature. Type 2 was characterized by irregular/absent pits and irregular vasculature (Fig. 4) and type 3 has equivocal features which may exhibit dilated vasculature but no change in caliber (Fig. 5). The performance of this classification to identify high grade dysplasia was tested in the Asia-Pacific Barrett's Consortium in 2014 involving endoscopists from 11 countries with varying experience from the Asia-Pacific region. They evaluated images of BE with dysplastic and non-dysplastic areas.

The results were affirmative for positive identification of high grade dysplastic BE areas with sensitivity of 90% and negative predictive value of 99% showing that such a classification could be readily adopted.\(^{32}\)

### ESOPHAGEAL SQUAMOUS CELL CARCINOMA

Esophageal squamous cell carcinoma (ESCC) remains at present the dominant histology of esophageal cancer worldwide.\(^{33}\) ESCC appears to be more prevalent in countries stretching from northeast China to the Middle East forming what is termed, the high risk "Asian esophageal cancer belt.\(^{34}\) It is 2 to 4 times more frequent in males.\(^{35}\) Risk factors for the development of ESCC include smoking and alcohol consumption. Smoking increases the risk of developing ESCC by 5-fold.\(^{36}\) The reported odds are also greater for former smokers when compared to those who

| Table 1. Nottingham Classification for Esophageal Adenocarcinoma |
|---------------------------------------------------------------|
| **Type** | **Description** | **Histology** |
|---|---|---|
| A | Round pits with regular microvasculature | Columnar mucosa |
| B | Villous/ridge pits with regular microvasculature | Intestinal metaplasia |
| C | Absent pits with regular microvasculature | Intestinal metaplasia |
| D | Distorted pits with irregular microvasculature | High grade dysplasia |

| Table 2. Asia-Pacific Barrett’s Consortium Classification |
|----------------------------------------------------------|
| **Type** | **Description** | **Histology** |
|---|---|---|
| 1 | Regular pits and/or vasculature | No dysplasia |
| 2 | Irregular/absent pits and irregular vasculature | High grade dysplasia |
| 3 | Equivocal, area may exhibit dilated vasculature but no change in caliber | Not clear/unsure |

**Fig. 1.** Round pits and regular microvasculature (no dysplasia).

**Fig. 2.** Ridged/villous pits and regular microvasculature (no dysplasia).

**Fig. 3.** Absent pits and regular microvasculature (no dysplasia).
never smoked. With alcohol, the inherited enzyme deficiency of aldehyde dehydrogenase 2 which is involved in alcohol metabolism has been associated with increased risk of ESCC.\textsuperscript{37} It was reported that an estimated 36% of East Asians (Japanese, Chinese, and Koreans) are deficient in this enzyme leading to the “Asian Flush or Glow” response characterized by symptoms of facial flushing, nausea and tachycardia after alcohol consumption.\textsuperscript{38} Other risk factors include dietary aspects such as the consumption of food with high concentration of nitrogenous compounds,\textsuperscript{36} hot beverages like coffee, tea and mate (caffeine-rich infused drink)\textsuperscript{39} and genetic related etiologies such as tylosis.\textsuperscript{40}

It is well known from epidemiologic studies that ESCC risk factors differ from geographical regions and ethnicity making the implementation of a uniformed screening program challenging.\textsuperscript{34-36} Additionally, early diagnosis without a screening program is not possible given that most patients remain asymptomatic. Therefore, certain communities and countries provide a tailored approach focused heavily on the community who are at risk. This has been performed in China where endoscopic screening is employed to detect dysplastic lesions in high-risk asymptomatic patients.\textsuperscript{41} The risk of developing ESCC not surprisingly is strongly associated with increasing grades of dysplasia (relative risk, 15.3 to 52.4 for severe dysplasia) and carcinoma \textit{in situ} (relative risk, 16.6 to 71.4).\textsuperscript{42}

**NBI OF SQUAMOUS CELL CARCINOMA**

Unlike gastric, intestinal, colonic or Barrett’s mucosa, squamous epithelium lacks mucosal contours where “disturbances” in pit patterns can be visualized. Hence NBI’s ability to provide high contrast images of the microvascular structure plays a key role in describing ESCC at different stages. Numerous studies to date have identified and characterized the intrapapillary capillary loops (IPCL) patterns as major determinants of early ESCC characteristics. This remarkable observation was spearheaded by Inoue and colleagues where a classification of IPCL patterns identified by NBI with magnification was developed to correlate with the intraepithelial neoplastic changes and the depth of invasion.\textsuperscript{43-45}\textsuperscript{45} The progressive abnormal changes of IPCL patterns include the degree of dilatation, tortuosity, caliber and variation in shapes. Such progressive abnormal changes allow the endoscopist to predict the degree of neoplasia, its invasion depth and allows determination of the feasibility of endoscopic resection as a curative procedure.\textsuperscript{45-47}

In normal mucosa, IPCL arises perpendicularly or vertically in thin loop like structures. With reflux esophagitis, the IPCL pattern may become slightly more elongated and dilated. In the presence of tissue atypia or dysplastic changes, the IPCL becomes remarkably dilated with irregularity of the vessel caliber. The risk of developing ESCC not surprisingly is strongly associated with increasing grades of dysplasia (relative risk, 15.3 to 52.4 for severe dysplasia) and carcinoma \textit{in situ} (relative risk, 16.6 to 71.4).\textsuperscript{42}

Fig. 4. Irregular/absent pits and irregular microvasculature [high-grade dysplasia].

Fig. 5. Dilated vasculature but no change in caliber [equivocal].
area of 2.5 mm compared to ESCCs invading deeper than 200 µm into submucosa which showed a mean avascular area of 9 mm.

Goda et al. 49 demonstrated non-inferior diagnostic potential of magnifying endoscopy with NBI when compared to non-magnifying high-resolution WLE and high-frequency endoscopic ultrasonography with no significant difference in sensitivity and specificity. In comparison, NBI with magnification also reduced invasion depth over-estimation translating to the right therapeutic intervention which has significant implications on morbidity and mortality.

Aside from observing IPCL patterns, Ishihara and colleagues made observations of brownish dots and epithelial appearance on NBI as another feature associated with high grade intraepithelial neoplasia or invasive cancer. 49 Analysis into these findings were investigated and reproduced in a study by Kanzaki et al. 50 where brownish epithelium were confirmed to be significantly associated with high grade intraepithelial neoplasia and ESCC. 50 The appearance of brown colored neoplastic area was as a result of epithelial and keratinous layer thinning due to the neoplastic process. This inspired a retrospective observation which validated the usefulness and ease of NBI over WLE in detecting superficial esophageal neoplasia (<10 mm diameter lesions) with brownish dots and epithelium. 51 The potential of NBI in highlighting superficial cancers continues to be recognized over conventional WLE and chromoendoscopy with Lugol’s iodine. 52,53 The chances of overlooking suspicious lesions could certainly be minimized by the supplemental utility of NBI given the astounding report that as high as 7.8% of esophageal cancer lesions were missed in the United Kingdom. These were patients had an endoscopy done 3 to 36 months preceding diagnosis. 54

In regards to the usefulness of NBI in assessing depth of invasion for superficial ESCC, there was a conflicting multicenter prospective study published in 2015 by Ebi et al. 55 It reported that magnifying endoscopy with NBI to be not superior to conventional endoscopy in assessing the depth of invasion of superficial ESCC. The authors reported accuracy of magnifying endoscopy with NBI (65.3%) versus accuracy of conventional endoscopy (71.4%, p=0.375). However, this study had a relatively small sample size involving 49 lesions. The result comparing the accuracy between magnifying endoscopy with NBI and conventional endoscopy with a p-value of 0.375 was not statistically significant.

In 2017, another prospective multicenter study using magnifying endoscopic NBI (ME-NBI) in predicting the invasion depth of superficial squamous cell carcinoma was published by Oyama et al. 56 The study involved 211 patients with superficial ESCC. The study found that ME-NBI had an overall accuracy rate of 90.5% for predicting cancer invasion depth based on type B1, B2 and B3 microvessels pattern. The authors concluded that ME-NBI is of sufficient high accuracy for clinical use.

The study by Oyama et al. 56 was based on the recently revised 2017 Japan Esophageal Society guideline classifying superficial ESCC with magnifying endoscopy and NBI. This was a simplified classification proposed for estimating the depth of invasion of superficial ESCC by observing the microvascular patterns. The AB classification (Table 3 and Figs 6-9) is composed of two types of vessels with type A vessels (lacking severe irregularity) indicating non-cancer subtype while type B vessels are related to cancer.

Type B microvessels are sub-classified further into three groups, B1, B2 and B3. B1 predicts invasion depth confined within the epithelium or lamina propria where endoscopic resection is still feasible. B2 however suggests invasion into muscularis mucosa or beyond to the superficial layers of submucosa where endoscopic resection could still be a

Table 3. AB Classification for Esophageal SCC

| Vessel type | Definition | Invasion depth |
|-------------|------------|----------------|
| A           | Normal or abnormal IPCL without severe irregularity | Noninvasive |
| B1          | Severe irregularity/dilatation of IPCL with loop-like formation | High grade intraepithelial neoplasia or SCC limited to lamina propria |
| B2          | Severe irregularity/dilatation of IPCL with loss of loop-like formation | SCC involving muscularis mucosa and <200 µm depth of submucosa |
| B3          | Highly dilated irregular vessels more than 3 times B2 vessel | >200 µm depth of submucosal invasion |

SCC, squamous cell carcinoma; IPCL, intrapapillary capillary loop.

Fig. 6. Diagram showing invasion depth according to different types of intrapapillary capillary loop patterns. EP, epithelium; LP, lamina propria; MM, muscularis mucosa; SM, submucosa; MP, muscularis propria.
relative indication. B3 subtype marks the contraindication for endoscopic resection as it traverses more than 200 µm into the submucosa where risks of lymphovascular invasion increases exponentially.56

CONCLUDING REMARKS

The role of NBI in the esophagus continues to impact major clinical decision in early cancer management. Its effectiveness in terms of ease of use, high sensitivity and accuracy has been proven. However, the interpretation of mucosal surface and microvasculature pattern requires training. Classifications have been revised to encourage uniformity and simplify ease of learning. As a recommendation, the simplified Asia-Pacific Barrett’s Consortium Classification for BE and the AB classification by Japanese Esophageal Society for ESCC could be adopted.

CONFLICTS OF INTEREST

No potential conflict of interest relevant to this article was reported.

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