Narrow-band imaging and white-light endoscopy with optical magnification in the diagnosis of dysplasia in Barrett’s esophagus: results of the Asia-Pacific Barrett’s Consortium

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Introduction

Barrett’s esophagus (BE) is the only known pre-malignant precursor of esophageal adenocarcinoma, a tumor that is rapidly increasing in the developed world [1]. At present, endoscopic surveillance with white-light endoscopy (WLE) and random 4-quadrant biopsies is considered the standard of care for detecting intestinal metaplasia and neoplasia in BE. However, current recommended guidelines for surveying patients with BE are time consuming and poorly adhered to [2]. Over the last decade, new endoscopic imaging technologies have shown great promise in detecting dysplasia and early neoplasia in BE [3–8]. Histology can be predicted with high accuracy potentially making surveillance cost-effective. Recent data have emerged that a targeted biopsy approach using Narrow-Band Imaging (NBI) could be considered in patients undergoing surveillance for BE [9]. Given the low prevalence in BE, endoscopists from the Asia Pacific region may be less experienced with this condition. On the other hand, they may be well versed with image enhancing endoscopic modalities given the high prevalence of gastric cancers, especially in East Asia. We embarked on a study to evaluate the performance characteristics of WLE with optical magnification (WLE-z), NBI with optical magnification (NBI-z) and a combination of both modalities given the high prevalence of gastric cancers, especially in East Asia. We embarked on a study to evaluate the performance characteristics of WLE with optical magnification (WLE-z), NBI with optical magnification (NBI-z) and a combination of both modalities in the assessment of patients presenting for surveillance endoscopy for BE amongst a large group of endoscopists from the Asia Pacific region.

Methods

A total of 28 endoscopists from 11 countries (Asia-Pacific Barrett’s Consortium) who have an interest in BE were invited to participate in an online study. Fifty images of WLE-z and 50 corresponding images of NBI-z in BE with corresponding histopathology were prospectively collected.

Objective
The advent and utility of new endoscopic imaging modalities for predicting the histology of Barrett’s esophagus (BE) in real time with high accuracy appear promising and could potentially obviate the need to perform random biopsies where guidelines are poorly adhered to. We embarked on evaluating the performance characteristics of white-light endoscopy with magnification (WLE-z), narrow-band imaging with magnification (NBI-z) and a combination of both modalities.

Design
This was a prospective online study with 28 endoscopists from 11 countries (Asia-Pacific Barrett’s Consortium) who have an interest in BE were invited to participate in an online study. Fifty images of WLE-z and 50 corresponding images of NBI-z in BE with corresponding histopathology were prospectively collected.

Results
NBI-z and a combination of both were calculated. The overall Acc for WLE-z, NBI-z and a combination of both were calculated. The high NPV (99%) when both WLE-z and NBI-z images were placed side by side, the Acc increased to 90.3%. The Sn, Sp, PPV, and NPV of WLE-z were 48%, 92%, 45%, and 93% while with NBI-z, these improved to 89%, 89%, 56%, and 98%, respectively. When both imaging modalities were viewed together, they improved further to 93%, 90%, 61%, and 99%.

Conclusion
The high NPV (99%) when both WLE-z and NBI-z were used simultaneously indicates that areas with regular appearance that are diagnosed with confidence can effectively be left alone and not biopsied when performed at a skilled resource center. This approach could potentially lead to a paradigm shift of how patients with BE are assessed.
**Image acquisition**

All images were taken by a single gastroenterologist (RS) with expertise in advanced endoscopic imaging at a single Australian center, which has the required equipment. All procedures were performed using the commercially available Olympus 160Z endoscope with optical magnification (×115) capability. A transparent cap was used and adjusted to a length of 2mm at the tip of the endoscope. After obtaining informed consent, all patients were offered local pharyngeal anesthesia with xylocaine spray followed by induction and maintenance of deep sedation with intravenous Midazolam and Propofol. It is imperative to visualize the mucosa clearly, hence patients were given a mucolytic agent, N-acetylcysteine, and a defoaming agent, Simethicone, mixed with water (50mL) to drink before the procedure. Following intubation of the esophagus, an additional 10–20mL of the mucolytic agent was flushed to rid the surface of any adherent mucus. This was followed by flushing the surface with dilute adrenaline (1:20 000)(10–20mL) to reduce interference with blood and improve mucosal visualization after biopsies had been taken. To minimize movement artifacts and, if deemed necessary by the endoscopist, sedation was titrated further. If the esophagus exhibited excessive peristaltic activity, an antispasmodic agent, hyoscine-N-butyrbromide (Buscopan, 10–20mg), was given intravenously. The esophagus was first examined in the overview mode (without activation of NBI or optical magnification function). The endoscopist recorded the length of Barrett’s segment according to Prague’s C&M criteria and any macroscopically evident lesions. Only inconspicuous, flat areas or areas adjacent to macroscopically visible lesions in the Barrett’s segment were further assessed in the NBI-z mode. We ensured that the imaged view was the area biopsied by applying light suction pressure to the mucosa, thus enabling the cap to fix to the imaged area. All images were then stored as high definition JPEG files (200–300 kb, 1280×1024 pixel array, 32-bit color). Corresponding biopsies of each imaged area were placed in separate labeled pots filled with 10% buffered formalin and sent for histopathological analysis by a single gastrointestinal pathologist with expertise in Barrett’s esophagus. Only clear images with corresponding histopathology were selected. A total of 100 images were obtained from 35 patients.

**Learning phase**

A short learning Microsoft PowerPoint presentation was given to all of the endoscopists at the Asia Pacific Digestive Week 2011 held in Singapore. This session lasted for 1 hour. In addition, a total of 30 sample images were shown to the participants. They were also given sample PowerPoint slides as a guide after the education session. Subsequently, all participants were invited to predict the histology of the images, which were presented on a web portal for direct online assessment.
Classification
A simplified classification based on the microstructural pattern and microvascular architecture was used to grade the images:
(1) Regular pits and/or vasculature (R): no dysplasia (Fig. 1–3)
(absent/round/ovar/linear/cerebiform/villous pits with regular vasculature).
(2) Irregular/absent pits and irregular vasculature (IR): high grade dysplasia (Fig. 4).
(3) Equivocal (E): not clear/unsure. Area may exhibit dilated vasculature but no change in caliber (Fig. 5).
(1) and (2) were graded with confidence whereas (3) was graded no confidence.

Image evaluation
The 100 images were then randomly mixed and displayed as 100 single images on an online portal which consisted of three parts:
- Part-I: participant’s information.
- Part-II comprising of 100 WLE-z/NBI-z images, and
- Part-III comprising of 50 NBI-z images placed alongside the corresponding WLE-z images. This last exercise was done to simulate real-time endoscopy where both imaging modalities are often interchanged. No clinical data was revealed to the evaluators. The endoscopists were given specific instructions to complete the study within 1 month of the learning session (online portal webpage was closed thereafter). The overall Accuracy (Acc), Sensitivity (Sn), Specificity (Sp), Positive Predictive Value (PPV), and Negative Predictive Value (NPV) of WLE-z, NBI-z and the combination were then determined.

Results
Endoscopist characteristics
A total of 28 endoscopists representing 11 countries in the Asia Pacific region with a diverse range of experience participated in the study. Twelve (43%) had more than 10 years of experience as a consultant, 3 (11%) 8 – 10 years, 2 (7%) 5 – 7 years, 6 (21%) 2 – 4 years and 5 (18%) less than 2 years. The number of endoscopies performed by the evaluators ranged from 500 to 3000 per year with 4 (14%) doing more than 3000 procedures per year and 5 (18%) performing less than 500 per year. Table 1 summarizes the average number of BE seen by endoscopists per year.

Image evaluation: overall
A total of 4200 images were available for evaluation (28 endoscopists; 150 images each). These images were obtained from 43 non-dysplastic and 7 dysplastic areas. The assessors were able to grade 3370 images with confidence (80.2%). The overall Acc for WLE-z and NBI-z images were 87.1% and 88.7%, respectively. When images from the two modalities were placed side by side, the Acc increased to 90.3%. The Sn, Sp, PPV, and NPV of WLE-z were 48%, 92%, 45%, and 93%, while with NBI-z, these values improved to 89%, 89%, 56%, and 98%, respectively. When both imaging modalities were viewed together, these values improved further to 93%, 90%, 61%, and 99% (Table 2).

Table 1 Average number of Barrett’s esophagus (BE) patients seen yearly by assessors.

<table>
<thead>
<tr>
<th>Average number of BE seen/year</th>
<th>Number</th>
</tr>
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<tbody>
<tr>
<td>&lt;20</td>
<td>19</td>
</tr>
<tr>
<td>21 – 50</td>
<td>5</td>
</tr>
<tr>
<td>51 – 100</td>
<td>3</td>
</tr>
<tr>
<td>&gt;100</td>
<td>1</td>
</tr>
</tbody>
</table>

Table 2 Sensitivity (Sn), Specificity (Sp), Positive Predictive Value (PPV) and Negative Predictive Value (NPV) in the assessment of high grade dysplasia vs. no dysplasia with white-light endoscopy and optical magnification (WLE-z), narrow-band imaging and optical magnification (NBI-z) and a combination of both (95% confidence interval).

<table>
<thead>
<tr>
<th></th>
<th>WLE-z</th>
<th>NBI-z</th>
<th>Combination</th>
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<tbody>
<tr>
<td>Sn, %</td>
<td>48.4</td>
<td>89.1</td>
<td>93.4</td>
</tr>
<tr>
<td>Sp, %</td>
<td>92.2</td>
<td>89.3</td>
<td>90.3</td>
</tr>
<tr>
<td>PPV, %</td>
<td>45.6</td>
<td>56.5</td>
<td>61.7</td>
</tr>
<tr>
<td>NPV, %</td>
<td>93.5</td>
<td>98.3</td>
<td>99.2</td>
</tr>
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</table>

Table 3 Narrow-band imaging (NBI) experience of participants and accuracy in detecting dysplasia (95% confidence interval).

<table>
<thead>
<tr>
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<th>Not experienced with NBI</th>
<th>Experienced with NBI</th>
</tr>
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<tbody>
<tr>
<td>WLE-z</td>
<td>86.7 (84.2 – 88.9)</td>
<td>88.3 (84.0 – 91.8)</td>
</tr>
<tr>
<td>NBI-z</td>
<td>87.9 (85.5 – 90.0)</td>
<td>90.9 (87.2 – 93.8)</td>
</tr>
<tr>
<td>Combination</td>
<td>89.0 (86.7 – 91.9)</td>
<td>94.6 (91.0 – 97.0)</td>
</tr>
</tbody>
</table>

Table 4 Participant experience with optical magnification and accuracy in detecting dysplasia (95% confidence interval).

<table>
<thead>
<tr>
<th></th>
<th>Not experienced with optical magnification</th>
<th>Experienced with optical magnification</th>
</tr>
</thead>
<tbody>
<tr>
<td>WLE-z</td>
<td>86.4 (83.2 – 89.2)</td>
<td>87.7 (84.8 – 90.3)</td>
</tr>
<tr>
<td>NBI-z</td>
<td>88.5 (85.6 – 91.0)</td>
<td>88.8 (86.1 – 91.2)</td>
</tr>
<tr>
<td>Combination</td>
<td>87.3 (83.9 – 90.2)</td>
<td>92.7 (90.4 – 94.7)</td>
</tr>
</tbody>
</table>

Image evaluation: by experience with NBI and optical magnification
Twenty participants (71%) routinely use NBI in daily practice. The overall Acc in detecting dysplasia in BE for experienced endoscopists was 95% in contrast to 89% for endoscopists not experienced with NBI when both modalities were used (P<0.05) (Table 3). Fourteen participants (50%) have experience with optical magnification technology. An Acc of 93% was achieved by endoscopists with experience in optical magnification when the combined modalities of NBI-z and WLE-z were used compared to 87% with endoscopists who were not experienced with this technique (P<0.05) (Table 4).
Intraobserver agreement
The intraobserver agreement (kappa) for WLE-z, NBI-z, and the combination was 0.31, 0.49, and 0.56, respectively.

Discussion
This study looked at two endoscopic imaging modalities diagnosing dysplasia in patients who had BE. Current surveillance and practice guidelines for BE are not widely followed as there is a marked variability in the technique and interval of surveillance [10, 11]. The identification of dysplasia may be difficult and distribution is multifocal leading to sampling error in even the most thorough biopsy surveillance protocols [12]. The American Society of Gastrointestinal Endoscopy’s Preservation and Incorporation of Valuable endoscopic Innovations guidelines [13] on imaging technology in BE recommend that an imaging technology with targeted biopsies should have a per-patient sensitivity of 90% or greater and a negative predictive value (NPV) of 98% or greater for detecting high grade dysplasia/early esophageal adenocarcinoma. In the present study, a high NPV of 99% could be achieved when WLE-z and NBI-z were used, although it must be taken into context that this was an analysis performed on a per-image rather than a per-patient basis.

NBI with optical magnification has been extensively studied in the past with a myriad of different classification systems [14–16]. Baldaque-Silva et al. [17] recently concluded that structured learning in endoscopic assessment of BE using magnification and NBI did not have an impact on accuracy based on the Amsterdam classification system. The authors obtained sensitivity for prediction of neoplasia ranging between 62% and 90%. The same group also compared three different classification systems (Nottingham, Kansas, and Amsterdam) for BE using NBI and magnification with nine endoscopists from Europe and Japan [18]. The global accuracy for detection of non-dysplastic and dysplastic lesions using these three classifications was rather disappointing at 46%, 47%, and 51%, respectively. Our study differs from that of the above as a simplified classification system was used. The evaluators in this study also had a learning phase and appeared to be well versed with the classification system before embarking on the online module, although the image acquisition process and selection bias as described previously could have also led to the higher NPV in this study.

This study has some limitations. A single endoscopist with extensive experience in mucosal imaging obtained all of the images. This may not be generalizable to the community Gastroenterologist. However, with the advent of newer endoscopic imaging devices, obtaining clear, crisp images is becoming increasingly easier. Another drawback is that the images were evaluated ‘post hoc’ and not in real time. There were also no low grade dysplastic (LGD) lesions evaluated in the study. This was deliberately done as NBI-z does not reliably diagnose LGD. It is well known though that there is significant interobserver variability in the diagnosis of LGD even amongst expert gastrointestinal pathologists [19].

In conclusion, this study demonstrated that a high NPV in the prediction of non-dysplastic areas in BE can be achieved with the application of a combination of WLE-z and NBI-z. It may thus be plausible to refrain from biopsying regular appearing areas, resect irregular appearing areas (both of which when diagnosed with high confidence and performed at skilled resource center), and biopsy only equivocal areas. This cost and time saving approach could potentially lead to a paradigm shift in how patients with BE are surveyed. Real-time prospective evaluation of dysplasia in BE will further consolidate and support the findings of this study.

Competing interests: None

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