The Role of Acetic Acid Chromoendoscopy and Narrow Band Imaging in Diagnosis of Esophageal Barrett Dysplasia

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ABSTRACT

Background: Barrett’s esophagus (BE) is an important risk factor for the occurrence esophageal adenocarcinoma. Endoscopic surveillance for Barrett’s dysplasia is indicated in all patients with BE. Recent data found that with the use of advanced endoscopic techniques there is an increased diagnostic yield for dysplasia.

Aim: The aim of this study is to assess the accuracy of acetic acid assisted narrow band imaging (AA-NBI) endoscopy in detection of Barrett’s dysplasia. Patients and Methods: Forty patients with BE surveyed for Barrett’s dysplasia by AA-NBI endoscopy. Patients with difficult sampling (n=1) or not proved to have Barrett’s mucosa by histopathology (n=5) were excluded. Patients with positive and negative dysplasia underwent targeted and random endoscopic biopsy, respectively. The diagnosis of BE based on Prague endoscopic classification. The pathological assessment of Barrett’s dysplasia based on Vienna classification. Results: A total of 34 patients were included; their mean age was 39.03±14.23 years; they were 19 (55.88%) males and 15 (44.11%) females. Of them 28 (82.35%) had cardiac type BE, 3 (8.82%) had fundic type BE and 3 (8.82%) had intestinal type BE. Five patients were depicted as Barrett’s dysplasia by AA-NBI. Low grade dysplasia was proved in 3 of them by histopathology. The sensitivity, specificity, positive predictive value, negative predictive value and accuracy of AA-NBI in diagnosis of Barrett’s dysplasia were 100%, 93.5%, 60%, 100%, and 94.1% respectively. Conclusion: Nine percent of patients with BE are positive for dysplasia. AA-NBI has promising sensitivity, specificity and accuracy in prediction of Barrett’s dysplasia.

Keywords: Barrett’s esophagus. Dysplasia. Acetic acid chromoendoscopy. Narrow band imaging. esophageal adenocarcinoma.

INTRODUCTION

The incidence of esophageal cancer is rising 1 representing the seventh most common cancer worldwide. It is well known that Barrett’s esophagus (BE) is considered an important risk factor for the occurrence of esophageal adenocarcinoma (EAC) and is present in up to 1.6% of the general population 2 and in up to 20% of patients suffering from gastroesophageal reflux disease (GERD) 3.

It is postulated that the development of EAC in Barrett occurs in a progressive fashion from intestinal metaplasia (IM) to low grade dysplasia (LGD) to high grade dysplasia (HGD) and then EAC. The annual rate of transformation into EAC in BE patients without dysplasia is estimated to be between 0.07% and 0.82% 4. While the annual rate of progression from LGD to HGD or EAC is as high as 8.8% 5 and from HGD to EAC is 12% to 40% 6.

The aim of endoscopic surveillance for Barrett dysplasia is to change the natural history of the disease through recognizing dysplasia at an earlier stage and thus introducing beneficial endoscopic treatment. Established surveillance protocols suggest Seattle protocol with taking targeted biopsies of visible lesions and random four quadrant biopsies (4QBS) every 1-2 cm which seemingly offers the maximum yield of dysplasia in comparison with other biopsy protocols 7. However, there are several disadvantages to this Seattle protocol including only 13% of early neoplastic lesions appearing as visible nodules 8, a major proportion of Barrett’s neoplasia is not visible on high definition white-light endoscopy alone, with reported sensitivity in the range 40%-64% and specificity 98%-100% 9. In addition, 4QBS are time-consuming, costly and poorly accepted by patients. These drawbacks have encouraged evaluation of more effective techniques to improve the diagnostic accuracy for the detection of IM and early Barrett’s dysplasia used in surveillance 10.

PATIENTS AND METHODS

The study was performed on 40 consecutive adults with BE who required endoscopic examination at El-Hussein University Hospital between December 2017 and April 2019, Department of Internal Medicine, Gastroenterology and Hepatology Unit.

Patients with advanced esophageal cancer, previous esophageal resection, on-going treatment with antiplatelet medication, or anticoagulant medication, and those with hemorrhagic diseases, were excluded.

Ethics and Patient Consent

The study was approved by the Ethics Board of Al-Azhar University and an informed written consent was taken from each participant in the study.

Endoscopic procedure:

Surveillance of BE was conducted by white light imaging (WLI), and narrow band imaging (NBI). Endoscopic diagnosis of BE was based on the detection of
the characteristic “tongues” or salmon-pink colored extensions of mucosa that grow into the esophagus above the gastroesophageal junction using Prague criteria to determine circular and maximum length of Barrett segment. \(^7\) Assessment of Barrett’s dysplasia was carried on by magnified WLI, magnified NBI and acetic acid chromoendoscopy. The magnified images were analyzed with respect to pit-patterns, regular mucosal patterns (circular, ridge/villous, tubular), irregular mucosal patterns (absent or irregular) and vascular patterns (regular or irregular) using BING criteria \(^11\). Following this, 10–20 mL of 2% acetic acid (AA) was sprayed by using spray catheter via the instrument channel over the lower esophagus and Barrett’s epithelium. At first Barrett’s mucosa took on a white appearance (the aceto-white reaction), while dysplastic mucosa was turned red earlier than the non dysplastic Barrett’s mucosa \(^12\).

The positions of the lesions detected by WLE, NBI or AA were recorded and photographed to confirm the accuracy of the biopsies obtained. At least one targeted biopsy was collected separately from the endoscopic lesions suspicious of Barrett’s dysplasia by AA-NBI or WLE. In absence of Barrett’s dysplasia 2-4 random biopsies were taken from the different sites of barrette’s mucosa. All real time endoscopic assessment was carried by one author.

**Assessment of video records and photographs:**

A second session of assessment was conducted by two authors using the video records and photographs to confirm the results of real time assessment session. In case of conflict between both sessions, agreement was reached by consensus.

**Histopathology:**

All biopsy specimens were fixed in 4% formalin and embedded in paraffin and the slides were processed by using hematoxylin and eosin (H&E) stains. All histologic analyses were performed by single experienced pathologist who was uninformed about the results of endoscopy. The biopsy specimens were analyzed for the presence of specialized intestinal metaplasia, dysplasia and/or cancer by using Vienna classification \(^13\).

**Statistical Analysis**

For the per-patient analysis, the lesions suspected by endoscope in one patient were considered one unit of analysis in this evaluation. The sensitivity, specificity, positive predictive values, negative predictive values and accuracy for the prediction of Barrett’s dysplasia in the AA-NBI were calculated using histology as a reference value.

**RESULTS**

This study included 40 patients diagnosed as having Barrett’s Esophagus by endoscopy 5 of them were false positive endoscopic results for BE (negative by histopathology), the causes of this false results were chronic active reflux esophagitis (n=3), chronic active reflux esophagitis with ulceration (n=1) and wrong sampling (n=1) and 1 case was difficult to be biopsied. These 6 cases were excluded from the final analysis. The remaining 34 patients were diagnosed to have BE by endoscopy, which was confirmed by histopathology.

**Demography and clinical data:**

Among the demographic and clinical data of 34 patients with BE the mean age was 39.03±14.23 years; they were 19 (55.88%) males and 15 (44.12%) females. Of them 15 patients (44.1%) had no history of GERD while 19 patients (55.9%) had history of GERD. 10 patients (52.6%) had GERD < 1 year ago, 6 patients (31.6%) had GERD 1 – 3 years ago and 3 patients (15.8%) had GERD > 3 years ago. As regard indications of endoscopy, abdominal (epigastric) pain was recorded in 19 patients, vomiting was recorded in 9 patients, bleeding per rectum was recorded in 1 patient, chronic diffuse abdominal pain was recorded in 1 patient, heart burn was recorded in 6 patients, screening for malignancy was recorded in 2 patients, melena was recorded in 1 patient, hematemesis was recorded in 1 patient, iron deficiency anemia was recorded in 2 patients, regurgitation was recorded in 2 patients and esophageal dilation was recorded in 1 patient. Smoking were recorded in 11 patients (32.4%) while 23 patients (67.6%) were non-smoker.

**Endoscopic data:**

As regarding the endoscopic data of all patients with BE (34 patients); 30 (88.2%) had regular mucosal pattern, 4 (11.8%) had irregular mucosal pattern, 29 (85.3%) had regular vascular pattern and 5 (14.7%) had irregular vascular pattern on NBI by using Barrett’s International NBI Group (BING) criteria, on AA-WLE and AA-NBI 29 (85.3%) had BE with no dysplasia and 5 (14.7%) had Barrett’s dysplasia with no difference between AA-WLI and AA-NBI. Also GERD was present in 15 (44.1%) and 19 (55.9%) had no GERD on endoscopy. Among 15 patients with GERD 8 (53.3%) had GERD grade B, 4 (26.7%) had GERD grade A and 3 (20%) had GERD grade C (Figure 1). 5 Patients were diagnosed to have Barrett dysplasia by endoscopy 2 of them had GERD and 3 had no GERD.
**Histopathological data:**

As regarding the histopathological data: patients with BE and no dysplasia were 31 (91.2%) and among 5 patients diagnosed with Barrett’s dysplasia by AA-NBI 3 (8.8%) of them had low grade dysplasia proved on histopathology. The type of Barrett epithelium identified on histopathology was 28 (82.35%) had cardiac type BE, 3 (8.82%) had fundic type BE and 3 (8.82%) had intestinal type BE. Among patients with low grade dysplasia all had gastric cardiac type (100%) of BE, chronic active reflux esophagitis and cardiac H. Pylori infection were recorded in 28 (82.4%) and 8 (23.5%) respectively, and among patients with low grade dysplasia all had chronic active reflux esophagitis with no H. pylori recorded in any case. The sensitivity, specificity, positive predictive value, negative predictive value and accuracy of AA-NBI in diagnosis of Barrett’s dysplasia were 100%, 93.5%, 60%, 100%, and 94.1% respectively (Figure 2).

In this study we used NBI BING criteria for identification of dysplasia in patients with Barrett esophagus according to the mucosal pit pattern and vascular pattern and we almost found results similar to results of study done by Sharma et al., which concluded that BING criteria can identify patients with Barrett dysplasia with 85% accuracy, 80% sensitivity, 81% positive predictive value (PPV), and 85% negative predictive value (NPV). These values were 92%, 91%, 89%, and 95%, respectively when dysplasia was evaluated with high confidence.

![Figure 2](image)

**DISCUSSION**

Our study illustrates that acetic acid chromoendoscopy and narrow band imaging have a promising role in assessment and prediction of esophageal Barrett dysplasia. 34 patients with BE were included in our study; 5 of them were diagnosed by AA-NBI as Barrett dysplasia based on magnified mucosal pit pattern, magnified vascular pattern and aceto-whitening reaction after acetic acid spray over the mucosa. Among the 5 cases diagnosed by endoscopy 3 of them confirmed by histopathology. In contrast, no any cases diagnosed as dysplasia on histopathology and missed by endoscopy.

In our study results demonstrated that the sensitivity and specificity of AA-NBI were 100% and 93.5% respectively which are in agreement with a study by Vázquez-Iglesias, et al., which reported high sensitivity 100% and specificity 97.7% of detection of dysplasia by acetic acid chromoendoscopy. However, there were only 13/100 of the patient included in the study had dysplasia.

Our findings also consistent with large series of studies by Longcroft-Wheaton, et al., which included 119 patients and concluded that acetic acid-assisted evaluation of Barrett’s esophagus detected dysplasia/neoplasia more than white light endoscopy, and showed sensitivity 95.5% and specificity 80% which were identical to that of histopathological analysis.

Our study results illustrated that the diagnostic accuracy of AA-NBI for Barrett dysplasia was 94.1% which is similar to those of Coletta, et al., who reported that acetic acid chromoendoscopy (AAC) has great diagnostic accuracy for recognizing dysplastic Barrett esophagus patients. This study included 13 prospective studies with pooled sensitivity, specificity, positive likelihood ratio, and negative likelihood ratio for all included studies (9 studies, 1379 patients) were 0.92 (95% confidence interval [CI], 0.83-0.97), 0.96 (95% CI, 0.85-0.99), 25.0 (95% CI, 5.9-105.3), and 0.08 (95% CI, 0.04-0.18), respectively.

In contrast, our findings don’t meet the findings of study by Curvers, et al., who found that addition of acetic acid chromoendoscopy and narrow band imaging to white light endoscopy didn’t increase the yield of recognizing dysplasia/early neoplasia in Barrett esophagus patients.

Sharma et al., reported that narrow band imaging improves detection of dysplasia more than white light image which meets our study results.

Kara et al., also found that narrow band image is superior than standard WLI in detection of Barrett dysplasia in descriptive study including 67 patients using histology as reference standard. In their study, detection of high grade dysplasia had sensitivity 94%, specificity 76%, PPV 64% and NPV 98% which are similar to results of our study. But when compared with high-definition white light endoscopy (HDWLE) revealed no differences in the detection rate, however NBI demanded fewer biopsies.

In this study we used NBI BING criteria for identification of dysplasia in patients with Barrett esophagus according to the mucosal pit pattern and vascular pattern and we almost found results similar to results of study done by Sharma et al., which concluded that BING criteria can identify patients with Barrett dysplasia with 85% accuracy, 80% sensitivity, 81% positive predictive value (PPV), and 85% negative predictive value (NPV). These values were 92%, 91%, 89%, and 95%, respectively when dysplasia was evaluated with high confidence.
In this study there were some limitations. First, many patients had esophagitis on endoscopy, which increased rate of false positive endoscopic results for BE. Second, the number of patients with Barrett esophagus included in this study was little (34 patients) and patients with dysplasia was also little (3 patients) making it difficult to draw significant conclusion from it with respect to rate of dysplasia detection.

The high diagnostic accuracy of AA-NBI for detecting and also for excluding Barrett dysplasia suggests that this procedure may be very beneficial to appliance in clinical practice for BE surveillance. AA-NBI should decrease the number of random biopsies which sometimes can miss dysplasia and facilitate target biopsies increasing yield of dysplasia. Acetic acid chromoendoscopy is safe, low cost, available and don’t need specialist training. NBI also is widely available, easily to use with visualization of mucosal and vascular pattern. So, and according to our study results AA-NBI can be used in Barrett esophagus surveillance for Barrett dysplasia.

Our study justifies the need for large study including large number of patients to make a strong conclusion for assessment of the role of AA-NBI in BE surveillance for Barrett’s dysplasia.

CONCLUSION

Nine percent of patients with BE are positive for dysplasia according to our results. Cardiac type columnar metaplasia represents the majority of patients with BE in our series. AA-NBI has promising sensitivity, specificity and accuracy in prediction of Barrett’s dysplasia. Our study justifies the need for large study to assess the role of AA-NBI in BE surveillance for detection of Barrett’s dysplasia.

REFERENCES