

# Effectiveness of computer-aided diagnosis of colorectal lesions using novel software for magnifying narrow-band imaging: a pilot study



## Authors

Naoto Tamai<sup>1,2</sup>, Yutaka Saito<sup>1</sup>, Taku Sakamoto<sup>1</sup>, Takeshi Nakajima<sup>1</sup>, Takahisa Matsuda<sup>1</sup>, Kazuki Sumiyama<sup>2</sup>, Hisao Tajiri<sup>3</sup>, Ryosuke Koyama<sup>4</sup>, Shoji Kido<sup>4</sup>

## Institutions

- 1 Endoscopy Division, National Cancer Center Hospital, Tokyo, Japan
- 2 Department of Endoscopy, The Jikei University School of Medicine, Tokyo, Japan
- 3 Department of Innovative Interventional Endoscopy Research, The Jikei University School of Medicine, Tokyo, Japan
- 4 Applied Medical Engineering Science, Graduate School of Medicine, Yamaguchi University, Yamaguchi, Japan

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## Corresponding author

Naoto Tamai, MD, PhD, Department of Endoscopy, The Jikei University School of Medicine, 3-25-8 Nishi Shinbashi, Minato-ku, Tokyo 105-8461, Japan  
Phone: +81-3-3433-1111 (Ext. 3181)  
Fax: +81-3-3459-4524  
[tamai-naoto@jikei.ac.jp](mailto:tamai-naoto@jikei.ac.jp)

## ABSTRACT

**Background and study aims** Magnifying narrow-band imaging (M-NBI) enables detailed observation of microvascular architecture and can be used in endoscopic diagnosis

of colorectal lesion. However, in clinical practice, differential diagnosis and estimation of invasion depth of colorectal lesions based on M-NBI findings require experience. Therefore, developing computer-aided diagnosis (CAD) for M-NBI would be beneficial for clinical practice. The aim of this study was to evaluate the effectiveness of software for CAD of colorectal lesions.

**Materials and methods** In collaboration with Yamaguchi University, we developed novel software that enables CAD of colorectal lesions using M-NBI images. This software for CAD further specifically divides original Sano's colorectal M-NBI classification into 3 groups (group A, capillary pattern [CP] type I; group B, CP type II+CP type IIIA; group C, CP type IIIB), which describe hyperplastic polyps (HPs), adenoma/adenocarcinoma (intramucosal [IM] to submucosal [SM]-superficial) lesions, and SM-deep lesions, respectively. We retrospectively reviewed 121 lesions evaluated using M-NBI.

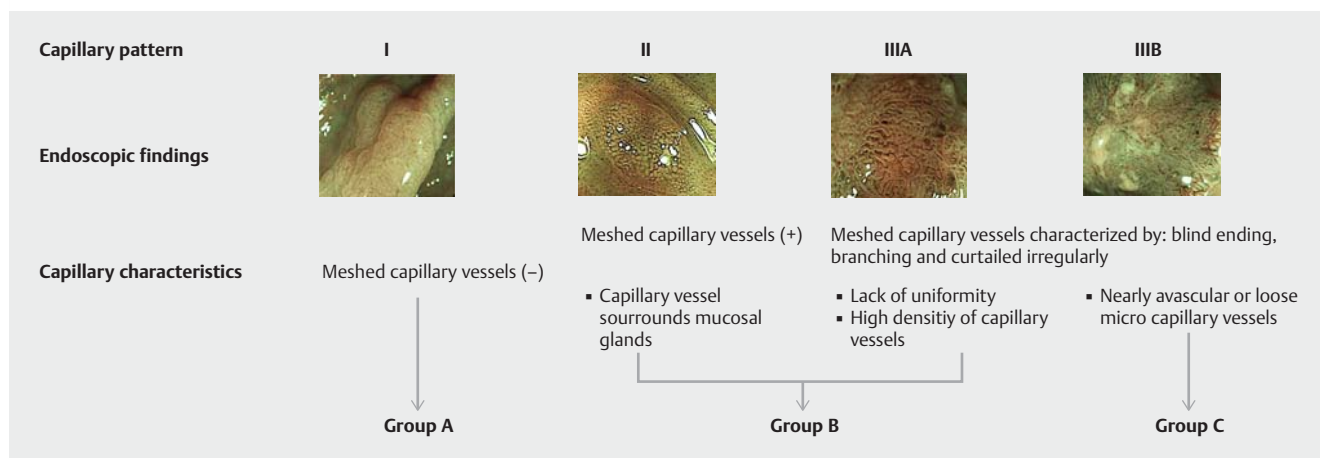
**Results** The 121 reviewed lesions included 21 HP, 80 adenoma/adenocarcinoma (IM to SM-superficial), and 20 SM-deep lesions. The concordance rate between the CAD and the diagnosis of the experienced endoscopists was 90.9%. The sensitivity, specificity, positive and negative predictive values, and accuracy of the CAD for neoplastic lesions were 83.9%, 82.6%, 53.1%, 95.6%, and 82.8%, respectively. The values for SM-deep lesions were 83.9%, 82.6%, 53.1%, 95.6%, and 82.8%, respectively.

**Conclusion** Relatively high diagnostic values were obtained using CAD. This software for CAD could possibly lead to a wider use of M-NBI in the endoscopic diagnosis of colorectal lesions.

## Introduction

Early detection and removal of colorectal neoplasia are essential in reducing mortality rates for patients with colorectal cancer [1, 2]. Although several modalities are available for colorectal cancer screening [3–6], colonoscopy is considered the most effective because it allows direct visualization and on-

site treatment of encountered lesions. Accurately determining the histological features of colorectal polyps using endoscopy could prevent unnecessary endoscopic treatments thus allowing the proposal of adequate surveillance recommendations [7]. Magnifying narrow-band imaging (NBI) enables detailed observation of the microvascular architecture of lesions and can be used for endoscopic differential diagnosis and estima-



► **Fig. 1** Sano's classification (modified for this study). In this study, Sano's classification was further divided broadly into three groups: group A, capillary pattern (CP) type I; group B, CP type II + CP type IIIA; group C, CP type IIIB.

tion of invasion depth of colorectal lesions [8,9]. We have developed software for computer-aided diagnosis (CAD) of colorectal lesions into nonneoplastic lesions (hyperplastic polyps [HPs]), adenoma/intramucosal (IM) cancers/submucosal (SM)-superficial cancers (invasion depth, <1000µm), or SM-deep cancers (invasion depth, ≥1000µm). The aim of this study was to retrospectively evaluate the effectiveness of CAD of colorectal lesions using our novel software and still images obtained from M-NBI. The aim of this study is to evaluate the effectiveness of CAD of colorectal lesions using novel software.

## Materials and methods

### Software for CAD

Using class-appropriate images obtained with front-view M-NBI, we determined the region of interest (ROI), converted the images into grayscale images, visualized the rough vessels through binarization using the moving average method, deleted all colors except the color of the vessel, and removed isolated points and holes. Subsequently, we used characteristics, such as the average width, length, total length, and concentration of the visualized vessel, and the fractal dimension to extract significant quantitative characteristics. Then, we extracted the average length, total length, and concentration of the visualized vessel as the effective quantitative characteristics for classifying colorectal lesions. In corroboration with Yamaguchi University, we developed software for CAD of lesions using the 3 quantitative characteristics. This software for CAD further broadly divides Sano's classification, which is the original colorectal NBI classification [10–12], into 3 groups (group A, capillary pattern [CP] type I; group B, CP type II + CP type IIIA; group C, CP type IIIB; ► **Fig. 1**), which describe hyperplastic lesions, adenoma/adenocarcinoma (IM-SM superficial) lesions, and SM-deep lesions, respectively.

The first step in using this software for CAD is to attach the M-NBI image to the central window. Subsequently, the ROI to be evaluated is determined using a cursor. Then, by clicking "Calculate" and "Classify" on the software for CAD interface, the M-NBI image can be automatically classified into 1 of 3 groups (► **Fig. 2**).

### Histological assessment

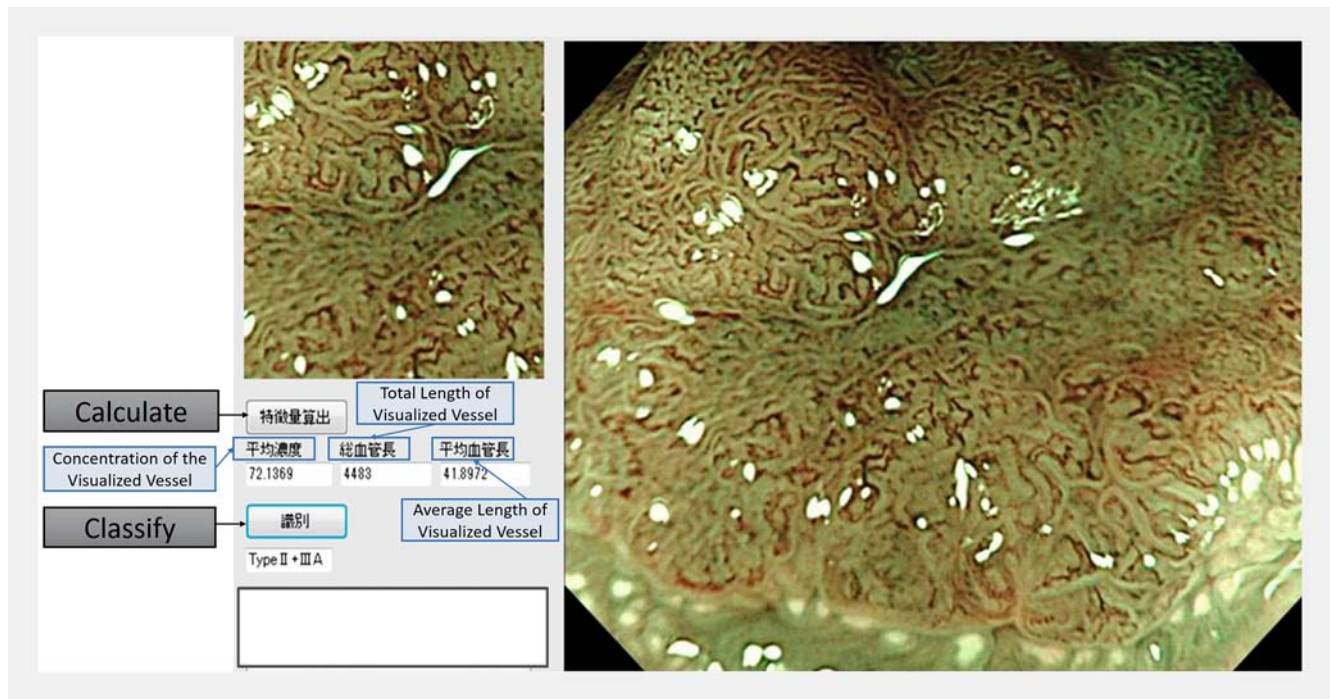
All specimens were fixed in 10% buffered formalin and cut into 2-mm slices. The specimens were then examined microscopically for depth of invasion and classified according to histological type. Histological diagnoses were based on the Japanese classification system of colon and rectal cancers and on the Vienna classification system.

### Statistical analysis

Statistical analysis was performed using SPSS for Windows (SPSS, release 6.0, 1993; SPSS Inc., Chicago, IL, USA). To determine differences in diagnostic accuracies between the CAD and diagnosis of experienced endoscopists for the differentiation and estimation of invasion depth of colorectal neoplasia, the Fisher exact and  $\chi^2$  tests were used. A *P* value <0.05 was considered statistically significant.

### Method

We retrospectively reviewed medical records of 103 consecutive patients with 121 lesions that were evaluated using magnifying colonoscopy (CF-H260AZI, CF-FH260AZI; Olympus Medical Systems) and M-NBI before endoscopic or surgical treatment of colorectal neoplasia at the Jikei University Hospital between June 2008 and March 2010. An experienced endoscopist who was blinded to this study selected the most representative M-NBI image of each colorectal lesion with the highest magnification and identified the ROI. Then, CAD, using our novel software, was performed on the ROI. Endoscopic diagnosis was also conducted using the selected M-NBI image, with the diagnosis made based on the agreement between 2 experienced endoscopists who were not involved in selection of M-NBI images



► **Fig. 2** The user interface of software for computer-aided diagnosis (CAD). The first step when using this software for CAD is to choose the region of interest using the cursor that appears on the attached narrow-band image (right-side window). Then, by clicking “Calculate” and “Classify,” the image is automatically classified into 1 of the 3 groups.

and identification of the ROI, and blinded to other endoscopic images of the lesions and the pathological information. All images were classified into the 3 groups described previously (groups A, B, or C). We then compared the results of the CAD using novel software and the diagnoses by the experienced endoscopists to evaluate the effectiveness of the CAD using novel software in the differential diagnosis and estimation of invasion depth of colorectal lesions.

## Results

Based on the histological assessment, the 121 reviewed lesions included 21 HPs, 80 adenoma/adenocarcinoma (IM to SM-superficial) lesions, and 20 SM-deep lesions. Among these lesions, 26 lesions were classified into group A, 79 were lesions were classified into group B, and 16 lesions were classified into group C by CAD. By contrast, 21 lesions were classified into group A, 82 lesions were classified into group B, and 18 lesions were classified into group C by the experienced endoscopists. The concordance rate between the CAD using our novel software and the diagnoses by the experienced endoscopists was 90.9% (► **Table 1**).

The sensitivity, specificity, positive and negative predictive values (PPV and NPV, respectively), and accuracy of the novel software for CAD in diagnosing neoplastic lesions were 95.0%, 100%, 100%, 80.8%, and 95.9%, respectively (► **Table 2**). For SM-deep lesions, the rates were 55%, 95.0%, 68.8%, 91.4%, and 88.4%, respectively (► **Table 3**).

► **Table 1** Concordance rates between computer-aided diagnosis using software and endoscopists' diagnosis.

	Group A (DE)	Group B (DE)	Group C (DE)
Group A (DS)	21	5	0
Group B (DS)	0	75	4
Group C (DS)	0	2	14

DE, diagnosis by the endoscopists; DS, diagnosis using the software for computer-aided diagnosis.

The concordance rate between computer-aided diagnosis using software and experienced endoscopists' diagnoses was 90.9% (110/121).

► **Table 2** Diagnostic value of the computer-aided diagnosis using software for neoplastic lesions.

	HP (nonneoplastic lesion)	Adenoma/adenocarcinoma (IM, SM-superficial, and SM-deep) (neoplastic lesion)
Group A (DS)	21	5
Group B + C (DS)	0	95

HP, hyperplastic polyp; IM, intramucosal; SM, submucosal; DS, diagnosis using the software for computer-aided diagnosis.

The diagnostic accuracy rate of the computer-aided diagnosis using software was 95.9% (116/121) for neoplastic lesions.

► **Table 3** Diagnostic value of the computer-aided diagnosis using software for SM-deep lesions.

	SM-deep lesions	Non-SM-deep lesions
Group A + B (DS)	9	96
Group C (DS)	11	5

SM, submucosal; DS, diagnosis using the software for computer-aided diagnosis.

The diagnostic accuracy rate of the computer-aided diagnosis using software was 88.4% (107/121) for SM-deep lesions.

► **Table 4** Diagnostic value of the endoscopists for neoplastic lesions.

	HP (nonneoplastic lesion)	Adenoma/adenocarcinoma (IM to SM-superficial)/SM-deep
Group A (DE)	21	0
Group B + C (DE)	0	100

HP, hyperplastic polyp; IM, intramucosal; SM, submucosal; DE, diagnosis by the endoscopists.

The diagnostic accuracy rate of the experienced endoscopists was 100% (121/121) for neoplastic lesions.

► **Table 5** Diagnostic value of the endoscopists for SM-deep lesions.

	SM-deep lesions	Non-SM-deep lesions
Group A + B (DE)	6	97
Group C (DE)	14	4

SM, submucosal; DE, diagnosis by the endoscopists.

The diagnostic accuracy rate of the experienced endoscopists was 91.7% (111/121) for SM-deep lesions.

In the diagnoses by experienced endoscopists, all HPs were classified as type A and all neoplastic lesions were classified as type B or C (► **Table 4**). The sensitivity, specificity, PPV, NPV, and accuracy of their diagnoses of SM-deep lesions were 70.0%, 96.1%, 77.8%, 94.2%, and 92.6%, respectively (► **Table 5**).

No significant difference in the accuracies of differential diagnosis of colorectal lesions was observed between the CAD and the endoscopists' diagnosis ( $P > 0.05$ ). Moreover, no significant difference in the diagnostic accuracy of invasion depth was found between the CAD and the endoscopists' diagnosis ( $P > 0.05$ ).

## Discussion

Accurately differentiating between neoplastic and nonneoplastic lesions and estimating the invasion depth of colorectal neoplasia are essential in determining appropriate endoscopic treatment. Currently, magnifying chromoendoscopy with dye (indigo carmine/crystal violet), which has high diagnostic accuracy in differentiating colorectal lesions and in estimating the



► **Fig. 3** Lesions diagnosed differently between the endoscopists and the software for computer-aided diagnosis. **a** Lesion diagnosed as group B by the endoscopists, and diagnosed as group C by the software for computer-aided diagnosis Macroscopic Type: Is, Pathological diagnosis: Intramucosal adenocarcinoma in tubular adenoma. **b** Lesion diagnosed as group B by the endoscopists, and diagnosed as group C by the software for computer-aided diagnosis Macroscopic Type: Is + IIa, Depth of invasion: Intramucosal adenocarcinoma in tubulovillous adenoma. **c** Lesion diagnosed as group C by the endoscopists, and diagnosed as group B by the software for computer-aided diagnosis Macroscopic Type: Ip + IIc, Pathological diagnosis: Intramucosal adenocarcinoma with adenoma component. Depth of invasion: Submucosal Deep (3600  $\mu$ m). **d** Lesion diagnosed as group C by the endoscopists, and diagnosed as group B by the software for computer-aided diagnosis Macroscopic Type: Is, Depth of invasion: Intramucosal adenocarcinoma in tubular adenoma.

invasion depth of colorectal lesions, should be considered as the standard criterion [13–16]. However, chromoendoscopic observation requires much time, compared not only with white light observation but also with NBI observation [17]. M-NBI enables detailed observation of microvascular architecture and can be used for endoscopic differential diagnosis and estimation of invasion depth of colorectal lesions. However, magnifying observation of colorectal lesions has not been standardized in many Western countries. In addition, use of M-NBI in clinical practice requires knowledge and experience; thus, endoscopic differential diagnosis and estimation of invasion depth of colorectal lesions based on the findings of M-NBI has not become widespread in Western countries [18]. Therefore, a simple and automated interpretation using an objective standard of M-NBI findings would be beneficial for clinical practice.

The results of this study show that relatively high identification rates were obtained with CAD of M-NBI images for differential diagnosis and estimation of invasion depth of colorectal lesions, and that the diagnostic values were not significantly different from those of experienced endoscopists, although several lesions were diagnosed differently by the endoscopists and CAD using software (► **Fig. 3**). In addition, this software for

CAD can be further improved by strengthening the reliability of each quantitative characteristic and seeking new ones.

This study has several limitations. The number of images used in this study was limited, sample size was not calculated, and the images were obtained from a single center. In addition, determination of ROI was conducted by a single experienced endoscopist, thus precluding determination of interobserver agreement. The 2 lesions diagnosed as Group C by the CAD using software but diagnosed as Group B are presented (► **Fig. 3a**, ► **Fig. 3b**). One of them had a villous component that often presents blood pooling (► **Fig. 3b**). Considering this result and the software that uses visualized vessel findings for the diagnosis of the lesions, novel software for CAD might have limitations for diagnosis of a villous lesion.

## Conclusion

With further improvement of the software used in the current study, CAD as aided by this novel software could possibly lead to a wider use of M-NBI in the endoscopic diagnosis of colorectal lesions.

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## Competing interests

None

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