Artificial Intelligence-Based Colorectal Polyp Histology Prediction by Using Narrow-Band Image-Magnifying Colonoscopy

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Background/Aims: We have been developing artificial intelligence based polyp histology prediction (AIPHP) method to classify Narrow Band Imaging (NBI) magnifying colonoscopy images to predict the hyperplastic or neoplastic histology of polyps. Our aim was to analyze the accuracy of AIPHP and narrow-band imaging international colorectal endoscopic (NICE) classification based histology predictions and also to compare the results of the two methods.

Methods: We studied 373 colorectal polyp samples taken by polypectomy from 279 patients. The documented NBI still images were analyzed by the AIPHP method and by the NICE classification parallel. The AIPHP software was created by machine learning method. The software measures five geometrical and color features on the endoscopic image.

Results: The accuracy of AIPHP was 86.6% (323/373) in total of polyps. We compared the AIPHP accuracy results for diminutive and non-diminutive polyps (82.1% vs. 92.2%; \( p=0.0032 \)). The accuracy of the hyperplastic histology prediction was significantly better by NICE compared to AIPHP method both in the diminutive polyps (n=207) (95.2% vs. 82.1%) (\( p<0.001 \)) and also in all evaluated polyps (n=373) (97.1% vs. 86.6%) (\( p<0.001 \)).

Conclusions: Our artificial intelligence based polyp histology prediction software could predict histology with high accuracy only in the large size polyp subgroup. Clin Endosc 2022;55:113-121

Key Words: Artificial intelligence; Colorectal polyps; Histology prediction; Narrow band imaging; Narrow-band imaging international colorectal endoscopic classification

INTRODUCTION

Colonoscopy with polypectomy of early colorectal neoplastic lesions (polyps) is a proven and widely accepted method of reducing colorectal cancer mortality rates. Predicting histology prior to endoscopic colorectal polyp removal is useful especially for diminutive (1–5 mm) and small (6–10 mm) polyps. Colorectal polyp histology can be non-neoplastic (hyperplastic) or neoplastic (tubular, villous adenomas, or sessile serrated lesions [SSls]).

Non-neoplastic lesions, especially if they are diminutive hyperplastic polyps, may not require endoscopic polypectomy because of the negligible risk for developing malignancy.1-3 Evaluation of colorectal polyps using the narrow-band imaging (NBI) technique and the NBI International Colorectal Endoscopic (NICE) classification are useful to predict the histology during endoscopy.5,7 However, NBI and magnification based polyp histology prediction needs training and endoscopic experience. Moreover, the final and objective diagnosis still requires histology.

Therefore, we have been developing artificial intelligence-based polyp histology prediction (AIPHP) software to automatically evaluate the magnified NBI colonoscopy images aiming the histology prediction of polyps. We report the development of our software that can evaluate colorectal polyps using selectively recorded still colonoscopic images.
We aimed to analyze the AIPHP and NICE classification-predicted histology results and compare the histological predictive accuracy of the two methods.

MATERIALS AND METHODS

We examined 373 colorectal polyps (207 polyps, \( \leq 5 \) mm; 103 polyps, 6–9 mm; and 63 polyps, \( \geq 10 \) mm) obtained from 279 patients. Polyps were removed by traditional polypectomy or with mucosectomy or by cold-snare technique.

All endoscopic procedures and histological examinations were performed at the Petz Aladar University Teaching Hospital between October 2015 and November 2018.

Colonoscopies were performed with Olympus EXERA III CFHQ190 I (Olympus, Tokyo, Japan) high-resolution NBI colonoscope, providing 65x optical magnification.

Colorectal polyps were detected first by high definition colonoscopy then by NBI at the optical maximum magnification (65 \( \times \)). All studied polyps were photo-documented. The stored NBI photos were analyzed by the NICE classification and AIPHP parallel system (Fig. 1).

The still NBI-magnifying colorectal polyp images were taken before polyp removal by an endoscopist having > 20-year experience. The stored NBI images were classified as type I or II-III with high-confidence prediction by three experienced endoscopists using NICE. All the three endoscopists were blinded to the histology and AIPHP results. In particular polyp cases when the NICE class assessment differed among the evaluating endoscopists, the majority (two-thirds) classification was accepted as final. The NICE classification divides pit patterns and microvessels at the surface structure in NBI images into types I, II, or III.¹

This classification correlates with the most likely pathology (Figs. 2, 3). NICE I corresponds to serrated lesions, such as hyperplastic polyps and SSLs, whereas NICE II and III correspond to adenomatous polyps.

Histological examination methods

Pathological examinations were performed by a single pathologist who was blinded to both the NICE and AIPHP system diagnosis. To determine the alterations, we used the WHO classification of colorectal polyps.⁸

We used histology as the gold standard reference in statisti-
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The two-class classifications were considered: hyperplastic or neoplastic (SSLs, tubular or villous adenomas, and invasive adenocarcinomas).

The study was approved by the Regional and Hospital Research Ethics Committee (ethical approval number: 76–1–20/2015) and it was performed at the Department of Endoscopy and Gastroenterology, Petz Aladar University Teaching Hospital, Gyor, Hungary, in accordance with the Declaration of Helsinki (clinical trial registration number: NCT04425941). Patients gave informed consent for the endoscopic procedure and the pathological examination study as well as for the NICE and AIPHP analysis of the resected specimen.

AIPHP software system

The AIPHP software was developed in Python language, using OpenCV module. The AIPHP software is based on the categorization of the vascular pattern and color of the polyps.

The main steps of AIPHP software development were the following: 1) feature vector calculation, 2) training of classifier module, and 3) AIPHP classifier testing (Fig. 4). The present AIPHP version cannot automatically find the area of interest of polyp in a colonoscopy image; therefore human interaction is needed to select it. A simple image editor program (GNU Image Manipulation Program; GIMP, USA) was used for this purpose. The next step is the pre-processing that contains

**Fig. 3.** Typical endoscopic view of a NICE classification II polyp.

**Fig. 4.** Main steps of artificial intelligence-based polyp histology prediction (AIPHP) training; feature vector calculation (left) and training of sub-classifiers (right). NBI, narrow-band imaging, HP, hyperplastic polyp; PI, polyp investigation; SSA, sessile serrated adenoma; SVM, support vector machine; TVA, tubulo villous adenoma.
automatic noise reduction, glare removal, and a brightness correction step.\textsuperscript{a}

Five features were used by our AIPHP software. Feature 1 is the relative standard deviation of the intensity diagram of the pre-processed image. Features 2 and 3 are the relative area of irregular bright and dark spots based on a method that classifies every spots as either "s1", "s2", "s3", or "s4" where "s1" is for approximately circular and "s4" for irregular shapes with branches (Fig. 5). Features 4 and 5 measure the color difference of the polyp surface and the surrounding area.

Training was performed using a cross-validation scheme with 10 subsets. All trained 10 classification methods were saved and used later in the testing phase.

The average time to assess polyp histology by AIPHP was 0.5 ± 0.2 s. A trained person had to mark the polyp surface on still images because the present AIPHP version cannot automatically find the area of interest. This maneuver took an additional 10–15 s (12.2 ± 6).

Statistical analysis

Medical parameters and histological data were collected in one Excel file, which was analyzed by a self-developed Python program using the Pandas module to calculate various statistical quantities in tables. Fisher’s exact test was applied for calculating $p$-values using Scientific Python software (an open source, community developed software) and $p < 0.05$ was considered statistically significant.

Linear regression was calculated to characterize the correlation between the size of polyps and accuracy of the AIPHP and NICE methods.

RESULTS

A total of 400 colorectal polyps were detected, photo-documented (NBI still images 65× magnification) and removed for histological analysis. Because of technical failures, like mechanical or thermal damage, 27 specimens were considered not suitable, hence a total of 373 polyps were characterized by AIPHP analysis and NICE classification as well as histologically (Fig. 6). The endoscopic, histological and NICE characteristics of the lesions are presented in Table 1.

Among the polyps, 143 (38.3%) and 230 (61.7%) were hyperplastic and neoplastic (among them, 151 tubular adenomas, 70 tubulovillous adenomas, 3 SSLs and 6 invasive adenocarcinomas), respectively. Of the diminutive polyps, 128 (61.8%) and 79 (38.2%) showed hyperplastic and neoplastic histology, respectively (Table 1).

Utility of AIPHP and NICE classification

The accuracy of AIPHP was 86.6% (323/373) (sensitivity, 92.2%; specificity, 77.6%; positive predictive value [PPV], 86.9; negative predictive value [NPV], 86.0%) in all the polyps.

We compared the AIPHP accuracy results for diminutive and non-diminutive polyps (82.1% vs. 92.2%; $p = 0.0032$), which showed a significantly higher accuracy in the non-diminutive polyp group (Table 2).

Further, we evaluated the NICE classification results in predicting hyperplastic or neoplastic polyp histology in different polyp size groups. The accuracy of NICE results predicting the correct histology was 95.2% (197/207) (sensitivity, 97.5%; specificity, 93.7%; PPV, 90.6%; NPV, 98.4%) in the diminutive
The accuracy of the hyperplastic histology prediction was significantly better by NICE compared to AIPHP method both in the diminutive polyps ($n=207$, 95.2% vs. 82.1%, $p<0.001$) and also in all evaluated polyps ($n=373$, 97.1% vs. 86.6%, $p<0.001$) (Table 4).

**Fig. 6.** Polyp analysis flowchart. AIPHP, artificial intelligence-based polyp histology prediction; NBI, narrow-band imaging; NICE, NBI international colorectal endoscopic; SD, standard deviation.

**Table 1.** Characteristics of the Polyps Analyzed in the Study

|                         | 1–5 mm ($n=207$) | 6–10 mm ($n=103$) | 11–100 mm ($n=63$) | Total number ($n=373$) |
|-------------------------|-------------------|-------------------|-------------------|-----------------------|
| **Localizations in the colon** |                   |                   |                   |                       |
| Left side               | 175 (84.5%)       | 74 (71.8%)        | 54 (85.7%)        | 303 (81.2%)           |
| Right side              | 32 (15.5%)        | 29 (28.2%)        | 9 (14.3%)         | 70 (18.8%)            |
| **NICE classification** |                   |                   |                   |                       |
| NICE I                  | 122 (58.9%)       | 13 (12.6%)        | 1 (1.6%)          | 136 (36.5%)           |
| NICE II-III             | 85 (41.1%)        | 90 (87.4%)        | 62 (98.4%)        | 237 (63.5%)           |
| **Shape of polyps**     |                   |                   |                   |                       |
| Pedunculated            | 1 (0.5%)          | 6 (5.8%)          | 15 (23.8%)        | 22 (5.9%)             |
| Semipedunculated        | 3 (1.4%)          | 18 (17.5%)        | 19 (30.2%)        | 40 (10.7%)            |
| Sessile                 | 203 (98.1%)       | 79 (76.7%)        | 29 (46.0%)        | 311 (83.4%)           |
| **Histology**           |                   |                   |                   |                       |
| Hyperplastic polyp      | 128 (61.8%)       | 14 (13.6%)        | 1 (1.6%)          | 143 (38.3%)           |
| Neoplastic lesion       | 79 (38.2%)        | 89 (86.4%)        | 62 (98.4%)        | 230 (61.7%)           |

NICE, narrow-band imaging international colorectal endoscopic.
Table 2. Comparison of Artificial Intelligence-Based Polyp Histology Prediction Results with Histology

| Polyp size (mm) | AIPHP vs. Histology | NICE vs. Histology |
|----------------|---------------------|---------------------|
|                | Neoplastic | Hyperplastic | Sens % | Spec % | PPV % | NPV % | Acc % |
| 1–5 (n=207)    | 70/79     | 100/128      | 88.6  | 78.1  | 71.4  | 91.7  | 82.1* |
| >5 (n=166)     | 142/151  | 11/15        | 94.0  | 73.3  | 97.3  | 55    | 92.2* |
| All polyps (n=373) | 212/230  | 111/143      | 92.2  | 77.6  | 86.9  | 86    | 86.6  |

*p<0.0032

Acc, accuracy; AIPHP, artificial intelligence-based polyp histology prediction; NPV, negative predictive value; PPV, positive predictive value; Sens, sensitivity; Spec, specificity.

Table 3. Comparison of NICE Classification Results with Histology

| Polyp size (mm) | NICE | I | II-III |
|----------------|------|---|--------|
|                | Sens % | Spec % | PPV % | NPV % | Acc % |
| 1–5 (n=207)    | 122   | 85  | 97.5  | 93.7  | 90.6  | 98.4  | 95.2  |
| 6–100 (n=166)  | 14    | 152 | 100   | 93.3  | 99.3  | 100   | 99.4  |
| 1–100 (n=373)  | 136   | 237 | 99.1  | 93.7  | 96.2  | 98.5  | 97.1  |

Acc, accuracy; NICE, narrow-band imaging international colorectal endoscopic; NPV, negative predictive value; PPV, positive predictive value; Sens, sensitivity; Spec, specificity.

Table 4. Hyperplastic Polyp Histology Predicting Data by NICE Classification and Artificial Intelligence-Based Polyp Histology Prediction

|                | Sensitivity (%) | Specificity (%) | Accuracy (%) |
|----------------|-----------------|-----------------|--------------|
| All evaluated polyps (n=373) | NICE | 99.1 | 93.7 | 97.1* |
|                | AIPHP | 92.2 | 77.6 | 86.6* |
| Diminutive 1–5 mm polyps (n=207) | NICE | 97.5 | 93.7 | 95.2** |
|                | AIPHP | 88.6 | 78.1 | 82.1** |

*p<0.001
**p<0.0001

AIPHP, artificial intelligence-based polyp histology prediction; NICE, narrow-band imaging international colorectal endoscopic.

Neoplastic and hyperplastic histology were correctly predicted by AIPHP in 92.2% (212/230) and 77.6% (111/143) of the specimens, respectively (p<0.0001).

Polyp sizes had influenced both the AIPHP/histology and NICE/histology agreement results. A detailed calculation was performed to study the connection between the polyp sizes and agreement accuracy values using five size groups (Fig. 7).

The accuracy of NICE prediction was close to 100% in all size groups whereas AIPHP/histology accuracy showed an increasing tendency with the polyp sizes. Linear regression correlation coefficient results indicate that AIPHP method was significantly more accurate in the bigger polyps than in the smaller polyps.

DISCUSSION

Colorectal polyp diagnostic methods have progressed very dynamically. The current dilemma is that diminutive polyps (≤5 mm) mostly show a hyperplastic histology which has a non-neoplastic behavior, and thus the rationale of polyp removal is under discussion.

High-definition endoscopy, including NBI technology, is able to predict the histology of colorectal polyps with high accuracy. However, this accuracy is significantly influenced by subjective elements, such as endoscopist technical skills and experience.6,10

Consequently, the number of artificial intelligence polyp classification endoscopic methods is rapidly increasing.11,17
The ideal aim of any AIPHP is to provide objective “semi-histological” diagnosis and ensure high diagnostic accuracy for non-expert endoscopists.

In our study, the still images were taken by an endoscopist experienced with the NBI-based histology prediction criteria and able to choose good-quality images with optimal position. The photo-documented and stored polyp images were collected, and the regions of interest were selected manually.

We previously reported that our pilot AIPHP program can distinguish between images of hyperplastic or neoplastic lesions in NBI-magnifying colonoscopy images. The Preservation and Incorporation of Valuable Endoscopic Innovation (PIVI) committee of the American Society for Gastrointestinal Endoscopy published a statement on establishing endoscopic diagnostic techniques. Several NBI-based and magnifying endoscopy methods like NICE and Hiroshima classifications are consistent with the PIVI recommendations of a ≥ 90% NPV and accuracy criteria.

Our present results show that the NICE classification fulfilled the PIVI criteria but the AIPHP program could only partly achieve PIVI recommendations among diminutive polyps (NPV, 91.7%; accuracy, 82.1%). However, in the case of non-diminutive polyps, the values fulfilled the PIVI criteria (sensitivity: 94.0%, accuracy: 92.2%). These findings indicate that the present AIPHP cannot offer the automatic computerized endoscopy-based histological diagnosis especially in those polyp size groups where the AI diagnosis would be most beneficial regarding cost-benefit considerations.

The question is why our AIPHP accuracy results, especially in the diminutive polyp subgroup, were worse than in the bigger polyp subgroups. One possible reason is that the diminutive polyps are mostly hyperplastic and are covered with mucous surfaces. The adherent mucous disturbs the ability to analyze the vascular pattern by AIPHP procedure. The mucous cover and the whitish shape is one reason why we got less precise histologically predictive results by AIPHP program compared to NICE classification results. Therefore, hyperplastic polyps can be differentiated better from adenomas using the NICE classification. Mucous surface and whitish characters hinder histological recognition by AIPHP program in which the polyp surface vascularity and the color differences are among the main features. In conclusion, we plan to add the mucus surface as a new feature for the next-generation AIPHP programs.

To the best of our knowledge, this is one of the first studies which analyzed the AI diagnostic results in different polyp sizes. We found that the AIPHP method is significantly more accurate in the bigger size polyps than in smaller ones. Polyp sizes had significantly influenced AIPHP accuracy but not the NICE/histology agreement results. This study is also pioneering in comparing the NICE classification with a special artificial intelligence-based polyp image analyzer. Both hyperplastic and SSLs correspond to NICE I, therefore it is almost impossible to distinguish the two by NICE classification. This problem limits the accuracy of NICE classification regarding the differentiation of non-neoplastic polyps and the neoplastic SSLs. In our study, the accuracy of NICE evaluation is high, which is partly due to the small number (n = 3) of SSLs analyzed in the study. The accuracy of NICE likely would decrease as the ratio of SSLs increases.

We comparatively analyzed our AIPHP results with those three studies in which software-based automatic polyp histol-
ogy analysis provided higher sensitivity and accuracy results than our outcomes. Takemura et al. and by Komiani et al. studied low numbers of hyperplastic polyps (47 and 45, respectively) which are notably less than the 143 hyperplastic polyps in our study. The critical point of the artificial intelligence (AI) analysis is to correctly identify the diminutive non-neoplastic lesions. To fulfill this requirement, an eligible number of diminutive hyperplastic lesions should be analyzed. In the study of Gross et al., 135 non-neoplastic diminutive polyps were analyzed by a computer with 94.3% NPV. These results are slightly superior to our findings showing a 91.7% NPV by AIPHP in 128 hyperplastic lesions.

There are several limitations of our study. Trainings are necessary both for NICE evaluation and production of high-quality stored polyp images. Diminutive polyps are typically hyperplastic with a mucous covering, interfering both photo-documentation quality and correct AIPHP recognition. The “one polyp, one picture” method for analysis was suitable for NICE evaluation but did not complete the correct AI software analysis requirements in diminutive and small polyps. The present AIPHP version cannot automatically find the area of interest on the polyp surface, hence additional human interaction is needed. We found that 10–15 s is sufficient for a trained person to mark the polyp surface on a still image. It could also be acceptable in routine endoscopy, provided that the selection marker could be integrated in the software. Such an integrated software is under development.

Further problems that may limit the generalizability of our findings are that there are many less experienced endoscopists in the real-world practice who may not take good-quality images on particular areas of interest. In addition, magnifying endoscopy is not widely used in general endoscopy.

In conclusion, our artificial intelligence system manipulating still images of colorectal polyps by NBI-magnifying colonoscopy could predict histology with high accuracy only for non-diminutive (>5 mm) polyps. The AIPHP accuracy significantly increased with larger polyp sizes. Similar to other studies, NICE classification results showed high sensitivity and accuracy in all polyp size groups.

Because our present AIPHP cannot substitute the subjective endoscopic virtual histology evaluations, especially in diminutive polyps, further development with automatic area-of-interest detection and parallel analysis of multiple images of the same polyp could assist automatic polyp histology predictions, even in smaller colorectal polyps.

**Conclusions**

Further problems that may limit the generalizability of our findings are that there are many less experienced endoscopists in the real-world practice who may not take good-quality images on particular areas of interest. In addition, magnifying endoscopy is not widely used in general endoscopy. Therefore, the development of an integrated software system is needed to enable automatic recognition of diminutive colorectal polyps. Further advancements in AI-based endoscopic virtual histology analysis could improve the accuracy and efficiency of polyp classification, allowing for more precise and timely diagnosis and management.

**Conflict of interest**

The authors have no potential conflicts of interest.

**Funding**

This study was supported by the GINOP-2.3.4-15-2016-00003 grant.

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**REFERENCES**

1. McGill SK, Evangelou E, Ioannidis JPA, Soetikno RM, Kaltenbach T. Narrow band imaging to differentiate neoplastic and non-neoplastic colorectal polyps in real time: a meta-analysis of diagnostic operating characteristics. Gut 2013;62:1704-1713.
2. Ignjatovic A, East JE, Suzuki N, Vance M, Guenther T, Saunders BP. Optical diagnosis of small colorectal polyps at routine colonoscopy (Detect InSpect ChAracterise Resect and Discard; DISCARD trial): a prospective cohort study. Lancet Oncol 2009;10:1171-1178.
3. Ignjatovic A, Thomas-Gibson S, East JE, et al. Development and validation of a training module on the use of narrow-band imaging in differentiation of small adenomas from hyperplastic colorectal polyps. Gastrointest Endosc 2011;73:128-133.
4. Hewett DG, Kaltenbach T, Sano Y, et al. Validation of a simple classification system for endoscopic diagnosis of small colorectal polyps using narrow-band imaging. Gastroenterology 2012;143:599-607.e1.
5. Hewett DG, Huffman ME, Rex DK. Leasing distal colorectal hyperplastic polyps in place can be achieved with high accuracy by using narrow-band imaging: an observational study. Gastrointest Endosc 2012;76:374-380.
6. Rastogi A, Rao DS, Gupta N, et al. Impact of a computer-based teaching module on characterization of diminutive colon polyps by using narrow-band imaging by non-experts in academic and community practice: a video-based study. Gastrointest Endosc 2014;79:390-398.
7. van den Broek FJC, Reitsma JB, Curvers WL, Fockens P, Dekker E. Systematic review of narrow-band imaging for the detection and differentiation of neoplastic and nonneoplastic lesions in the colon (with videos). Gastrointest Endosc 2009;69:124-135.
8. Ahadi M, Sokolova A, Brown I, Chou A, Gill AJ. The 2019 World health organization classification of appendiceal, colorectal and anal canal tumours: an update and critical assessment. Pathology 2021;53:454-461.
9. Horváth A, Spindler S, Szalay M, Rácz I. Preprocessing endoscopic images of colorectal polyps. Acta Technica Jaurinensis 2016;9:65-82.
10. Neumann H, Vieth M, Fry LC, et al. Learning curve of virtual chromoendoscopy for the prediction of hyperplastic and adenomatous colorectal lesions: a prospective 2-center study. Gastrointest Endosc 2013;78:115-120.
11. Tischendorf J JW, Gross S, Winograd R, et al. Computer-aided classification of colorectal polyps based on vascular patterns: a pilot study. Endoscopy 2010;42:203-207.
12. Gross S, Trautwein C, Behrens A, et al. Computer-based classification of small colorectal polyps by using narrow-band imaging with optical magnification. Gastrointest Endosc 2011;74:1354-1359.
13. Takemura Y, Yoshida S, Tanaka S, et al. Quantitative analysis and development of a computer-aided system for identification of regular pit patterns of colorectal lesions. Gastrointest Endosc 2010;72:1047-1051.
14. Alagappan M, Brown J RG, Mori Y, Berzin TM. Artificial intelligence in gastrointestinal endoscopy: the future is almost here. World J Gastrointest Endosc 2018;10:239-249.
15. Hassan C, Bisschops R, Bhandari P, et al. Predictive rules for optical diagnosis of <10-mm colorectal polyps based on a dedicated software. Endoscopy 2020;52:52-60.
16. Le Berre C, Sandborn WJ, Aridhi S, et al. Application of artificial intelligence to gastroenterology and hepatology. Gastroenterology 2020;158:76-94.e2.
17. Hassan C, Spadaccini M, Iannone A, et al. Performance of artificial intelligence in colonoscopy for adenoma and polyp detection: a systematic review and meta-analysis. Gastrointest Endosc 2021;93:77-85.e6.
18. Rácz I, Horváth A, Szalai M, et al. Sa1556 digital image processing software for predicting the histology of small colorectal polyps by using narrow-band imaging magnifying colonoscopy. Gastrointestinal Endoscopy 2015;81:AB259.
19. Rex DK, Kahi C, O’Brien M, et al. The american society for gastrointestinal endoscopy PIVI (preservation and incorporation of valuable endoscopic innovations) on real-time endoscopic assessment of the histology of diminutive colorectal polyps. Gastrointest Endosc 2011;73:419-422.
20. ASGE Technology Committee, Abu Dayyeh BK, Thosani N, et al. ASGE technology committee systematic review and meta-analysis assessing the ASGE PIVI thresholds for adopting real-time endoscopic assessment of the histology of diminutive colorectal polyps. Gastrointest Endosc 2015;81:502.e1-502.e16.
21. Cohen J, Bosworth BP, Chak A, et al. Preservation and incorporation of valuable endoscopic innovations (PIVI) on the use of endoscopy simulators for training and assessing skill. Gastrointest Endosc 2012;76:471-475.
22. Tanaka S, Kaltenbach T, Chayama K, Soetikno R. High-magnification colonoscopy (with videos). Gastrointest Endosc 2006;64:604-613.
23. Kanao H, Tanaka S, Oka S, Hirata M, Yoshida S, Chayama K. Narrow-band imaging magnification predicts the histology and invasion depth of colorectal tumors. Gastrointest Endosc 2009;69:631-636.
24. Tanaka S, Sano Y. Aim to unify the narrow band imaging (NBI) magnifying classification for colorectal tumors: current status in Japan from a summary of the consensus symposium in the 79th annual meeting of the Japan gastroenterological endoscopy society. Dig Endosc 2011;23(Suppl 1):131-139.
25. Oba S, Tanaka S, Sano Y, Oka S, Chayama K. Current status of narrow-band imaging magnifying colonoscopy for colorectal neoplasia in Japan. Digestion 2011;83:167-172.
26. Takemura Y, Yoshida S, Tanaka S, et al. Computer-aided system for predicting the histology of colorectal tumors by using narrow-band imaging magnifying colonoscopy (with video). Gastrointest Endosc 2012;75:179-185.
27. Kominami Y, Yoshida S, Tanaka S, et al. Computer-aided diagnosis of colorectal polyp histology by using a real-time image recognition system and narrow-band imaging magnifying colonoscopy. Gastrointest Endosc 2016;83:643-649.