Research Article

Value of Magnifying Chromoendoscopy and Magnifying Optical Enhancement Technology in Classifying Colorectal Polyps: A Prospective Controlled Study

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Background and Aims. Magnifying chromoendoscopy (ME-CE) through the observation of pit patterns is a productive way to distinguish between neoplastic and nonneoplastic polyps. Magnifying optical enhancement technology (ME-OE) is an emerging virtual chromoendoscopy imaging technology and appeared to be a promising approach. However, this information is currently not available. This study is aimed at comparing the differential diagnostic value of ME-CE and OE for neoplastic and nonneoplastic polyps. Patients and Methods. Consecutive patients undergoing colonoscopy were randomized (1:1) into examination by ME-OE or ME-CE. Histopathological findings were utilized as the reference standard. Accuracy, sensitivity, specificity, and positive and negative predictive values of two endoscopy methods were compared using ME-OE (were classified according to the JNET classification) and ME-CE (were classified according to the Kudo pit pattern classification), respectively, and the time to predict the histological polyp type was compared. And the agreements between the pathological and clinical diagnosis by ME-OE or ME-CE were analyzed. Results. A total of 365 polyps were found in the 220 patients included (ME-OE: 185; ME-CE: 180.202 had nonneoplastic polyps, 163 had neoplastic polyps). The diagnostic accuracy of ME-OE was higher than that of ME-CE (93% vs. 92%, p > 0.05). The average diagnosis time was lower in ME-OE than ME-CE (83 ± 26.4 s vs. 194 ± 17.7 s, p < 0.001). The agreements between the pathological and clinical diagnosis were at least substantial in both groups. Conclusion. ME-OE was superlative to ME-CE in predicting the histology of polyps. OE devoted classification would possibly similarly enhance the endoscopist performance. The trial is registered with ChiCTR2000032075.
1. Introduction

In 2018, an additional 1.8 million new cases of colorectal cancer were diagnosed, with 881,000 deaths. Colorectal cancer accounts for one in 10 cancer cases and deaths, the 3rd incidence and the 2nd mortality [1, 2]. Although the incidence in China is lower than the world average, the number of new cases and deaths in China is the highest in the world. The adenoma-carcinoma series is the classic colorectal cancer (CRC) development paradigm, in which CRC starts as an adenoma. The endoscopic resection that follows will stop the condition from spreading and may also be a solution for intramucosal adenocarcinoma [3]. However, white light (WLE) colonoscopy alone is insufficient to distinguish neoplastic from nonneoplastic polyps, possibly resulting in removing a significant number of lesions that were not necessary [4]. Multiple endoscopic modalities have been recorded to be beneficial for colorectal polyp assessment [5].

Furthermore, since diminutive polyps account for the majority of polyps found during the colonoscopy, the ability to predict polyp histology in real time is clinically significant [6]. The JNET (The Japan NBI Expert Team) classification consists of four categories, types I, 2A, 2B, and 3, based on vessel and surface pattern findings. The morphological appearance of each JNET type is then correlated with the histology, from benign hyperplastic polyps to advanced carcinomas [7].

Since the 1970s, Japanese endoscopists have used chromoendoscopy, and the Kudo proposed pit pattern classification has been generally adopted. Using the Kudo pit pattern classification system, chromoendoscopy with or without magnification has been used to discern neoplastic from nonneoplastic polyps [8]. The optical enhancement (OE, Pentax Medical, Tokyo, Japan) technology was developed as a new system for visualizing the morphology of mucosal surface patterns. The early i-Scan system is utilized as it uses white light as a source of brightening. However, it does not fully meet current needs. OE is a novel technique of electronic chromoendoscopy. The innovated optical filters may achieve higher overall transmittance by connecting the peaks of the hemoglobin absorption spectrum (415 nm, 540 nm, and 570 nm), generating a continuous wavelength spectrum. There are two modes with different OE filters (mode 1 and mode 2) [9]. At present, there are only a few studies in the esophagus or stomach for OE [10–26]. It is appropriate to choose mode 1 because of the similar principle to NBI (narrow band imaging) [10–26]. Compared with NBI, there are only a few studies on classifying colorectal polyps currently available. Therefore, in this study, we evaluated and compared the detection efficacies of ME-OE and ME-CE for classifying colorectal polyps.

2. Method

This was a prospective, randomized, and single-center study conducted at a teaching hospital in Zhengzhou, China. All participating patients provided written informed consent. The Ethics Committee approved Zhengzhou Central Hospi-
Figure 1: Endoscopic images in the magnifying mode: (a) nonneoplastic polyps (JNET type 1); (b) neoplastic polyps (JNET type 2A); (c) nonneoplastic polyps (pit pattern types I); (d) neoplastic polyps (pit pattern type IV).

Figure 2: Flow chart of the examinations: 519 patients enrolled, 220 patients eligible (299 patients excluded: no polyps: \(n = 111\), coagulation disorders: \(n = 92\), inadequate bowel preparation: \(n = 43\), melanosis coli: \(n = 20\), previous colon resection: \(n = 24\), polyposis syndrome: \(n = 5\), and incomplete colonoscopy: \(n = 4\)).
the surface of polyps using a spray tube, and excess solution was aspirated after the spraying, making a magnified observation, and the prediction was reported to the assistant. The assistant recorded the time from inserting the spray tube to reaching the diagnosis and withdrawing the spray tube. During the whole process, any communication forms between the endoscopists were not allowed. The location was estimated by the anatomic landmarks. The size was evaluated by comparison with the span of open biopsy forceps. Polyp mated by the anatomic landmarks. The size was evaluated by the kappa test. The larger the kappa, the higher the agreement.

2.3. Pathological Polyp Evaluation. The tissue specimens collected during endoscopies were placed into formalin solution for 24 hours, subjected to conventional dehydration, paraffin embedding, sectioning, and then staining using the hematoxylin-eosin (H/E) staining method. Two experienced pathologists reviewed the slides independently and reached histological conclusions without knowing the endoscopic findings. If the results of the two pathologists were notably different, a third pathologist was consulted. The diagnostic histopathological criteria were based on the Vienna classification [10].

2.4. Outcome Measures. With histopathological evaluation as a reference standard, we evaluated the accuracy, sensitivity, specificity, and positive and negative predictive values of ME-OE and ME-CE endoscopy for polyp histology prediction, as primary outcomes.

The secondary outcome measures included comparing the time needed to predict polyp histology and the agreements between the pathological and clinical diagnosis by ME-OE or ME-CE.

2.5. Sample Size Estimation. The overall diagnostic accuracy in differentiating neoplastic from nonneoplastic lesions in their series was 80% [11]. We have considered clinically relevant an absolute difference of 15% [7]. All statistical tests were two-sided with a significance level of 5%, and statistical power, 1-β, is set to be 80%. With 10% expected dropouts, the total sample size was 220, who prudentially assume a detection of only one polyp. Combined with my endoscopic center, the detection rate of polyps is about 35%. We planned to enroll 628 patients.

2.6. Statistical Analysis. Statistical analysis was performed using SPSS 26.00 statistical software. Frequencies with percentages were used to represent qualitative variables, whereas means and standard deviations were used to describe quantitative variables. Comparisons of qualitative variables were conducted using Fisher’s exact probability test or chi-squared test, while for comparisons of continuous variables, we employed a t-test. Consistency analysis was evaluated by the kappa test. The larger the kappa, the higher the agreement.

### Table 1: Clinical characteristics of the subject subjects.

|               | ME-OE | ME-CE | Overall | p value |
|---------------|-------|-------|---------|---------|
| Number        | 110   | 110   | 220     | NA      |
| Gender (M/F)  | 56/54 | 53/57 | 109/111 | 0.686   |
| Age (mean ± SD) (y) | 48.8 ± 14.1 | 51.7 ± 14.3 | 50.3 ± 14.2 | 0.121   |
| Indication    |       |       |         | 0.154   |
| Screening     | 21    | 14    | 35      |         |
| Symptoms      | 43    | 34    | 77      |         |
| Surveillance  | 23    | 35    | 58      |         |
| Examination   | 23    | 27    | 50      |         |
| Bowel cleansing (BBPS) | 6/7 | 7/17 | 6/32 | 13/49 | 0.08 |

ME-CE: magnifying chromoendoscopy; ME-OE: magnifying optical enhancement technology; BBPS: Boston Bowel Preparation Scale; NA: not applicable; SD: standard deviation.

### 3. Results

#### 3.1. Patients and Polyps. From August 2019 to May 2020, we screened 519 patients, and after, 229 patients were excluded (Figure 2). A total of 220 patients were included (ME-OE: 185 polyps in 110, ME-CE: 180 polyps in 110). There were similarly no differences in age, sex, bowel preparation, or indications for colonoscopy between the two groups' (Table 1) accessibility, size, morphology, anatomical location, and pathologic diagnosis (Table 2).

#### 3.2. Primary End-Point. Our main targets here are to compare the accuracy of polyp diagnosis between groups.

The overall accuracies were 93% vs. 92% for ME-OE and ME-CE, respectively, p > 0.05. The accuracy of ME-OE in stage 1 and 2 was 90% vs. 96%, p > 0.05. The accuracy of ME-CE in stages 1 and 2 was 90% vs. 93%, p > 0.05 (Table 3). In stage 1, the overall accuracies were 90% and 90% for ME-OE and ME-CE, p > 0.05. In stage 2, the overall accuracies were 96% for ME-OE and 93% for ME-CE, p > 0.05 (Table 4).

#### 3.3. Secondary End-Points. In addition to our objectives of primary interest, we examined the following secondary purposes. The average diagnosis time between the two groups and various stages in the same groups was compared: 83 s ± 26.4 s for ME-OE and 194 s ± 17.7 s for ME-CE (p < 0.001), 96 s ± 26.8 s and 70 s ± 19.0 s (p < 0.001) in two stages of ME-OE, and 20.5 s ± 13.9 s and 183 s ± 14.0 s (p < 0.001) in two stages of ME-CE (Table 3).

The average diagnosis time between the various groups in the same stages was compared: 96 s ± 26.8 s for ME-OE and 205 s ± 13.9 s for ME-CE (p < 0.001) in stage 1 and 70 s ± 19.0 s for ME-OE and 183 s ± 14.0 s for ME-CE (p < 0.001) in stage 2 (Table 4).

The agreements between the pathological and clinical diagnosis between the two groups and various stages in the same groups were compared: both almost perfect for ME-OE (κ = 0.859) and ME-CE (κ = 0.827), both almost perfect.
Table 2: Descriptive per-polyp analysis.

|                  | ME-OE (%) | ME-CE (%) | Overall (%) | p value |
|------------------|-----------|-----------|-------------|---------|
| Number           | 185       | 180       | 365         | NA      |
| Size (mm)        |           |           |             |         |
| ≤5               | 104 (56.2)| 119 (66.1)| 223 (61.1)  | 0.157   |
| 5-10             | 76 (41.1)| 58 (32.2)| 134 (36.7)  |         |
| ≥10              | 5 (2.7)  | 3 (1.7)  | 8 (2.2)     |         |
| Location         |           |           |             | 0.389   |
| Right side of colon | 73 (39.5)| 64 (35.6)| 137 (37.5)  |         |
| Transverse colon | 35 (18.9)| 26 (14.4)| 61 (16.7)   |         |
| Descending colon | 40 (21.6)| 44 (24.4)| 84 (23.1)   |         |
| Sigmoid colon/rectum | 37 (20.0)| 46 (25.6)| 83 (22.7)   |         |
| Morphology       |           |           |             | 0.064   |
| Ip               | 23 (12.5)| 33 (18.3)| 56 (15.3)   |         |
| Is               | 62 (33.5)| 73 (40.6)| 135 (37.0)  |         |
| Ila              | 77 (41.6)| 61 (33.9)| 138 (37.8)  |         |
| IIb              | 23 (12.4)| 13 (7.2) | 36 (9.9)    |         |
| Histology        |           |           |             | 0.120   |
| Nonneoplastic    | 95 (51.4)| 107 (59.4)| 202 (55.3) |         |
| Size (mean ± SD) (mm) | 3.1 ± 1.1 | 3.3 ± 1.2 | 3.2 ± 1.1 |         |
| Neoplastic       | 90 (48.6)| 73 (40.6)| 163 (44.7)  |         |
| Size (mean ± SD) (mm) | 5.9 ± 1.7 | 5.8 ± 1.4 | 5.8 ± 1.6 |         |
| Time (mean ± SD) (s) | 83 ± 26.4 | 194 ± 17.7 | 138 ± 60.2 | 0.001   |

ME-CE: magnifying chromoendoscopy; ME-OE: magnifying optical enhancement technology; NA: not applicable; SD: standard deviation.

Table 3: Diagnostic accuracy and time in two methods.

| Overall | ME-OE n = 185 | ME-CE n = 180 | Stage 1 n = 92 | Stage 2 n = 93 | Stage 1 n = 90 | Stage 2 n = 90 |
|---------|---------------|---------------|----------------|----------------|----------------|----------------|
| Sensitivity (%) | 91 | 85 | 96 | 90 | 87 | 84 |
| Specificity (%)  | 95 | 96 | 85 | 1 | 93 | 98 |
| PPV (%)          | 94 | 94 | 86 | 1 | 93 | 96 |
| NPV (%)          | 92 | 90 | 95 | 93 | 88 | 92 |
| Accuracy (%)     | 93 | 92 | 90 | 96 | 90 | 93 |
| p value          | 0.639 | 0.145 | 0.0145 | 0.418 | 0.800 | 0.848 |
| K                | 0.859 | 0.827 | 0.804 | 0.912 | 0.800 | 0.848 |
| Time (mean ± SD) (s) | 83 ± 26.4 | 194 ± 17.7 | 96 ± 26.8 | 70 ± 19.0 | 205 ± 13.9 | 183 ± 14.0 |
| p value          | 0.001 | 0.001 | 0.001 | 0.001 | 0.001 | 0.001 |

PPV: positive predictive value; NPV: negative predictive value; ME-CE: magnifying chromoendoscopy; ME-OE: magnifying optical enhancement technology; SD: standard deviation.

(k = 0.804 and 0.912) in 2 stages of ME-OE, and substantial (k = 0.800) for stage 1 and nearly perfect (k = 0.848) for stage 2 in ME-CE (Table 3).

4. Discussion

Three aspects were evaluated in our prospective randomized study. First, the diagnostic accuracy of ME-OE was higher than that of ME-CE. Second, the average diagnosis time was lower in ME-OE than in ME-CE. Third, the agreements between the pathological and clinical diagnosis at least were substantial in both groups.

In older studies, the accuracy of polyps categorized as nonneoplastic or neoplastic lesions by conventional endoscopy was shown to be 80% by the Kudo classification [16]. In the subsequent research, Calderwood et al. showed that ME-CE could distinguish between neoplastic and nonneoplastic polyps with an accuracy of 80.1% [15]; Hirata et al. reported that differentiation between neoplastic and non-neoplastic lesions was possible with a 92% sensitivity and a
Table 4: Diagnostic accuracy and time in two stages.

|                | Overall  | Stage 1  | Stage 2  |
|----------------|----------|----------|----------|
|                | ME-OE  | ME-CE    | ME-OE  | ME-CE    | ME-OE  | ME-CE    |
|                | n = 365 | n = 180  | n = 182 | n = 90   | n = 93  | n = 90   |
| Sensitivity (%)| 91      | 85       | 96      | 87       | 0.90    | 84       |
| Sensitivity (%)| 95      | 96       | 85      | 93       | 1       | 98       |
| PPV (%)        | 94      | 94       | 86      | 93       | 1       | 96       |
| NPV (%)        | 92      | 90       | 95      | 88       | 93      | 92       |
| Accuracy (%)   | 93      | 92       | 90      | 90       | 96      | 93       |
| $p$ value      | 0.639   | 0.961    | 0.705   |
| $K$            | 0.859   | 0.827    | 0.804   | 0.800    | 0.912   | 0.848    |
| Time (mean ± SD) (s) | 83 ± 26.4 | 194 ± 17.7 | 96 ± 26.8 | 205 ± 13.9 | 70 ± 19.0 | 183 ± 14.0 |

$PPV$: positive predictive value; $NPV$: negative predictive value; ME-CE: magnifying chromoendoscopy; ME-OE: magnifying optical enhancement technology; SD: standard deviation.

73.3% specificity [17]. Overall, the diagnostic accuracy in differentiating neoplastic from nonneoplastic lesions was 88.4%. In meta-analysis [18], the Kudo classification has great significance in the identification of colorectal neoplasm. Tung et al. showed that ME-CE is a reliable tool to predict and differentiate between neoplastic and nonneoplastic polyps [20]. Optical enhancement (OE) was recently developed by PENTAX, and few studies have been conducted in this field. However, several studies suggest that the JNET classification can provide immediate histological diagnosis and more reliable estimation of the depth of invasion by magnifying endoscopy [19–23].

Compared to the accuracy of the two groups, the diagnostic accuracy of ME-OE was significantly higher than that of ME-CE (93% vs. 92%, $p > 0.05$). In stage 1, the overall accuracies were 90% and 90% for ME-OE and ME-CE, $p > 0.05$. In stage 2, the overall accuracies were 96% for ME-OE and 93% for ME-CE, $p > 0.05$. Whether from the overall accuracy or the same group of different stages, the same stage of the other groups’ accuracy is more than 90% accurate. The agreements between the pathological and clinical diagnosis are healthy at all times. It can be seen that both groups have effective methods to diagnose the nature of polyps. This point needs attention. The performance of ME-OE in classifying colorectal polyps is much better than ME-CE at all times. Especially in OE stage two, the specificity was 100%, and there were no cases with misdiagnosis. While the ME-CE is also excellent, the ME-OE seems to be the better choice.

In terms of diagnosis time, ME-OE was also faster and convenient than ME-AAC (83 s ± 26.4 s vs. 194 s ± 17.7 s). This superiority continues until the time point before the last. This may be related to the operation process of pigment endoscopy, which requires not only delivering the spray tube to the intestinal cavity and fixing the spray tube to a specific position to spray accurately but also removing the excess solution. Besides, there are differences between different stages within the same group. In both groups, the second stage was faster than the first. This might be due to a lack of experience in the early. In a study of endoscopists with different experiences, after 20 minutes of NBI teaching [17], 37 doctors’ accuracy in distinguishing adenomas from proliferative polyps rose from 47.6% to 90.8%, suggesting that short education can improve doctors’ judgment on the nature of polyps. In our study, no matter in the OE group or the CE group, after the first stage, the accuracy of judging the nature of polyps by endoscopists in the second stage has been significantly improved, which seems to show that practical learning plays an essential role in enhancing the ability of doctors. According to the European Society of Gastrointestinal Endoscopy (ESGE) [17, 18], it is significant for optical diagnostics by self-directed learning or practical courses. It is feasible that virtual chromoendoscopy and dye-based chromoendoscopy can be used for optical diagnosis of limited to diminutive polyps ($\leq 5\,mm$) without the need for pathologic confirmation, but only if adequately photodocumented which would be required to support endoscopists’ claims of adenoma detection, and also emphasized experienced endoscopists who are adequately trained.

This study also had some limitations. Namely, this was a single-center study with a small sample size, and thus, further in-depth, large-scale, and multicenter studies are required in the future. Besides, no specific classifying colorectal polyps for ME-OE are available thus far, and we hope that a ME-OE diagnostic standard can be established shortly. The examination was performed by experienced endoscopists, which may lead to selection bias. As the limitation of the experimental specimens, we did not collect all the polyp types, which may have an individual impact on the conclusion.

Overall, the study adds to our understanding of classifying colorectal polyps by ME-OE. And the ability of ME-OE was significantly better than that of ME-CE. The method not only has high accuracy but also is simple to perform with ME-CE and saves time. Furthermore, practical learning is necessary to raise diagnostic accuracy. In the future, we should further verify the ability to use OE to identify various polyp types based on JNET classification.
Data Availability
No additional data are available.

Additional Points

Clinical Trial Registration Statement. This study was registered at chictr.org.cn. The registration identification number is ChiCTR2000032075.

Ethical Approval
All procedures performed in studies involving human participants were in accordance with the ethical standards of the institutional and national research committee and with the 1964 Helsinki declaration and its later amendments or comparable ethical standards. The research protocol was approved by the Ethics Committee of Zhengzhou Central Hospital Affiliated to Zhengzhou University (202032).

Consent
All participating patients provided written informed consent.

Conflicts of Interest
All authors declare no potential conflicting interests related to this paper.

Authors’ Contributions
Song YH, Xu LD, Qu YL, Xu RX, and Xing MX were responsible for the study conception and design, data analysis and interpretation, and manuscript drafting; Li KK, Xiao XG, Zhang Y, Xiao YJ, Jia BH, Ma YJ, and Wu HL critically revised the article for important intellectual content; all the authors reviewed and approved the final version to be published. Ying-Jie Ma, Bao-Hui Jia, and Hui-Li Wu contributed equally to this work.

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