The feasibility and safety of disposable endoscope vs. conventional endoscope for upper gastrointestinal tract examination: a multicenter, randomized, parallel, non-inferiority trial

Die Durchführbarkeit und Sicherheit von Einweg-Endoskopen bei der Untersuchung des oberen Gastrointestinaltrakts im Vergleich zu konventionellen Endoskopen: Eine multizentrische, randomisierte, parallele, Nichtunterlegenheits-Studie

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ABSTRACT

Background A disposable upper gastrointestinal endoscope can effectively decrease infectious outbreaks associated with endoscope reuse. In the present study, we aimed to evaluate the feasibility and safety of a disposable endoscope for upper gastrointestinal examination.

Methods In a prospective, randomized trial, 144 upper endoscopic procedures were allocated to either the disposable endoscope group or the conventional endoscope group. The primary outcomes were rates of excellent and good image qualities and maneuverability satisfaction. The second outcome included procedure duration, endoscopic diagnosis, and adverse events.

Results A total of 144 subjects were enrolled in the present analysis and prospectively randomized to 2 study groups. Finally, 70 and 69 subjects were enrolled in the novel disposable endoscope group and the conventional endoscope group, respectively, due to the schedule cancellation of 5 subjects. The baseline characteristics of the patients were similar in both groups. The excellent and good image quality rates and maneuverability satisfaction of the novel disposable endoscope were not inferior to the conventional endoscope (p = 0.99 and p = 0.99, respectively). Moreover, no significant between-group difference was observed in the endoscopic results and adverse events (p = 0.30 and p = 1, respectively). However, the procedure duration in the novel disposable endoscope was longer compared with the conventional endoscope (8.40 ± 4.28 min vs. 5.12 ± 2.65 min, p < 0.001).

Conclusions The novel disposable endoscope was as safe, effective, and maneuverable as a conventional endoscope. However, the novel disposable endoscope was associated with a longer procedure duration.
Methods and Materials

Study design
This was a multicenter, randomized, parallel, non-inferiority trial, and patients were recruited from 2 hospitals in China between December 2019 and June 2020. The study was approved by the Human Ethics Committees, and our research protocols complied with the Declaration of Helsinki. The study was registered online at http://www.chictr.org.cn (No. ChiCTR2000029945). Written informed consent was obtained from all subjects before their enrolment.

Novel disposable upper gastrointestinal endoscope
The novel disposable upper gastrointestinal endoscope (PR-IPD-002) was made by Shenzhen PengRui Intelligent Image Co., Ltd., and consists of a computer monitor, image processor, control body, and endoscope. The details are available in Appendix 1.

Patients
The inclusion criteria were set as follows: (1) patients aged between 18 and 75 years; (2) patients scheduled for upper gastrointestinal endoscopy, and (3) patients who provided informed consent.

Patients with the following conditions were excluded from the present study: (1) severe cardiopulmonary disease, (2) gastrointestinal perforation, (3) shock, (4) severe laryngeal disease, (5) acute erosive esophagitis (such as misuse of strong acid or strong bases), (6) thoracoabdominal aortic aneurysm, (7) history of mental illness or cognitive impairment, (8) allergic to anesthetic drugs, (9) coagulation dysfunction, (10) hemoglobin < 50 g/L, (11) pregnancy or lactation, and (12) participation in other clinical trials.
Endoscopists

All endoscopists were adequately trained for the use of the novel disposable upper gastrointestinal endoscope and the image quality grading system at the initiation of this study (Supplement Table S1). Many multiple upper gastrointestinal images were independently reviewed and scored by each endoscopist. All results were summarized, and discrepancies were discussed and resolved through consensus. All procedures were performed by experienced endoscopists who had performed more than 10,000 upper gastrointestinal endoscopies.

Randomization

Routine laboratory tests results of patients, including electrocardiogram, blood counts, HBV, HIV, human chorionic gonadotrophin, and COVID-19 real-time PCR, were within the normal ranges. The patients were randomly allocated to either the disposable endoscope group or the conventional endoscope group. The randomization numbers were generated using a computer random number generator and placed in sealed opaque envelopes, which were opened at the beginning of the upper gastrointestinal endoscopy. However, due to the nature of this study, it was impossible to blind endoscopists and participants to the randomized allocations.

Upper gastrointestinal endoscope procedure

All patients fasted for 8 h before the procedure, and a mixture of 100 mL water, 20,000 U pronase (Beijing Tide Pharmaceutical Company, Beijing, China), 1 g sodium bicarbonate (Beijing Tide Pharmaceutical Company, Beijing, China), and 10 mL simethicone (Berlin-Chemie AG, Berlin, Germany) was given to the patients before the procedure. All patients received anesthesia through intravenous injection of 5 mg midazolam and 50 mg pethidine.

Both the novel disposable endoscopy and conventional endoscopy were performed by experienced endoscopists from Shenzhen People’s Hospital and the University of Chinese Academy of Sciences Affiliated Shenzhen Hospital. A GIF-260 (Olympus, Japan) was used for the conventional endoscopy.

Definition

Comprehensive evaluation standards of image quality included observation, preservation, and analysis of digital image quality. The detailed evaluation contents are listed in Table S1.

Comprehensive evaluation standards for operability evaluation included operation satisfaction, visual fatigue, and physical fatigue. The detailed evaluation contents are listed in Table S2.

Procedure duration was defined as the time elapsed from the entry of the endoscope to the removal of the endoscope. Adverse

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Note: SMT, submucosal tumor.
events included throat injury, Mallory-Weiss syndrome, bleeding, and perforation during 1-day follow-up.

Outcomes
The primary outcomes were the rates of excellent and good image qualities. The secondary outcomes included assessment of maneuverability, procedure duration, endoscopic detection, and adverse events.

Statistical analysis
The following formula was used to calculate the sample size.

\[
\begin{align*}
\eta = & \frac{(Z_{1-\alpha/2} + Z_{1-\beta})^2 [P_T (1 - P_T) + P_C (1 - P_C)]}{(|D| - \Delta)^2} \\
\eta & = n_t = n_c
\end{align*}
\]

\( P_T \) indicates that the predicted excellent and good rates of image quality in the novel endoscope group were 96%. \( P_C \) indicates that the predicted excellent and good rates of image quality in the conventional group were 96%. \( |D| = |P_T - P_C| \). \( \Delta \) was the non-inferiority margin defined as \(-0.1\). Seventy-two cases were needed in each group according to an alpha of 0.05, a power of 10%, and a dropout rate of 15%.

All analyses were calculated using SPSS 23.0 software package (SPSS Company, Chicago, IL, USA). Categorical variables were expressed as frequencies (percentage), while continuous variables were expressed as mean ± standard deviation or median (interquartile range) due to the distribution. Between-group differences in categorical variables were analyzed using the Chi-squared test or Fisher’s exact test, and continuous variables were analyzed using the Student’s t-test or a Mann-Whitney test.

Results
Baseline characteristics
A total of 144 eligible subjects were enrolled in the present study and randomly allocated to the 2 study groups. However, 2 subjects in the novel disposable endoscope group and 3 subjects in the conventional endoscope group were excluded because of the schedule cancellation. Finally, there were 70 and 69 subjects in

![Fig. 1](image_url) Endoscopic images of the same anatomical regions of the upper gastrointestinal tract between the novel disposable endoscope and traditional endoscope.
the novel disposable endoscope group and the conventional endoscope group, respectively. There was no further dropout or loss to follow-up in both groups.

Table 1 shows that there was no significant difference in terms of gender and age between the 2 groups (p = 0.55 and p = 0.71, respectively). The indications for upper gastrointestinal endoscopic examination were similar between the 2 groups (p = 0.99). Moreover, no significant between-group difference was observed in the endoscopic results (p = 0.30). Indeed, there were no adverse events in either group (p = 1). However, the procedure duration in the novel disposable endoscope group was longer compared with the conventional endoscope group (8.40 ± 4.28 min vs. 5.12 ± 2.65 min, p < 0.001) (Table 1).

Image quality and maneuverability assessment

Fig. 1 shows endoscopic images at the same anatomical regions of the upper gastrointestinal tract in both groups. Fig. 2A shows that the rates of excellent and good image qualities were similar between the 2 groups (98 % vs. 100 %, p = 0.99). Upper gastrointestinal tract examination was successfully accomplished in both groups, and the maneuverability satisfaction of the novel disposable endoscope was not inferior to the conventional endoscope (98 % vs. 100 %, p = 0.99) (Fig. 2B).

Discussion

In the present study, we, for the first time, compared the feasibility and safety of a novel disposable endoscope and a conventional endoscope for upper gastrointestinal tract examination. We found that the image quality, maneuverability satisfaction, endoscopic diagnosis, and adverse events were comparable between the 2 approaches. However, the procedure duration for the novel disposable endoscope was longer compared with the conventional endoscope.

Previous studies have reported carbapenemase-producing Klebsiella pneumonia outbreaks in hospitals because of contaminated reusable duodenoscopes during ERCP procedures [11, 12]. Duodenoscopes have been an important issue particularly due to the difficult and often incomplete cleaning of the “elevator function” of this scope. The incidence of endoscope-related infectious transmission has been reported to be 3.7 and 1.6 per 1,000 gastrointestinal endoscopic procedures [1, 2]. Moreover, it estimated the cost of post-endoscopic infection is nearly $12,574.28 per hospitalization in the US, leading to a huge health insurance burden[13]. The cost of reusable endoscopes may be between $101.16 and $238.71 per endoscopic procedure including purchase, maintenance, reprocessing, and repair [13, 14]. Furthermore, even with strict compliance to all reprocessing and maintenance instructions, such as cleaning, inspection, and repair of the reusable endoscope, device-related infections persist [15].

In the present study, we demonstrated that the use of a novel disposable endoscope was as feasible and safe as the conventional endoscope for upper gastrointestinal tract examination. Moreover, there were several advantages of the novel disposable endoscope. First, the novel disposable endoscope may be more effective in preventing endoscope-related infections. Second, the novel disposable endoscope may be cost-effective in terms of reprocessing, personnel, maintenance, and repair cost. The cost of the novel disposable endoscope may be between $100 and $150 in the future. However, the estimated cost of reusable endoscopes is between $101.16 and $238.71 per examination [13, 14]. Therefore, the cost of the novel disposable endoscope may be relatively lower compared with the conventional endoscope once such an approach becomes widely utilized. Third, time spent on reprocessing will be saved in high-volume endoscopic procedure centers once the novel disposable endoscope becomes more widely used. Although the procedure duration of the novel disposable endoscope was longer compared with the conventional endoscope, this could potentially ameliorate with a growing expertise. Fourth, the disposable endoscope may also be beneficial to patients living in remote areas that do not have stringent reprocessing facilities.

Our study has some limitations. First, the sample size in this study was relatively small, and the feasibility and safety study of the novel disposable endoscope must be assessed further in larger cohorts. Second, the patient’s comfort level with the novel disposable endoscope could not be assessed as all patients were sedated during the procedure. Third, the effect of intra-observer and inter-observer bias could not be removed despite all endoscopists being adequately trained before the procedure. Fourth, the novel disposable endoscope was not capable of performing electronic staining and magnification, which may restrict the early screening of upper gastrointestinal tract cancer. However, as no premalignant or malignant lesions were detected in either group, it was difficult to assess whether the novel disposable endoscope was as good as the conventional endoscope in detecting premalignant or malignant lesions. Fifth, it is still unknown whether the novel disposable endoscope can be used for therapeutic procedures. Indeed, this would be a very important quality measure in the future.
Acknowledgement

Conflict of Interest

Funding

Author contributions

Li-sheng Wang and Jun Yao were responsible for the design of the study and reviewed the manuscript. Tao Dong and Yong-de Cai trained for the use of the novel disposable gastrointestinal endoscope. De-Feng Li performed data analysis and drafted the manuscript. Rui-yue Shi and Yan-hui Tian recorded data. Li-sheng Wang and Jun Yao were responsible for revising the manuscript. Jun Yao and Li-sheng Wang were responsible for revising the manuscript. All authors have read and approved the final manuscript.

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