Current Status and Research into Overcoming Limitations of Capsule Endoscopy

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Endoscopic investigation has a critical role in the diagnosis and treatment of gastrointestinal (GI) diseases. Since 2001, capsule endoscopy (CE) has been available for small-bowel exploration and is under continuous development. During the past decade, CE has achieved impressive improvements in areas such as miniaturization, resolution, and battery life. As a result, CE is currently a first-line tool for the investigation of the small bowel in obscure gastrointestinal bleeding and is a useful alternative to wired enteroscopy. Nevertheless, CE still has several limitations, such as incomplete examination and limited diagnostic and therapeutic capabilities. To resolve these problems, many groups have suggested several models (e.g., controlled CO₂ insufflation system, magnetic navigation system, mobile robotic platform, tagging and biopsy equipment, and targeted drug-delivery system), which are in development. In the near future, new technological advances will improve the capabilities of CE and broaden its spectrum of applications not only for the small bowel but also for the colon, stomach, and esophagus. The purpose of this review is to introduce the current status of CE and to review the ongoing development of solutions to address its limitations.

Key Words: Capsule endoscopy; Small bowel; Colon; Esophagus

INTRODUCTION

Capsule endoscopy (CE) has been available in clinical practice for the evaluation of small-bowel disease since 2001. CE has most commonly been used in cases of obscure gastrointestinal bleeding (OGBB). Various small-bowel capsules (PillCam, Given Imaging, Yoqneam, Israel; EndoCapsule, Olympus, Tokyo, Japan; MiroCam, IntroMedic, Seoul, Korea; OMOM, Jinshan Science, Chongqing, China; CapsoCam, CapsoVision, Saratoga, CA, USA) are now available worldwide. CE has many advantages compared to conventional wired endoscopy, such as convenience and less invasiveness. However, CE still has several significant technical limitations that need to be addressed technically. First, the gastrointestinal (GI) lumen is not inflated in CE, and only passive images can be obtained while the capsule passes through the GI tract. Therefore, lesions may be missed in CE, and the ampulla of Vater or ileocecal valve could be undetected in small-bowel CE. Moreover, CE often produces obstructed images because of bile, mucus, and other factors. It takes too long to administer CE and interpret capsule images. Second, CE cannot be used to take a biopsy specimen, nor does it have therapeutic capabilities. To overcome these limitations, several groups have suggested alternatives, and these alternatives are currently in development. Here, we introduce results from recent research about the future of CE.

CURRENT STATUS OF CAPSULE ENDOSCOPY

Small-bowel capsule endoscopy

With the advancements in CE technology, CE has rapidly expanded the indication for investigations of the GI tract. Given Imaging has installed more than 4,250 centers in about 60 countries and has sold >650,000 capsules. CE is a particularly
useful tool for patients with suspected small-bowel disease, including OGIB, iron deficiency anemia, Crohn’s disease (CD), tumors, polyposis syndromes, and celiac disease. A recent clinical guideline has recommended CE as a first-line investigation tool in patients with OGIB. The first available video capsule, brand name M2A (mouth to anus; Given Imaging), was approved by the Food and Drug Administration (FDA) in 2001 as an adjunctive tool for small-intestine imaging. At present, there are five small-bowel capsule endoscopy models on the market worldwide. Capsule endoscopy models with U.S. FDA approval consist of PillCam, EndoCapsule, and MiroCam. Although the various capsules are similar in size and shape, they differ in dimensions, frame rates, operating time, field of view, image sensor, and optical enhancements (Table 1). PillCam and MiroCam capture images by using a complementary metal oxide silicon sensor, whereas EndoCapsule and the OMOM capsule use a charge-coupled device. In 2013, the third generation of PillCam SB3 was launched and received FDA clearance. This capsule system has improved image detail and adaptive frame rate technology (two to six frames per second [fps]), leading to increased visualization of the small bowel and improved efficiency. The CapsoCam has four cameras that provide a 360° field of view, 12 to 20 fps, and 15 hours battery life. The CapsoCam camera takes images at a rate of 3 fps for the first 2 hours and thereafter at a rate of 3 fps, resulting in 12 and 20 fps, respectively. Smart Motion Sense Technology also enables the capsule to activate its cameras only during capsule motion.

To improve the specificity of small-bowel CE findings, fecal calprotectin (FC) is considered a noninvasive, “gold standard” marker of GI inflammation. An FC level of >100 μg/g is a good predictor of positive small-bowel CE findings, and FC >200 μg/g is associated with higher small-bowel CE yield (65%) and confirmed CD in 50% of patients.6

### Colon and esophagus capsule endoscopy

Colorectal cancer (CRC) screening programs in high-risk populations were reported to result in a 90% decrease in CRC incidence. However, no more than 25% compliance has been achieved in screening programs.7 Colon CE could be a good alternative in patients refusing conventional colonoscopy or when conventional colonoscopy is inappropriate or not possible. In a series of 328 consecutive cases, the rate of complete colon visualization before the end of the lifetime of the battery was 92.8%.8 In two prospective studies with the newer colon capsule endoscope PillCam COLON 2 (Given Imaging), the sensitivity reached 84% and 89% for detecting polyps >6 mm.9,10 Although colon CE showed similar detection capabilities when compared with conventional colonoscopy in some studies, conventional colonoscopy remains more accurate than colon CE, and allows the simultaneous removal of polyps. The colon capsule has a potential future for CRC screening; however, more data are needed to answer many pending uncertainties such as the best preparation method, best scoring method (CECDAI [CE CD Activity Index] or Niv score), best booster, and suitable indications.

In 2004, the PillCam ESO (Given Imaging), which is also capable of studying the esophagus, was developed and approved by the FDA. It has two lenses at both ends and takes 18 images per second over approximately 30 minutes to maximize visualization. Although CE can be used for detecting esophageal diseases such as esophageal varices, Barrett’s esophagus, and esophageal cancer, conventional upper GI endoscopy is still the gold standard.
Improvement of detectability during small-bowel investigation

In vitro chromoendoscopy

The Fujinon intelligent chromoendoscopy (FICE) system is a new virtual chromoendoscopy technique that processes reflected photons to reconstruct virtual images with a choice of different wavelengths by using computerized spectral estimation technology. The addition of the FICE technology to small-bowel CE may improve diagnostic yield. However, there is some controversy concerning its effectiveness. In a study by Gupta et al., FICE-assisted small-bowel CE analysis was no better than analysis with white light for the diagnosis and characterization of significant lesions in the evaluation of OGIB. Matsumura et al. found that although there was no improvement in diagnostic yield, FICE detected a significantly higher number of small-bowel lesions per examination than did conventional imaging (2.5±2.1 and 1.8±1.7, respectively). Krystallis et al. compared FICE and white light in a total of 167 images. FICE was ineffective in improving endoscopic images except in the blue mode. Blue filter provided image improvement in 83% of images when compared with white light. Imagawa et al. suggested the usefulness of FICE for visualizing small-bowel lesions such as angioectasia, erosions, ulcerations, and various tumors, in a retrospective study. However, in a prospective study, FICE improved the detectability of only angioectasia.

Efficient microcancer detection in the small intestine can be realized with infrared fluorescence endoscopy. Infrared fluorescence levels emitted by the fluorophore indocyanine green (ICG) at different concentrations are able to discriminate low concentrations of ICG in early cancer in the small intestine.

Three-dimensional reconstruction

In recent years, research has been carried out to produce a three-dimensional (3D) reconstruction of the GI tract; 3D imaging in CE is not currently feasible because of hardware limitations (i.e., packaging and size constraints, and power consumption). As an alternative method, a software-based approach (shape from shading) that enables 3D reconstruction from monocular 2D images has become available. Koulaouzidis et al. determined enhanced visualization for 56% of vascular and <10% of protruding structures (p=0.007 and p=0.008, respectively). Rondonotti et al. showed that the adjunction of 3D reconstructions to their standard 2D counterparts does not enhance the performance of expert small-bowel CE readers (p=0.245), although it significantly improves the performance of novices in differentiating masses from bulges (p=0.045).

LIMITATIONS OF CURRENT CAPSULE ENDOSCOPY

CE has many advantages, but also has several drawbacks (Table 2). Fortunately, many promising solutions have been proposed to resolve these problems (Table 3).

Incomplete small-bowel examination

Air insufflation

During its transition along the GI tract, the capsule endoscope is limited by air inflation in exposing the entire mucosa. Several groups have developed a novel device to achieve untethered controlled carbon dioxide (CO₂) insufflation suitable for CE. This device shows the feasibility of controlled inflation to facilitate visualization.

Retention or delayed transition

The most common causes of incomplete examinations are delayed gastric emptying and prolonged small-bowel transit. These result in the exhaustion of the battery before the capsule reaches the cecum. Administration of water or intravenous metoclopramide could be used in an effort to overcome this problem; however, it is necessary to use this method carefully in patients with dysmotility, as rapid transit time may diminish the diagnostic yield. Use of the external real-time viewer to check the progress of the capsule significantly improved the completion rate (86% vs. 66%, p=0.002) and the rate of positive findings (80% vs. 67%, p=0.04) compared with the

### Table 2. Advantages and Disadvantages of Capsule Endoscopy

| Advantages                        | Disadvantages                        |
|----------------------------------|--------------------------------------|
| Convenience                      | Incomplete small-bowel examination   |
| No need for sedation             | Uncontrolled air insufflation        |
| Simple examination for patient   | Retention or delayed transition      |
| Less invasiveness                | Limited battery life                 |
| High diagnostic yield comparable | Impossible to maneuver               |
|                                 | to other imaging modality            |
|                                 | No therapeutic or biopsy capability  |
Table 3. New Devices and Future Development

| Disadvantage                              | Promising solution                           |
|-------------------------------------------|----------------------------------------------|
| Incomplete small bowel examination        | FICE, IRFE, 3D reconstruction                 |
| Low quality image                         | Untethered controlled CO₂ insufflation       |
| Uncontrolled air insufflation             | FICE, IRFE, 3D reconstruction                 |
| Retention or delayed transition           | Untethered controlled CO₂ insufflation       |
| Limited battery life                      | External real-time image viewer              |
| Location                                  | Frame rate modulation                        |
|                                           | Video compression                            |
|                                           | Impulse Radio-Ultra-Wideband                 |
| Location                                  | Software using 3D triangulation               |
| Impossibility of maneuver                 | Magnetic navigation system                   |
|                                           | Mobile robotic platform                      |
| Therapeutic or biopsy capability          | Tagging, biopsy and therapeutic equipment    |
|                                           | Targeted drug delivery                       |
| Delayed time of the interpretation        | Frame rate modulation                        |
|                                           | Video compression                            |
|                                           | Impulse Radio-Ultra-Wideband                 |

FICE, Fujinon Intelligent Color Enhancement; IRFE, infrared fluorescence endoscopy; 3D, three-dimensional.

Fig. 1. New capsule endoscope proposed for self- or external ordinary positioning or propulsion. (A) A capsule that emits magnetic force. (B) A capsule with legs for mucosal ambulation. (C) A capsule that involves use of a paddling stroke. (D) A capsule with four propellers. Permissions for all pictures were obtained. (A) Adapted from Lucarini et al.\textsuperscript{28} (B, C) Adapted from Quirini et al.\textsuperscript{29} and Kim et al.\textsuperscript{30} with permission from Elsevier, respectively. (D) Adapted from Tortora et al.\textsuperscript{31} with permission from Taylor & Francis.
nonviewer group. In addition, retention can be detected by localization using a capsule emitting a magnetic field or electromagnetic waves. Olympus Medical Systems Corporation has developed new software with 3D triangulation. In a study on this method, the average total spatial error with attenuation was 13.26 cm.

**Low battery life**

The capsule battery life is usually 8 to 15 hours. Size reduction and battery life extension have become important challenges, as novel capsule endoscopes and accessory tools have been developed to improve diagnostic yield and perform therapeutic work. Frame rate modulation decreases the frame rate outside of the targeted area and saves battery power, yielding a longer operating time. Consequently, completion rate and diagnostic yield has increased significantly. Consumption can also be reduced by video compression and transmission technology such as compressed sensing theory and impulse radio ultra-wideband. One group has proposed primary magnetic coils in a power-generating device outside of the body to send power to a capsule within the body to save space.

**Controlled locomotion and positioning of the capsule endoscope**

Currently, the movement of the capsule is absolutely dependent on gravitational and peristaltic force. Thus, many methods have been proposed for self- or external ordinary positioning or propulsion (Fig. 1). Although there are differences among capsule models, they are classified into three types: magnetic force, motion of leg or paddle, and propeller. The magnetic-enabled PillCam and magnetic upper GI Olympus capsules have been devised for this purpose. The magnetic-enabled PillCam was based on the Niobe magnetic navigation system (Stereotaxis, St Louis, MO, USA). The core of this system consists of focused-field permanent magnets, made of a neodymium-iron-boron compound. These large magnets are mounted on automatically operated arms to be easily arranged and oriented on either side. In an in vivo experiment, the capsules showed an accuracy of 1° and a localization error of 1 mm. In another study, the camera could be rotated in steps of 1.8°. Full 360° visualization was possible in the stomach but only a 45° visual field was possible in the colon. In 52 human subjects, a feasibility study on a magnetically guided capsule from Olympus demonstrated that visualization of the antrum, body, fundus, and cardia were at 98%, 96%, 73%, and 75%, respectively, and the feasibility of gastric examination was shown.

Endoscopic devices with flexible legs for ambulation have also been proposed. Legged locomotion mimicking inchworm motion has several advantages, including better adaptability to different geometries of the GI tract, higher velocity, and simplified adhesion induced by friction between the device and the tissue. A new prototype with eight legs has been devised. The two leg sets (each leg set consists of four legs) open independently in opposite directions; the rear legs are useful for propulsion, whereas the front legs are useful for stopping. In in vitro tests, the eight-leg capsule showed a speed of 6 cm per minute, and capabilities of backward and vertical locomotion. A paddling-based locomotion mechanism enhanced CE by using a paddling stroke. It provided fast locomotion speed and long travel distances. In one study, the mean velocity was 37.5 cm per minute in the extracted porcine colon and 17 cm per minute in the colon of a living pig. Another study reported that CE with feedback controlled paddling had higher locomotion speed, showing an increase of 58% compared with the previous control method based on a given timer value. A capsule endoscope with an integrated propeller has been developed, and recently, a novel capsule endoscope with four propellers has allowed for a reliable 3D locomotion if the capsule has neutral buoyancy. The number of blades per propeller usually varies from one to five, although three-blade propellers are commonly used. However, this type was useful only for situations in which a large amount of liquid was present. Therefore, liquid intake is required for filling the GI cavity.

For better locomotion and steering, some groups have suggested a hybrid of the magnetic system and self-propelled capsule endoscope, as well as of the magnetic system and legged locomotion. However, these methods require further development concerning the need for bulky and complex equipment for generating the magnetic field, and concerning accuracy and stabilization levels, which remain relatively low.

**Beyond detecting lesions (ongoing projects)**

A tagging module can mark the precise location of a target lesion for future surgical or wired endoscopic therapy. One prototype biopsy module consists of a trigger with a paraffin block, a rotating tissue-cutting razor with torsion spring, and a controller. It is constructed to operate sequentially so that the tissue sampling, sealing, and fixing are performed in one operation. Another form with a microactuator has been designed to perform biopsy. A microspike was incorporated into the capsule endoscope to obtain biopsy specimens. Experimental tests demonstrated that the developed microactuator with microspike successfully extracted tissue samples from a pig’s small intestines. One study reported that a magnetically maneuvered capsule with a nitinol clip-releasing mechanism successfully clipped an iatrogenic bleeding lesion in a pig model. Two new capsules, Intellisite
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(Innovative Devices, Raleigh, NC, USA) and Enterion (Phaeton Research, Nottingham, UK), have been developed for the collection of absorption data in the GI tract and can be used in the future for drug delivery. Nonvideo capsules that can deliver drugs with a pH-activated or temperature-activated release mechanism have also been evaluated. Wood et al. reported that a capsule endoscope prototype with pH, temperature, and pressure sensing, and a pin for anchoring with a holding mechanism, demonstrates the feasibility for targeted drug delivery.

The NEMO (nano-based CE with molecular imaging and optical biopsy) project is developing a new capsule that combines optical and maneuvering technologies, biosensing, and nanotechnologies to enhance the diagnostic and therapeutic potential of CE. The VECTOR (versatile endoscopic capsule for GI tumor recognition and therapy) project is in the process of developing a minirobot for the screening and surveillance of GI cancer, and for magnetic and legged motion, drug delivery, and tissue sampling. A prototype coagulation capsule that employs an exothermic chemical reaction to generate heat through the interaction of calcium oxide and water has been tested. This may be potentially useful for hemostasis through thermal coagulation. However, improvements in capsule maneuvering capabilities are necessary before these capsules can be further developed.

CONCLUSIONS

CE has evolved very rapidly to become an important tool for the visualization of the gut mucosa. Small-bowel CE is recommended as the first-line investigation technique in patients with OGIB and seems sufficiently accurate as an alternative tool in other small-bowel diseases such as CD, small tumors, celiac disease, unexplained abdominal pain, and/or diarrhea. For complete and perfect small-bowel investigation, several technical limitations must be addressed. Fortunately, many methods have been proposed and are in development, such as enhanced image modality, controlled air insufflation, decreased battery consumption, and several therapeutic and biopsy tools (Fig. 2). With technological developments, advanced CE could become the standard method of endoscopy.

Fig. 2. Schematic illustration of future imaginary capsule endoscopy based on current research. This illustration was made by the present author, Won Gun Kwack. FICE, Fujinon intelligent chromoendoscopy; IRFE, infrared fluorescence endoscopy; 3D, three-dimensional; LED, light-emitting diode; DSP, digital signal processing.
for many GI diseases in the future.

Conflicts of Interest

The authors have no financial conflicts of interest.

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