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**Small bowel capsule endoscopy: Where are we after almost 15 years of use?**

Van de Bruaene C *et al*. Almost 15 years SB CE

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**Abstract**

The development of Capsule Endoscopy (CE) in 2001 has given gastroenterologists the opportunity to investigate the small bowel in a non-invasive way. CE is most commonly performed for obscure gastrointestinal bleeding, but other indications include diagnosis or follow-up of Crohn’s Disease, suspicion of a small bowel tumor, diagnosis and surveillance of hereditary polyposis syndromes, Nonsteroidal anti-inflammatory drug-induced small bowel lesions and Celiac Disease. Almost fifteen years have passed since the release of the small bowel capsule. The purpose of this review is to offer the reader a brief but complete overview on small bowel CE anno 2014, including the technical and procedural aspects, the possible complications and the most important indications. We will end with some future perspectives of CE.

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**Key words:** Capsule endoscopy; Small bowel; Preparation; Procedure; Technology; Complications; Features; Enhancements; Indications; Future

**Core tip:** This review covers all the relevant aspects of Small Bowel Capsule Endoscopy anno 2014. The current techniques, procedures, analyses, indications and future perspectives are discussed thoroughly. Easy-to-use flowcharts are provided to help the readers in their decision-making when confronted with small bowel pathology.

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**NTRODUCTION**

Wireless Video Capsule Endoscopy (CE) was invented by Gavriel Iddan[1] in the mid-1990s. Being able to visualize the entire small bowel in a noninvasive, well-tolerated way, CE has closed the diagnostic gap between conventional gastroduodenoscopy and colonoscopy. Since the official release of CE in 2001, almost 15 years have passed, and CE has revolutionized the diagnosis and treatment of various small intestinal diseases. This review aims to provide state of the art on CE in gastrointestinal diseases. Both the evolution in technique and in indications will be discussed.

**TECHNICAL PRINCIPALS, PROCEDURE AND ANALYSIS**

***Capsule Definition***

The wireless CE system consists of 4 main parts: (1) the single-use wireless Video Capsule; (2) sensor arrays or a sensor belt attached to the patient; (3) the data recorder attached to the belt; and (4) the computer workstation with the application software[2-4] as can be seen in Figure 1 by Pan G *et al*[4].

The capsule weighs less than 4 g and measures about 11mm in diameter x 26 mm in length. It is made of plastic, biocompatible and resistant to digestive fluids. The capsule contains a short focal lens and a miniature video camera: a charge-coupled device or Complementary Metal Oxide Semiconductor, which focuses the image. The gastrointestinal tract is illuminated by white light LED’s. The capsule is powered by two mercury free silver oxide batteries with a life span of 8-12 h. More than 5,000 images are transmitted during this battery life at a rate of 2-6 fps. Capsule features have evolved since the release of the first capsule and nowadays standards are a 156-170° field of view, a high resolution and sharpness with a minimum size of detection of 0.07 mm, a 1:8 magnification, a more homogenous light exposure and a depth of view of at least 20-30 mm[5]. The captured data are sent to the sensor arrays and belt worn by the patients by either ultra-high frequency band radio telemetry or human body communications, using the human body as conductor.

At present, there are three main companies supplying wireless CE systems by approval of the FDA. Given Imaging (Ltd, Yoqneam, Israel) supplying the PillCam® SB 3, Olympus America (Inc, Center Valley, Pennsylvania) supplying the EndoCapsule® and Intromedic Company (Ltd, Seoul, Korea) manufacturing the MiroCam®. Although not approved by the FDA, another Chinese company Jianshan Science and Technology (Group) Co., Ltd., Chongqing, has developed its own capsule: the OMOM capsule. The capsule has been approved by the State Food and Drug Administration of the People’s Republic of China in March 2004 and is since then being used in China, Southeast Asia and some European countries[6]. The first large clinical trials have reported a yield and completion rate similar to the PillCam, but the major advantage of the OMOM capsule is without doubt its price, which could be reduced by fifty percent[6,7] (Table 1 and Figure 2)

Even though the three main capsules approved by the FDA differ in technical specifications, several trials have shown that they offer a comparable diagnostic yield, image quality and completion rate, as was stated in the systematic review by Koulaouzidis A *et al*[8].

***Small bowel preparation***

To ensure a clear view on capsule endoscopy, the patient is asked to start fasting 12 h before the small bowel Capsule Endoscopy procedure[2,3]. However, due to bubbles, small intestinal fluid and biliary secretions coming from the major duodenal papilla, the visualization by the VCE can deteriorate. Furthermore, limited battery life span can hamper a complete intestinal examination in patients with delayed gastric emptying and small bowel transit, which necessitates the use of additional small bowel preparation[3]. However, not all patients are eligible for small bowel cleaning. The 2012 Consensus guidelines for the safe prescription and administration of oral bowel-cleansing agents[9] states that small bowel preparation is contraindicated in patients with gastrointestinal obstruction, perforation, ulceration, ileus, gastric retention or Inflammatory Bowel Diseases (IBD), in patients with a reduced level of consciousness, swallowing disorders, hypersensitivity to the used agent and in patients having an ileostomy. The use of small bowel cleansing agents is relatively contraindicated in patients with Chronic Kidney Disease or undergoing dialysis, in patients with a renal transplant, congestive heart failure, liver cirrhosis or ascites and in patients taking Renin-angiotensin blockers, diuretics or Nonsteroidal anti-inflammatory drugs (NSAIDs). In these patients the utility of small bowel cleaning should be reconsidered and the choice of cleaning agent is of main importance: PEG is normally preferred over Sodium Phosphate. Patients taking Renin-angiotensin blockers, diuretics or NSAIDs are advised to discontinue their medication temporarily and their hydration and electrolyte status should be checked prior to the small bowel preparation. In a recent systematic review, Kotwal *et al*[10] compared the results of various randomized-controlled trials regarding improvement of vision quality (VQ), diagnostic yield (DY) and completion rate (CR) by small bowel preparation. In this review, administration of 2L polyethylene glycol (PEG) the evening before VCE was found to be superior to two doses of 45 mL Sodium Phosphate before VCE regarding VQ and DY improvements. Another study by Kantianis *et al*[11] showed that 2 L as well as 4 L PEG did not differ in small bowel cleansing and CR. Therefore 2 L should be preferred as regimen before VCE. Furthermore Kotwal VS *et al*[10] stated that simethicone, an antifoaming agent, significantly improved the VQ by decreasing the bubbles without implications on CR. Yet no significant improvement in VQ and DY were observed by combining Simethicone with PEG. After meta-analysis prokinetics did not show a significant improvement in CR, so they are not recommended.

***Procedure***

After bowel preparation, the patient gets eight sensor arrays attached to his body and a sensor belt fastened around his waist. The data recorder is attached to the belt before capsule ingestion. The capsule is ingested with a glass of water and fluid restriction is needed till 2 h after ingestion. After 4 h, fasting can be stopped. Daily activities do not need to be interrupted during capsule endoscopy.

Capsule propulsion needs to be followed by real-time viewing during the first hour to make sure the capsule passes the stomach. If not, gastroscopy is performed to deposit the capsule in the duodenum.

The sensor arrays and belt are removed once the capsule has been expelled into the colon (as verified by real-time viewing), or when the battery life has expired. Images can be downloaded from the recorder to the workstation. The capsule itself passes naturally with bowel movement and is usually excreted within 24 to 72 h.

***Analysis***

After downloading the data from the recorder to the workstation, images can be reviewed by gastroenterologists using the application software. Reading time and interpretation are around 40-120 min[2,12] which can be, compared to conventional endoscopy, a time-consuming activity. A solution to this problem might be to train nonphysicians in prereading the images. A study by Bossa *et al*[12] found that a nurse with expertise in endoscopy might be able to shorten the time needed by the endoscopist to read a capsule. Moreover the prereading of the CE by the nurse endoscopist led to a more careful approach of the physician in the flagged areas, which enhanced the accuracy of the CE investigation. Another recent study by Dokoutsidou *et al*[13] confirmed these findings and stated that despite a longer reading time, a nurse is perfectly capable of prereading and subsequently flagging aberrant images. However, another possibility to lower reading time is the use of special software to select aberrant images, which can be revised afterwards. With the introduction of Quickview by Given Imaging, reading time could be reduced significantly. The Quickview software samples sites of interest for review at a chosen rate, but unfortunately missed lesions occur far more often, which is unacceptable[14]. However, in certain clinical settings, such as overt obscure gastrointestinal bleeding (OGIB) in an urgent inpatient setting and suspected Crohn Disease or occult OGIB in outpatient setting, Koualouzidis *et al*[15]found that Quickview could be used confidently without clinical consequences. To enhance the yield of CE, virtual chromoendoscopy was developed by adding colour filters to the images. Fuji Intelligent Colour Enhancement (FICE) was seen to be superior in detecting small bowel lesions and in particular angioectasias compared to conventional CE[16]. In another trial by Krystallis *et al*[17] Blue Mode (BM) was found superior to FICE in detecting lesions of the small bowel. Adding BM to Quickview studies however did not show any diagnostic advantage and is therefore not recommended 15].

**COMPLICATIONS AFTER CE**

***Capsule retention***

Although very popular for its non-invasive character, capsule endoscopy can be the cause of unnecessary treatment due to complications. One of the most feared complications is capsule retention. It is defined as the presence of the capsule in the bowel lumen for a minimum of 2 wk after ingestion, or when the capsule is retained for an unspecified period of time unless targeted medical, endoscopic or surgical intervention is started[18].

According to a systematic review by Liao *et al*[19] overall retention rates are as low as 1.4%, which makes the procedure acceptable, regarding the high overall diagnostic yield of 59.4%. Furthermore, the retention rate differs according to the underlying pathology, with up to 2.6% in known Crohn’s Disease (CD) and 2.1% in patients with Neoplastic Lesions[19]. This can be explained by the fact that capsule retention is usually caused by masses, strictures and stenotic areas resulting from neoplastic lesions, CD, NSAID consumption or post-operative adhesions, which narrow the small bowel lumen and favors retention[20]. In this regard, known small bowel obstruction, strictures and extensive CD are a contraindication for CE[21]. In a large study by Höög *et al*[22] risk factors for capsule retention were identified. OGIB and suspected CD were associated with the lowest chance of capsule retention, whereas known CD and small bowel tumors had a higher chance of retention. These findings were also confirmed by other authors[19,23].

Most of the time, retained capsules are asymptomatic, but intestinal obstruction, partial or complete, may occur, especially in case of known CD or neoplastic lesions. In the 2009 consensus by the European Society of Gastrointestinal Endoscopy it is recommended with a grade B evidence level to precede the CE by small bowel imaging or a patency capsule (cf. infra) in patients with suspected or established CD to rule out potential strictures. As said earlier, known small bowel obstruction is a contraindication for CE and patients at risk for a small bowel obstruction should therefore be carefully investigated by their physician before a CE procedure[24].

Evacuation of the retained capsule can be spontaneously, medically or by surgery. The latter is unfortunately the most frequent intervention, but is on the other hand safe and can be seen as a required diagnostic and therapeutic tool for treating the underlying small bowel condition. With the surgery, not only the capsule is removed, but also the responsible lesion can be resected, which reliefs the patient’s symptoms. However retention can also lead to unnecessary surgery of lesions caused by *e.g.* NSAID or CD, for which a medical solution would also have been an option[25].

In recent years, an endoscopic approach of capsule retention has become more popular as a less invasive alternative for surgery. Before capsule retrieval a radiographic localization of the capsule is done to determine whether an upper or lower gastro-intestinal and a standard (gastroduodenoscopy, Push Enteroscopy or colonoscopy) or advanced endoscopic approach (device assisted enteroscopy) is needed. In this regard, surgery can be considered when endoscopic approaches did not manage to retrieve the capsule or when the patient presents with symptoms of toxicity[26]. In a study by Van Weyenberg *et al*[27] DBE showed to be an adequate tool to retrieve a retained capsule. Moreover the DBE was capable to aid in pre-operative staging by histological sampling. In conclusion DBE can prevent unnecessary surgery as well as determine the cause of the capsule retention before the operation, which is beneficial both for physician and patient.

***Capsule perforation***

Another yet very rare complication is perforation of the small bowel. Usually it results from capsule retention. In the few cases that are reported, CD was the most frequent underlying pathology causing the perforation[20, 28-31]. In a study by Repici *et al*[28] a possible explanation for this complication is given. CD affects the tissue of the small bowel wall and makes it vulnerable. By the complete luminal occlusion due to the entrapment of the capsule and the high peristaltic activity the fragile tissue of the small bowel wall distends just above the capsule and leads to fissuration and possible perforation of this area. One study by Gonzalez *et al*[32] reported perforation after CE in a patient with a history of surgery with subsequent adhesions. Because of the major implications, perforation should be acknowledged as a possible complication after CE in patients with known or suspected CD.

***Capsule interference***

One of the relative contra-indications for CE is the presence of implantable cardiac devices, such as pacemakers, ICDs, pulsatile and nonpulsatile LVADs. Interference may arise during the CE procedure resulting in an alteration of atrial or ventricular assistance[33]. This is however a theoretical assumption without clinical significance, because few studies show actual interference between cardiac devices and CE. Moreover, previous studies have already suggested that CE can be used safely in patients with these devices[34-41]. Only one study by Dubner *et al*[42] reported oversensing of an ICD due to interference with the CE procedure, which resulted in an inappropriate shock by the cardiac device. In another case report by Guyomar *et al*[37] interference between pacemaker and video capsule occurred, resulting in a failure of recording by the capsule when close to the pulse generator. Harris *et al*[43] found similar results in a 2013 study: all implantable devices proved to be safe for the patient, but LVAD had the tendency to interfere with image capture and therefore a CE lead position as far away from the LVAD as possible is required. On the other hand, in the article by Cuschieri *et al*[40] no loss of images was observed. In conclusion, close monitoring is recommended in patients with implantable cardiac devices but the risk for complications seems to be extremely low.

***Capsule aspiration***

Some cases have reported the existence of bronchial aspiration of the capsule. It is a very rare complication, which occurs in one out of every 800 investigations and can be asymptomatic[44]. Capsule endoscopy aspiration can resolve spontaneously[45], but often necessitates immediate radiological investigations to localize the capsule, followed by bronchoscopy to retrieve it with the aid of a Roth Net[44]. To prevent this unnecessary invasive procedure, screening for patients at risk should be done. Risk factors include aging, neurological or swallowing disorders and patients with a weak or absent cough[46,47]. Direct placement of the capsule in the gastrointestinal tract should be considered in these patients [44-46,48]. If not, the Real Time Viewer should be used during the ingestion of the capsule to make sure that the capsule reaches the gastrointestinal tract[46-48].

Until now, only one fatality has been reported due to intracerebral haemorrhage resulting from capsule aspiration[47]. The reason for this low mortality rate is hypothesized in a study by Lucendo *et al*[44] stating that the size of the capsule is not capable to block the total lumen of the trachea and therefore still allows adequate oxygenation after capsule aspiration. However Koulaouzidis *et al*[49] found that the CE size might be correlated with the chance of aspiration.

**ADDITIONAL FEATURES AND ENHANCEMENTS IN THE FIELD OF CE**

***Suspected Blood Indicator***

In 2003 Given Imaging introduced the Suspected Blood Indicator (SBI) as an aid in diagnostics. The new feature highlights images suspected for redness or blood, which makes it easier for physicians to identify possible bleeding sites accurately. The software is activated when the capsule has reached the duodenum and operates only during its stay in the small bowel[50].

Sensitivity of the SBI software is determined by the presence of active bleeding. In studies, sensitivity ranges from 20% to 56.4% and increases up to 58.3% to 93% in case of active bleeding[50-52]. However, sensitivity and specificity of SBI remains too low, so complete review by a gastroenterologist is still required and the SBI only serves as rapid screening tool for actively bleeding lesions[51,52].

The detection rate of the SBI is affected by background color of the small bowel as by velocity of the capsule[50]. This is also a possible explanation for the variation in sensitivity observed among different studies. The background of the small bowel differs according to patient’s condition and small bowel preparation[50]. In experimental small intestine models, a very pale magenta background showed the highest detection rate, followed by burnt sierra and yellow. Lowest detection rates were observed in small bowel sites with colors significantly different from the normal small bowel color or when the capsule reached a high velocity. In an interesting study by Buscaglia *et al*[53] SBI was found to be an inferior screening tool for sites of potential bleeding with a sensitivity below 60% even in active bleeding. Yet they found that in CD the SBI could be used as a screening tool for detection of aberrant mucosa with high sensitivity. Another study by D’Halluin *et al*[54]also rejected the SBI software as an useful tool for screening the small bowel stating that the detection rate was poor, independent of the type of lesion. Furthermore they found that the SBI missed certain lesions while tagging few others and that irrelevant flagging might unnecessarily prolong the reading time of the CE. However, in a recent study Tal *et al*[55] stated that SBI is a reliable aid in excluding active bleeding or major lesions, but that the role of the endoscopist could not be neglected. In summary we can conclude that SBI might improve the interpretation and thereby the yield of CE by tagging areas for a second review, but can certainly not replace the gastroenterologist’s review.

***Patency capsule***

To address the problem of capsule retention, the Agile Patency Capsule (PC) was developed by Given Imaging. The patency capsule with the same size as a video capsule, serves as dummy to assess the patency of the small bowel prior to CE examination. As one of the major contraindications for CE is suspicion of small bowel stenosis, routine administration of PC’s could enable safe CE use in a larger patient population by ruling out possible stenoses[56]. The PC system consists of two main parts: the capsule itself with a radiofrequency identification tag (RFID tag) and an external detector system to capture radio-frequency signals.

The patency capsule is made of lactose and 10% barium, which dissolves when coming into contact with intestinal fluids through the window at the edge of the capsule, also known as timer plug. To insure that the timer plug is not blocked by capsule impaction in a stricture, the second generation PC’s consist of two timer plugs. If excretion does not occur, dissolution starts at 30 h. After 35 h, 38 percent of the capsules are dissolved and all are dissolved within 72 h[57]. After dissolution, the remains of the capsule encounter no difficulties to pass the small bowel strictures.

The detector system receives the radio-frequency signals coming from the RFID tag and reconstructs the exact capsule position. This can also be done by using radiography, which visualizes the PC by its radiopaque RFID tag or 10% barium[56]. Localization can be complicated by overlap of intestines so subsequent fluoroscopy or CT scan can be warranted. One drawback of the RFID tag system is the probability of impaction in a stricture, which can lead to small bowel ileus[58]. Recently a new tag-less PC was developed by Given Imaging to overcome this issue and proved its usefulness as found by Nakamura *et al*[58].

The PC procedure is not as strict as the CE procedure. The capsule can be swallowed without previous food restrictions. If the capsule is not excreted in 33 h, further examination is warranted to localize the PC and make a distinction between a small bowel and a colonic localization. The latter is still an indication for VCE. The subsequent capsule endoscopy has to be done quickly after PC so a possible change of stricture status and subsequent capsule retention is avoided[56].

The use of the patency capsule still remains controversial. Although some authors have reported its utility[59-61], others have found that the capsule was not capable of confirming stenoses, which were found on CT or small bowel follow through[62]. In conclusion, patients still benefit from a CT investigation prior to CE to exclude possible stenosis and strictures.

***CapsoCam capsule***

Over the last decade, a new player entered the field of capsule endoscopy. CapsoCam by Capsovision renewed the concept of capsule endoscopy by offering a capsule with a 360° view and on-board storage, which enables the retrieval of images wire-free after interception of the capsule in the feces. The capsule contains four cameras, which offer high resolution images and a frame rate up to 20 fps max. Furthermore, two new technologies were developed, Smart Motion Sense Technology and Auto Illumination Technology. Smart Motion Sense Technology enables the capsule to activate its cameras only during capsule motion. When the capsule is stationary, a sensor is used to compare the current frame with the previous frame to control reactivation. Auto Illumination Technology controls the 16 white LED’s to provide the optimal level of illumination. When the capsule is located nearby the walls, a low light intensity is optimal to capture the best images. A position further away from the wall necessitates a higher light intensity. By adding these software features, battery life is sustained up to 15 h. The first clinical trial that used the CapsoCam accepted it as a safe and efficient tool in small bowel evaluation[63]. In a recent French study by Pioche *et al*[64] the concordance between the PillCam SB2 and CapsoCam was evaluated in terms of diagnostic yield and image quality. A kappa value of 0.63 was obtained, which confirms the good concordance between the two capsules. Although the reading time of the CapsoCam was longer, the CapsoCam detected significantly more lesions in a per lesion analysis.

***Three-dimensional representation***

In recent years, three-dimensional representation is becoming a hot issue. By reproducing the depth information lost by camera recording, diagnosis can be facilitated, because the texture and the abnormalities of the mucosa are highlighted. 3-D rendering can be software- or hardware-based. The latter is limited by the technological possibilities of the capsule, so software based 2-D to 3-D conversion is used[65,66]. Software-based 3-D rendering uses algorithms to recreate the third dimension. In a study by Karargyris *et al*[67] four Shape-from-Shading (SfS) algorithms were compared. The Tsai’s SfS algorithm excelled the other algorithms in visualization improvements, but we may not forget that the evaluation criterion was subjective in origin. However, the Tsai’s SfS algorithm is especially adapted to bright and round surfaces, therefore perfectly applicable in small bowel endoscopy.

***Lesion Localization***

Apart from the image quality, accurate lesion localization is one of the key elements of capsule endoscopy, because further therapeutic steps, non-invasive and invasive, can depend on the exact localization of the lesion[68]. Lesion localization is currently estimated according to the transit time and the use of pylorus and caecum as landmarks, but lacks precision.

Exact localization can be determined by using a capsule emitting a magnetic field or electromagnetic waves. Both methods have their advantages and drawbacks. Magnetic-field-strength-based localization is not attenuated by the human body and the capsule does not have to be aligned with the detectors to be detected. As a drawback, interference of the magnetic fields for capsule localization and the magnetic fields for active capsule movement in the future (cf. infra) may occur. On the other hand electromagnetic waves localization, such as the previously mentioned RFID tag, is based on radio-frequency waves, which are attenuated by the human body and therefore may lose precision. A promising step forward in capsule localization is the development of a new software by Olympus Medical Systems Corporation (Tokyo, Japan), which uses a 3D-triangulation. The exact capsule position is calculated by determining its distance from the 6 radiofrequency sensors using radiofrequency signal strength. In the study by Marya *et al*[69] an average localization error of 13.26 cm3 by attenuation was observed, especially in patients with an increased BMI.

Finally, in 2010 the Capsule-odometer, a conceptual CE design, was proposed by Karargyris *et al*[70] which in theory offers a more accurate lesion localization. The capsule has two protruding wheels attached to a spring-mechanism, so the wheels can adapt to the diameter of the intestinal lumen, serving as a micro-odometer with subsequent accurate lesion localization, calculated from the onset of the capsule investigation. This design also offered a greater stability, avoiding non-forward movement through the gastrointestinal tract. Further experiments and research are needed on this subject.

**MAJOR INDICATIONS FOR SMALL BOWEL CAPSULE ENDOSCOPY**

Capsule endoscopy has been approved for various indications. These include (1) overt and occult obscure gastrointestinal bleeding; (2) suspected CD; (3) surveillance in patients with polyposis syndromes and detection of small bowel tumors; (4) screening and evaluation of NSAID side-effects; and (5) suspected malabsorptive syndromes such as Celiac Disease. These indications will be explained further on in this paper. Relative contra-indications for capsule endoscopy include, like mentioned before: (1) known or suspected GI obstruction, strictures or fistulas; (2) cardiac devices; (3) swallowing disorders; and also (4) pregnancy.

***Obscure Gastrointestinal Bleeding***

Obscure Gastro-Intestinal Bleeding (OGIB) is defined as bleeding of unknown origin that persists or recurs following a bidirectional negative endoscopic evaluation of the gastrointestinal tract. OGIB is a common problem encountered by gastroenterologists, and accounts for approximately 5% of all GI bleedings[71]. OGIB can be overt (melena, hematochezia, hematotemesis) or occult (iron-deficiency anaemia, IDA, with or without a positive fecal occult blood test). OGIB is mostly caused by a lesion located in the small bowel, but can also originate from a lesion in the other parts of the GI tract as well, missed with conventional endoscopy because of intermittent bleeding or by human error[72]. The underlying pathology is age dependent. Under the age of 40, the most frequently detected lesion is a small bowel tumor, followed by Meckel’s diverticulum, Dieulafoy’s lesion and CD. Above the age of 40, vascular lesions such as angiodysplasia are most frequently observed, counting for up to 40% of the underlying lesions. NSAID-induced lesions (cf. infra) are the second most frequent finding on CE[71].

Since its development in 2000, capsule endoscopy has mainly been used for the indication of OGIB, accounting for 60% - 70% of the patients[8]. CE has proven superiority to all other diagnostic modalities in OGIB, such as barium contrast radiology, small bowel computed tomography (CT), magnetic resonance imaging (MRI), push enteroscopy and angiography, as can be seen in Table 2. The American Society for Gastrointestinal Endoscopy (ASGE) confirmed these findings in their guidelines presented in 2007[71]. Before using capsule endoscopy as a diagnostic tool, at least one gastroduodenoscopy and ileocolonoscopy have to be performed to rule out upper and lower gastrointestinal tract abnormalities. Repeating gastroscopies or colonoscopies immediately prior to CE in patients who have not had endoscopic investigations for more than 6 months, tends to have a low diagnostic yield and is not cost-effective[73]. Therefore, CE is recommended as the first-line investigation after negative bidirectional endoscopies. Younger patients however have a higher chance of IBD or tumours and a CT abdomen is indicated prior to CE to rule out stenosis[20]. A gynaecological etiology has to be considered in young females.

The overall yield of capsule endoscopy is between 35% and 83% for OGIB[19,71,72,74-80] with its mean around 60%[81,82]. Diagnostic yield is influenced by the type of bleeding. Patients with ongoing overt bleeding usually present with a higher diagnostic yield than patients with obscure-occult bleeding, presenting as IDA[72,83]. More factors associated with a higher diagnostic yield have been identified, including low hemoglobin measurements, transfusion need, older age and a short interval of less than 3 days between admission and the CE procedure[84-88]. CE is recommended in all cases of OGIB because of its diagnostic value and its impact on further management. A study by Albert *et al*[75] found that CE was able to determine the therapy in 66% of the cases and led to an alteration in management in 32.3% of the cases. This is in line of previous studies, which reported that CE could alter subsequent management in 23-66% of the cases[79,85,89-91]. Sidhu *et al*[84] found that this management alteration could be predicted by patient comorbidity or angiodysplasia findings on CE (Table 2).

The reason why CE has been recommended as first-line examination tool over DBE after initial negative upper and lower endoscopies is its noninvasive nature and ease of use, which makes it well-tolerated and feasible in an outpatient setting[92]. Furthermore its ability to visualize the whole small bowel in more than 80-85% of the cases[93,94] and the ability to determine the initial DBE approach makes it a helpful tool in OGIB diagnostics[92,95]. However, CE often fails to visualize lesions in the proximal small bowel, in a Roux-en-y loop and in patients presenting with diverticula[76].

If necessary, a CE procedure can be followed by a Double Balloon Enteroscopy (DBE) procedure[72]. DBE is the only diagnostic tool showing a similar diagnostic yield for OGIB as VCE, as can be seen in Table 2. However, the DBE procedure is more invasive, can be time-consuming, requires training, needs sedation or general anesthesia and can have a complication rate of up to 4.3% in therapeutic procedures as was reported by Mensink *et al*[96]. Moreover DBE is not always able to visualize the whole small bowel. A completion rate of only 62.5% was achieved in DBE, compared to 90.6% in CE as reported by Nakamura *et al*[94]. Yet, DBE is preferred over CE in patients requiring a biopsy or a therapeutic intervention such as argon plasma coagulation (APC). Also DBE tends to have an acceptable yield in patients with an initial negative CE and suspicion of a small bowel lesion[95,97], although it has been reported being much lower than the yield of DBE following a positive CE, respectively 28% and 75%[77].

Not only clinically, but also economically is CE recommended as first line investigation of OGIB. It has shown to be more cost-effective than DBE when only visualization of the small bowel is needed[98]. Negative CE investigations usually do not require further diagnostic work-up, which saves money in the long term, because reimbursement for CE is less than for DBE[93]. A mean cost-saving of €1738.07 was reported by Marmo *et al*[99] when CE was preferred over other modalities in OGIB and turned out to be positive. However, only reimbursement costs were evaluated, so the cost of the hospital and the personnel was not taken into account.

If a therapeutic intervention is needed with a probability of more than 25%, gastroenterologists should consider the use of DBE as initial therapeutic option to minimize costs[93]. Furthermore, cost equalization of DBE and CE was reached at 100 procedures for diagnostic DBE and 79 procedures for therapeutic DBE, which suggests that DBE is especially cost-effective in large-scale hospitals, with a substantial number of DBE procedures per year. Another study by Gerson *et al*[100] found that, regardless of the cost, DBE procedure was more cost-effective than CE-guided DBE procedure, because no additional costs were charged regarding further examinations and therapy could be given instantly. However, the workload for physicians would significantly increase if an initial DBE would be done and we may not forget that DBE is correlated with a higher rate of complications compared to CE. CE-guided DBE was associated with better outcomes in the long term because of fewer potential complications and fewer utilization of endoscopic resources. This can be explained by the high negative predictive value of capsule endoscopy, which leads to a reduction in the subsequent DBE procedures[92,100].

When CE is negative, the chance of rebleeding is low, so that further investigations can be deferred, even when a second test might be diagnostic[101-103]. Rebleeding was reported to be higher in CE-positive patients and patients using anticoagulants[103]. Nonetheless, gastroenterologists should consider close monitoring, alternative modalities in suspicious cases because the chance of rebleeding has been reported up to 28.4% and 35.3% during a median follow-up of respectively 23.7 mo and 31.7 mo[104,105]. Repeating the CE procedure however should only be considered if the bleeding presentation switches from occult to overt bleeding or the hemoglobin level drops with more than 4g/dL[106]. Diagnostic yield of a repeated CE was reported to be between 35 and 75 percent and a subsequent management change was reported in 39 to 62.5 percent of the patients[107,108].

To conclude this chapter about OGIB, we have made a flow-chart to represent the current knowledge in this field. For this purpose we included the reviewed articles in previous flow charts[18,109] (Figure 3)

***CD***

Non-stricturing CD is the second main indication for CE. CD is a type of chronic IBD which may affect the whole gastrointestinal tract and lead to mucosal and transmural damage. Categorization of patients with CD is done based on the disease presentation: solely the small bowel (30-35% of the patients), the small bowel and the large bowel (45-50%) or only the large bowel (20%)[110]. So even though it primarily affects the terminal ileum, the ileocecal region and the large bowel, one third of the patients presents with only small bowel inflammation which challenges gastroenterologists to diagnose the disease. Traditionally small bowel involvement was diagnosed by radiological procedures, small bowel barium radiography, CT, colonoscopy with ileoscopy or enteroscopy. But with the invention of CE, new possibilities in CD diagnostics have become available.

CE can be very helpful in the diagnosis of new cases of Crohn and in the evaluation of known CD, with regard to the activity and extent of the disease. CE is reserved however for cases with unexplained symptoms, when other investigations remain inconclusive or when CE would affect the management of the patient[111]. So both in suspected as established CD, CE usually is performed third after a negative colonoscopy and ileoscopy, thereby replacing the traditional modalities. Capsule endoscopy is considered positive for CD when more than 3 ulcerations are identified in the absence of NSAID use[111-113] or when 4 or more obvious clear ulcers, erosions, or a region with clear exudate and mucosal hyperemia and edema are seen[114].

Like in OGIB, CE also has shown a superior yield for detecting early inflammatory lesions in the small bowel comparing to all other modalities as can be seen in Table 3. The yield of CE in non-stricturing CD has been reported to be between 18% and 96%[81,95,115-119]. Triester *et al*[81] only found a significant difference in yield between CE and other modalities for diagnosing non-stricturing small bowel CD. However, a distinction should be made between suspected and established CD. The reported superior yield was only significant for evaluating established CD and was not reported for diagnosis of small bowel CD in patients with a suspected initial presentation of the disease. This was contradicted by Dionisio *et al*[115], who found that CE has a superior yield compared to Small Bowel Radiography (SBR), CT enterography (CTE) and colonoscopy with ileoscopy in the diagnosis of suspected CD patients. Although the yield of capsule endoscopy in CD is high, the proportion of CD patients diagnosed with CE is rather low (0-4%). Only in young patients presenting with abdominal pain plus diarrhea a 30% chance of diagnosing CD was achieved[114]. In established CD patients, CE was reported to be superior compared to SBR, CTE and PE, which was the same according to previous findings[115]. When compared to CT enterography and MR enterography, CE shows superior yields in the first two-thirds of the small bowel, but loses this superiority in the last portion of the small bowel by showing a yield similar to the comparators[120,121]. However, we may not forget that MR enterography is also capable of visualizing the small bowel surroundings, so that transmural and extra-intestinal manifestations can be diagnosed[122]. A recent study by Leighton *et al*[123] found that a combination of colonoscopy with ileoscopy and CE achieved a far more high yield than patients investigated with a combination of colonoscopy with ileoscopy and small-bowel follow-through (SBFT). They confirmed the role of CE as valuable third diagnostic option in diagnosis of suspected CD, when colonoscopy and ileoscopy turned out to be negative or inconclusive.

In patients with suspected CD, Girelli *et al*[119] found that, presuming a pre-test probability of CD of 50%, a positive CE was capable to raise the post-test probability up to 85% and if the CE was negative, it was capable to lower the probability to only 5%. In patients with established CD, the use of small bowel CE in monitoring therapy response is still a controversial issue. Many reports found that the clinical and biological response to treatment is not correlated with mucosal healing, which is monitored on CE, so it has not proven useful in this respect[124].

Caution should be taken when evaluating CEs positive for small bowel lesions. Because of the potential of CE to detect early lesions, CD-induced lesions are often non-specific and can be confused with NSAID-induced lesions. Both CD and NSAID-induced small bowel injury show endoscopically similar lesions and because of the inability of CE to take biopsies, the differential diagnosis remains inconclusive. Pathognomonic however for NSAID-induced lesions are the concentric diaphragmatic strictures in the ileum seen on endoscopy, which can lead to small bowel obstruction[125]. According to Doherty *et al* the problem of false positive capsules also overestimates the incremental yield of capsule endoscopy compared to other modalities, necessitating a diagnostic golden standard to overcome the problem of premature CD diagnoses. Currently, there are two scores available to assess and monitor mucosal disease activity on CE. The Capsule Endoscopy CD Activity Index (CECDAI or Niv score) and the Lewis score are only recently developed and still have to prove their usefulness in standardizing the diagnosis of CD on CE before being widely accepted in clinical practice as an objective tool of mucosal inflammation measurement[126,127]. Although the yield of CE has been questioned by these diagnostic problems, CE still remains a valuable tool in the diagnosis of CD: a recent study by Hall *et al*[128] found a very high negative predictive value in the long term despite the questioned yield in patients with suspected CD, which makes it capable of safely ruling out suspected CD.

As mentioned before, capsule retention is especially feared in patients with CD because of possible strictures and stenosis. The reported 2.6% by Sharaf *et al*[129] has made small bowel imaging a standard exam previous to the CE procedure[19]. MR is a useful tool to asses patency of the small bowel[130]. Another possibility is the use of the previously discussed Pillcam Patency Capsule (Given Imaging, Yoqneam, Israel), which indicates patency if the capsule is excreted intact or the scan has lost the RFID tag signal 30 h after ingestion[131], so the CE procedure can be done to evaluate the mucosal surface of the small bowel.

Cost analyses for CE in CD have been made and showed that colonoscopy with ileoscopy followed by a CT enterography was the most cost-effective choice among the different diagnostic options in patients suspected of CD[132]. Moreover, CE was proven to be not cost-effective as third diagnostic option, because of the high false positive rate, the diagnostic yield and the low pre-test probability of CD. Sharaf *et al*[129] confirmed these findings and concluded that CE is not a valuable option in patients with suspected CD. However, Leighton *et al*[133] found that CE did play a significant role in early diagnostics of CD, because it did not necessitate repeated procedures, physician visits and hospital stays, so direct costs could be reduced. Further investigation on this matter is needed.

In summary, CE has a superior diagnostic yield when compared to other modalities in suspected as well as established small intestinal CD. However, the question if this superior yield is due to false positive results remains unanswered. With the development of two scoring systems, this problem might be solved in the near future. Still, CE is a promising tool in CD diagnostics because of its capability to early diagnose small bowel lesions. We conclude with a flow chart based on the ICCE flowchart[130] with incorporation of new evidence[124,128] (Figure 4).

***Surveillance of polyposis syndromes and detection of small bowel tumors***

Small bowel tumors make up only 3-6% of the gastrointestinal neoplasm cases despite the 90% of the gastrointestinal tract surface the small bowel covers, which makes it a difficult entity to diagnose[134,135]. The most frequently observed tumors are adenocarcinoma, gastrointestinal stromal tumor, carcinoid, lymphoma and sarcoma[136-139]. Symptoms are rather unspecific and include anaemia or overt OGIB and later abdominal pain, nausea, vomiting, weight loss and anorexia[140,141]. Thereby tumors are mostly found on CE or DBE when investigating patients with OGIB [80,138,139,142]. A study by Sîngeap *et al*[136] reported a detection rate for small bowel tumors of 4.9% in patients presenting with OGIB or other nonspecific symptoms. Other studies have found a tumor detection rate of 6-12% on CEs done for OGIB[143]. The insidious process often is responsible for the delayed diagnosis of a patient, which impacts the further management of the patient[144]. Fast tumor detection is therefore very important, since management can be changed accordingly and outcomes can be improved even in malignant lesions if metastasis is absent[145]. Small bowel tumors can be benign, potentially malignant, malignant or metastatic. However the majority, 60%, of these tumors are malignant[143], and differentiation between benigna and malignant cannot be made on CE. Tumors mostly appear as masses or polyps, but also can present as ulcers and stenoses in a minority of the cases. Hereditary polyposis syndromes like Familial Adenomatous Polyposis (FAP) and Peutz-Jeghers Syndrome (PJS), are another entity and apart from the colon polyposis, patients often develop benignant small bowel pathology with a high tendency to evolve into cancer[146].

Capsule endoscopy was evaluated for small bowel tumors and hereditary polyposis syndromes and turned out to be a valuable diagnostic tool[147,148]. The pooled detection rate of CE was 55.9%[19]. Therefore, In patients with suspected small bowel tumors, CE can be the first choice in diagnostics[136]. In a study by Schulmann *et al*[146] it was stated that CE was capable of detecting small bowel polyposis, located in the distal jejunum and ileum beyond the reach of PE. These polyps could subsequently be removed by DBE, so surgery was avoided. However, most FAP patients with distal polyposis also presented with proximal polyposis, which was equally detected by CE as well as PE. Proximal jejunal polyposis is significantly correlated with the presence and severity of duodenal disease, which is one of the main locations for adenocarcinoma and subsequent mortality. Because CE was capable of detecting proximal small bowel polyposis and given its superior sensitivity and non-invasive nature, it was recommended as a surveillance tool in a subgroup of FAP patients with severe duodenal polyposis[146]. Duodenal polyposis itself is difficult to detect by CE, due to the rapid transit of the capsule in this part of the gastrointestinal tract. Another study by Plum *et al*[149] confirmed the superiority of CE compared to other modalities such as PE, ileoscopy and enteroclysis in patients with FAP. However, they also stressed on the fact that CE did not replace the other modalities, because CE sometimes missed lesions and did not manage to precisely localize the small bowel lesions. Also a study by Wong *et al*[150] confirmed the fact that CE could underestimate the number of small bowel polyps in FAP and a review by Koornstra[151] stated that CE cannot totally replace standard endoscopy in the surveillance of the proximal small bowel. A tool to overcome missed lesions might be the recently developed CICE tool, which enhances the contrast of the CE images and thereby improves the visibility in patients with small bowel polyposis. Although further evaluation is needed, a first trial showed that half of the adenomatous polyps could be better visualized and hamartomatous polyposis was better visible[152].

In PJS, CE was capable of detecting lesions with direct impact on further management. CE is the most accurate diagnostic tool to detect small bowel polyposis throughout the whole small bowel and can be seen as a safe alternative for the traditional modalities, such as PE and MR enteroclysis used in PJS and FAP[146]. The superiority of CE over MR enterography was also confirmed by Liao *et al*[19] who found that CE was capable of detecting smaller small bowel lesions. Urgesi *et al*[142] stated that CE could detect more lesions than the traditional endoscopy and radiological imaging in patients suspected for small bowel tumors. They concluded that CE played an important role in the diagnostic work-up of these patients[142]. Similarly another very recent study by Urquhart *et al*[153] found that CE was able to identify significantly more small bowel polyps compared with MRE in patients with PJS. Furthermore, Rahmi *et al*[154] found that CE was also useful in planning the DBE approach in patients in need of polypectomy. DBE, which achieves a similar yield as CE, is useful when biopsy, exact pre-operative localization or local therapy, such as stenting or balloon dilatation is needed[155].

CE also has its limitations in the detection of small bowel tumors. First of all it is not useful in an emergency setting, such as obstruction and peritonitis, because of the risk of capsule retention[144]. Furthermore, CE is not capable of treating locally or taking biopsies, needed to differentiate between benignant and malignant[156]. Finally CE is not able to differentiate a mucosal bulge from a smooth-walled tumor. To overcome the latter problem, the scoring system SPICE (Smooth, Protruding lesion Index on Capsule Endoscopy) has been developed. A score greater than 2 is suggestive of small bowel malignancy, but further validation is needed[157].

Just like in CD, the risk of capsule retention is present. Yet the rate is lower in patients with intestinal tumors compared to patients with CD[19]. Moreover, Bailey *et al*[145] stated that obstructions due to neoplasms were a positive complication because, since the tumor anyway needed to be treated by enteroscopy or surgery, the impacted capsule could serve as a guide. Like in CD, if the patient is suspicious for obstruction, imaging should be done before CE. Management of malignant small bowel tumors is primarily surgical. In selected cases, this can be performed laparoscopically. Adjuvant chemotherapy and radiotherapy may be needed, depending on the histology of the tumor[134].

We can conclude by stating that CE is a diagnostic tool with a big value regarding its yield in diagnosis and surveillance of small bowel tumors/polyps. However, it is complementary to the traditional modalities and can not substitute them. CE is recommended third after negative bidirectional endoscopy in patients with OGIB or other unspecific symptoms indicating a possible small bowel tumor. It can be used first as a complementary diagnostic tool in patients with established hereditary polyposis syndromes. We summarized the evidence in two flow-charts based on the Consensus statements for small−bowel capsule endoscopy, 2006/2007 by Mergener *et al*[109] and a study by Plum *et al*[149] (Figure 5 and 6).

***NSAID side-effects***

NSAIDs can inflict injury along the whole gastrointestinal tract, when used for a prolonged time. Although many publications have emphasized on the incidence of upper gastrointestinal lesions, fewer have mentioned lower gastrointestinal ones. However, as mentioned before, NSAIDs can also induce small bowel lesions, which can be observed on CE. In fact, these lesions are far more common than the NSAID-induced gastropathy[158]. Furthermore, complications in the lower gastrointestinal tract, such as perforation, bleeding, or obstruction are currently increasing while upper GI complications are decreasing[159], which necessitates the need of small bowel diagnostics in the field of NSAID side-effects.

In seventy percent of the patients using NSAIDs continuously, mucosal damage of the small intestine has been reported on CE or DBE[160,161]. Even a two-week NSAID-regimen with slow-release diclofenac resulted already in macroscopic injury of the small intestine in 68%-75% of the volunteers[162]. Different types of lesions have been observed ranging from mucosal redness and multiple petechiae to erosions, ulcers, loss of villi, diaphragm-like strictures, which are pathognomonic for NSAID-induced enteropathy, and even severe bleeding[163,164]. Most symptomatic patients present with OGIB with or without obstruction symptoms and are accordingly diagnosed[165,166].

Both CE and DBE have been evaluated for NSAID-induced lesions. They show a similar yield of 60% in diagnosis[165,167]. CE however is preferred for screening of NSAID-induced lesions and evaluation of further treatment because of its non-invasive character. DBE on the other hand is the first choice in patients suspicious of strictures. NSAID-use has been recognized as a risk factor for capsule retention and CE should therefore be avoided in these patients[19]. Furthermore DBE is preferred when further examination of the lesion, endoscopic or histologically, is needed or when local therapy has to be given, such as balloon dilatation of a stricture or endoscopic coagulation, clipping or injection of the bleeding site. Balloon dilatation of a stricture seems to be safe, since the muscularis propria remains intact and perforation is subsequently rarely observed[155]. A recent study by Tacheci *et al*[168] confirmed the high sensitivity of capsule endoscopy and further stated that subclinical small bowel damage also could be observed on CE. If NSAID enteropathy is found on CE or DBE, further investigation can be done using other modalities such as radiological examination, the permeability test, scintigraphy, the fecal excretion with 111In white blood cells and measurement of the calprotectine concentration in the faeces[163].

Just like in CD, scoring systems are available to classify lesions and to consider and evaluate further treatment[126,160,162,167]. However, no standard scoring system is thoroughly evaluated. Different therapeutic options are available. The first choice of therapy is a discontinuation in the use of NSAIDs, which in most cases is not possible due to the underlying pathology[160]. Cyclooxygenase-2 (COX-2) selective inhibitors, prostaglandin derivatives, a combination of NSAIDs and phosphatidylcholine, cytoprotective drugs and probiotics are all useful for the treatment of NSAID-induced small intestinal injuries[158]. Yet controversy remains around the use of selective COX-2 agents. A trial by Goldstein *et al*[169] reported that a 2-wk regimen of selective COX-2 agents caused fewer small intestine injuries than treatment with a nonspecific NSAID. This was confirmed in a big RCT by Chan *et al*[170]. However, Maiden *et al*[171] showed that COX-2 selective inhibitors caused the same amount of small bowel damage as long-term NSAIDs, which is interesting given the fact that they affect the gastroduodenal mucosa to a lesser extent[169]. So COX-2 might play a significant role in the maintenance of the small bowel integrity. We can conclude that it remains unclear whether selective COX-2 inhibitors truly prevent NSAID-induced enteropathy.

Also chronic Low-dose aspirin (LDA) users are at risk of small bowel enteropathy. The phenomenon was first described by Leung WK *et al*[172]in 2007 and the study by Endo *et al*[164] was the first to report the characteristics of the small bowel damage associated with long-term LDA use. The use of LDA however was less harmful than other types of NSAIDs[165]. These findings may have implications on treatment of the large group of patients requiring anti-inflammatory or antithrombotic drugs.

***Celiac disease***

Celiac disease is caused by an chronic auto-immune response of the intestines to gliadins in the diet and occurs in approximately 1% of the population in genetically susceptible persons[173]. It is characterized by duodenal folds, scalloping of folds, mucosal fissures, crevices or grooves, visible submucosal vessels, micronodules in the duodenal bulb and a mosaic pattern in the small bowel mucosa[174]. The lesions are visible on CE, which makes CE therefore a perfect tool to assess small bowel damage in these patients. In a large meta-analysis by Rokkas *et al*[175] sensitivity and specificity of CE in Celiac Disease have been reported up to 89% and 95% respectively. This was similar to a previous meta-analysis by El-Matary *et al*[176] which reported a sensitivity of 83% and a specificity of 98%. However, to confirm Celiac Disease in patients with positive serologic markers, a biopsy is needed, which is not possible with the current capsules. Therefore, the gold standard for the diagnosis of Celiac Disease remains the histological findings of a small bowel specimen obtained through gastroduodenoscopy.

The main indications to use CE are serological positive patients, who are unwilling to undergo gastroduodenoscopy or patients with antibody-negative villous atrophy. The latter group showed a higher yield on CE, compared with CE in serological positive patients with biopsy-proven Celiac Disease and persisting symptoms as was found by Kurien *et al*[177]. Also in patients with non-responsive Celiac Disease, defined by persistent or recurrent symptoms under treatment with a gluten-free diet, CE showed to be of use to detect complications, such as multiple erosions, ulcerations, ulcerative jejunitis and adenocarcinoma[178]. Tennyson *et al*[179]confirmed these findings, but emphasized that CE was not a necessary tool in the evaluation of non-responsive Celiac Disease when no alarm symptoms are present, such as weight loss and abdominal pain, or when no loss of T-cell antigens on intraepithelial lymphocytes or loss of clonality of the T-cell receptor gene was observed. In the latter situations, a combination of CE and CT or MR enterography should be performed. In all other cases, upper gastroduodenoscopy with biopsy remains the gold standard. A recent study by Van Weyenberg *et al*[180] found similar results stating that CE could be used in patients with non-responsive Celiac Disease to identify the cases who are at risk of complications. Capsule endoscopy might also be usefull in the follow-up of patients with celiac disease under treatment with a gluten-free diet, regarding mucosal healing, because the follow-up of duodenal histology is not representative for the mucosal healing more distally[181]. Finally, Akin *et al*[182] confirmed other authors by stating that CE was useful as an alternative to duodenal biopsy in patients unable or unwilling to undergo gastroduodenoscopy and further stated that CE could be of use in the diagnosis of Celiac Disease in elderly patients with unspecific symptoms.

In conclusion, gastroduodenoscopy remains the diagnostic tool of choice for celiac disease, but CE shows to be a useful adjunctive tool in specific situations.

**UPCOMING CHALLENGES**

To conclude this paper about the current knowledge of capsule endoscopy, we would like to offer an insight in its bright future. Since its release in 2001, optics, battery life, visualization and software have been improved, with consequences on yield, completion rate and reading time. We have already discussed some technological advances in CE, but we will now shortly focus on future expectations of this technology.

One major field of advancement will be the maneuverability. If a capsule endoscope would be steerable and could approach a site of interest, this could be a big step forward in the diagnosis and treatment of diseases of the whole gastrointestinal tract. With efficient movement, battery life could sustain during movement through the whole gastrointestinal tract and thereby could increase completion rate. Various studies have been done and many prototype active capsules, using different locomotion techniques are currently under investigation for human use[183-187]. However, in the near future, remote manipulation using magnetic forces will be the first to be commercially available. These capsules contain a magnet, which can then be mobilized with an externally handled magnetic paddle or with a joystick. Perspective of the camera also can be adjusted with this magnet, rendering the desired image[188]. Swain *et al*[189] was the first to document the use of a magnetic field to guide a capsule through the human oesophagus and stomach. Since this article, many studies have followed, especially focusing on investigation of the stomach[190-193]. To overcome the problem of capsule impaction and to improve mucosal visualization, especially in the colon, insufflation techniques have recently been described by several authors using a capsule with a magnetic controlled drug release system to create a basic chemical reaction forming CO2 in the lumen[194-196]. Another very interesting topic is a novel wireless platform able to measure and locate the force opposing capsule motion as a reflection of the gastrointestinal tract resistance[197]. It is the first platform for magnetic control of CEs that implemented this intermagnetic force measurement feature.

Another advancement might be the availability of a controlled drug release feature. This could help gastroenterologists in the local treatment of various gastrointestinal diseases, such as medical treatment of CD or even hemostasis in OGIB. Only one study introduced a capsule able to microposition a needle and to deliver 1 mL of a targeted medication, while resisting peristalsis with its holding mechanism[198].

The inability of taking biopsies is a third challenge CE faces. Together with an accurate maneuverability, this could enable CE to completely take the place of DBE in diagnostic and even therapeutic endoscopy of the small bowel. In 2008, Valdastri *et al*[199] was the first to successfully report an in vivo experiment with a capsule with built-in clip-releasing mechanism. The VECTOR project by the European Commission is currently developing a capsule for diagnosis and treatment of gastrointestinal cancer[200]. Another study also investigated the use of a large number of thermo-sensitive microgrippers in capsule endoscopy for this purpose of grabbing and retrieving tissue samples, which showed promising results[201].

Finally, to end this paper, we would like to reflect on the environmental consequences of capsule endoscopies, a subject that will become more important in the future, given the growing importance of CE. Although the ASGE consensus states that the capsule is ‘disposable and designed to be excreted’[3], considerations around this topic should be made, since the capsule contains many particles with potential biohazards[202]. Pezzoli *et al*[203] was the first to publish a small article on this matter in in 2011. They found that it was possible, after retrieval and cleaning, to reactivate used capsules with a 10 min procedure and a new battery cost of only 2 euro. Recycled capsules could then be given a second life in *e.g.* veterinary procedures[203].

**CONCLUSION**

This paper gives a brief but complete overview on small bowel CE anno 2014. As the technology is still evolving and new insights are still being published every year, we emphasize that healthcare-providers should continue to monitor the medical literature for recent data, in order to provide the best evidence-based care for their patients.

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**Table 1 Comparison between commercially available capsule endoscopy devices**

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| Capsule | PillCam® SB 3Given Imaging | EndoCapsule®Olympus America | MiroCam®Intromedic Company | OMOM®Jianshan Science and Technology |
| Size | Length: 26.2 mmDiameter: 11.4 mm | Length: 26 mm Diameter: 11 mm | Length: 24.5 mmDiameter: 10.8 mm | Length: 27.9 mmDiameter: 13 mm |
| Weight | 3.00 g | 3.50 g | 3.25 – 4.70 g | 6.00 g |
| Battery Life | 8 h or longer | 8 h or longer | 11 h or longer | 6-8 h or longer |
| Resolution | 340 x 34030% better than SB2 | 512 x 512 | 320 x 320 | 640 x 480 |
| Frames per second | 2 fps or 2-6 fps | 2 fps | 3 fps | 2 fps |
| Field of View | 156° | 145° | 170° | 140° |
| Communication | Radio Frequency Communication  | Radio Frequency Communication | Human Body Communication  | Radio Frequency Communication  |
| FDA approval | Yes | Yes | Yes | No |
| Price per capsule | $500 | $500 | $500 | $250 |

**Table 2 Comparison of different diagnostic modalities in obscure gastrointestinal bleeding**

|  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| Ref. | Country | Design | number of patients | Comparator | Yield of CE, n (%) | Yield of Comparator, n (%) | Significant difference? (yes/no) | CE superior? (yes/no) | Other |
| Triester *et al*[80] | United States | Meta-analysis and Systematic review | 396 | PE | 63 | 28 | yes | yes | NNT = 3 to yield one additional clinically significant finding with CE |
|  |  |  | 88 | SB radiography (barium contrast and enteroclysis) | 67 | 8 | yes | yes | NNT = 3 to yield one additional clinically significant finding with CE |
| Leighton *et al*[82] | United States | Meta-analysis and Systematic review | 396 | PE | 63 | 28 | yes | yes | Yield of significant findings: CE = 56% vs PE = 26%, NNT = 3 to yield one additional clinically significant finding with CE  |
|  |  |  | 88 | Barium radiography | 67 | 8 | yes | yes | Yield of significant findings: CE = 42% *vs* SB barium radiography = 6%, NNT = 3 to yield one additional clinically significant finding with CE  |
|  |  |  | 42 | Intraoperative enteroscopy | 83 | 83 | no | no |  |
|  |  |  | 17 | Mesenteric angiography | 47 | 53 | no | no |  |
| Chen *et al*[204] | China | Meta-analysis and Systematic review | 277 | DBE | 61 | 56 | no | no | CE was superior if no combination of oral + anal approach <-> DBE was superior when a combination of the two insertion approaches was done |
| Pasha *et al*[95] | United States | Meta-analysis and Systematic review | 397 | DBE | 24 | 24 | no | no | CE should be the initial diagnostic test for determining insertion route of DBE |
| Arakawa *et al*[76] | Japan | Retrospective Study | 162 | DBE | 54 | 64 | no | no |  |
| Teshima *et al*[77] | Canada, Netherlands | Meta-analysis and Systematic review | 651 | DBE | 62 | 56 | no | yes | yield of DBE after positive CE = 75,0% <-> yield after negative CE = 27,5% |
| Leung *et al*[205] | China | RCT | 60 | Mesenteric angiography | 53 | 20 | yes | yes | no significant difference in the long-term outcomes (transfusion need, hospitalization for rebleeding,mortality) |
| Wang *et al*[206] | China | Meta-analysis and Systematic review | 279 | CT | 53 | 34 | yes | yes | complementary role to CE and can be used as a triage tool prior to DBE in evaluating OGIB  |

PE: Push Enteroscopy; CE: Capsule Endoscopy; SB: Small bowel; NNT: Number needed to treat; DBE: Double balloon enteroscopy; RCT: Randomized controlled trial.

**Table 3 Comparison of different diagnostic modalities in Crohn’s Disease**

|  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| Ref. | Country | Design | number of patients | Comparator | Yield of CE, *n* (%) | Yield of Comparator, n (%) | Significant difference? (yes/no) | CE superior? (yes/no) | Other |
| Marmo *et al*[117] | Italy | RCT | 31 | SB radiography (enteroclysis) | 71 | 26 | Yes | Yes | Terminal ileum: yield 89% *vs* 37%Proximal SB: yield only 46% *vs* 13% |
| Chong *et al*[207] | Australia | Blinded prospective trial | 43 | SB enteroclysisPE | 7777 | 1914 | YesYes | YesYes | Results are in patients with a history of CD |
| Triester *et al*[81] | USA | Meta-analysis and Systematic review | 250 | SB barium radiography | 63 | 23 | Yes | Yes | NNT = 3 to yield one additional diagnosis with CE |
|  |  |  | 114 | colonoscopy with ileoscopy | 61 | 46 | Yes | Yes | NNT = 7 to yield one additional diagnosis with CE |
|  |  |  | 93 | CT enterography/CT enteroclysis | 69 | 30 | Yes | Yes |  |
|  |  |  | 84 | PE | 46 | 8 | Yes | Yes |  |
|  |  |  | 18 | MR enterography | 72 | 50 | No | yes |  |
| Solem *et al*[118] | United States | Blinded prospective trial | 41 | CT enterographyColonoscopy with ileoscopySmall bowel follow-through | 838383 | 837465 | NoNoNo | NoYesYes | specificity of CE (53%) was significantly lower than the other tests |
| Pasha *et al*[95] | United States | Meta-analysis and Systematic review | 343 | DBE | 18 | 16 | no | no |  |
| Dionisio *et al*[115]  | United States | Meta-analysis and Systematic review | 428 | SB barium radiography | 52 | 16 | Yes | Yes |  |
|  |  |  | 236 | colonoscopy with ileoscopy | 47 (71)1 | 25 (36) | Yes | Yes | Suspected CD (Established CD) |
|  |  |  | 119 | CT enterography | 68 (71) | 21 (39) | Yes | Yes | Suspected CD(Established CD) |
|  |  |  | 102 | PE | 66 | 9 | Yes | Yes | Established CD |
|  |  |  | 123 | MR enterography | 55 (70) | 45 (79) | No | Yes (no) | Suspected CD (Established CD) |
| Lu *et al*[116] | China | Retrospective Study | 50 | colonoscopy with ileoscopy | 96 | 66 | Yes | Yes | Combination of two methods showed a higher yield, but no significant differences were reported between each two examinations |
|  |  |  | 34 | CT enterography | 96 | 85 | Yes | Yes |  |
|  |  |  | 39 | Small bowel follow-through | 96 | 67 | Yes | Yes |  |

1Extra information between brackets is specific for Established Crohn’s Disease. PE: Push Enteroscopy; CE: Capsule Endoscopy; SB: Small bowel; NNT: Number needed to treat; DBE: Double balloon enteroscopy; RCT: Randomized controlled trial.

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**Figure 1 Main parts of the wireless capsule endoscope.**



**Figure 2 Types of small bowel capsule endoscopes.** A: PillCam SB 3 (Given Imaging, Yoqneam, Israel); B: MiroCam (IntroMedic, Seoul, Korea); C: Endo Capsule (Olympus America, Center Valley, PA); D: OMOM (Jinshan Science and Technology, Chongqing, China).



**Figure 3 Recommended approaches for diagnosis and treatment of Obscure Gastrointestinal Bleeding (OGIB).** 1DBE is recommended in (1) patients requiring a biopsy or a therapeutic intervention (2) patients with an initial negative CE and suspicion of a small bowel lesion (3) large scale hospitals or (4) hospitals where CE is not available; 2If a lesion is suspected, further work-up is needed; 3Rebleeding was defined as a change from occult to overt bleeding or a hemoglobin level drop more than 4g/dL. DBE: Double Balloon Enteroscopy; PE: Push Enteroscopy; IOE: Intraoperative Enteroscopy; Routine endoscopy: Uni/bidirectional endoscopy.



**Figure 4 Recommended approaches for diagnosis and treatment of Crohn’s Disease (CD).** 1Non-invasive markers have proven to be useful in giving baseline information about the presence of small bowel inflammation; 2If Capsule Endoscopy (CE) is negative, Crohn’s Disease can be ruled out due to the high negative predictive value of CE. In that case, other explanations should be considered. CRP: C reactive protein.



**Figure 5 Recommended approaches for diagnosis and treatment of Small bowel tumors.** DBE: Double Balloon Enteroscopy; PE: Push Enteroscopy.



**Figure 6 Recommended approaches or diagnosis of Small bowel hereditary polyposis.** DBE: Double Balloon Enteroscopy; PE: Push Enteroscopy.