Small-bowel capsule endoscopy and device-assisted enteroscopy for diagnosis and treatment of small-bowel disorders: European Society of Gastrointestinal Endoscopy (ESGE) Guideline – Update 2022

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MAIN RECOMMENDATIONS

MR1 ESGE recommends small-bowel capsule endoscopy as the first-line examination, before consideration of other endoscopic and radiological diagnostic tests for suspected small-bowel bleeding, given the excellent safety profile of capsule endoscopy, its patient tolerability, and its potential to visualize the entire small-bowel mucosa.
Strong recommendation, moderate quality evidence.

MR2 ESGE recommends small-bowel capsule endoscopy in patients with overt suspected small-bowel bleeding as soon as possible after the bleeding episode, ideally within 48 hours, to maximize the diagnostic and subsequent therapeutic yield.
Strong recommendation, high quality evidence.

MR3 ESGE does not recommend routine second-look endoscopy prior to small-bowel capsule endoscopy in patients with suspected small-bowel bleeding or iron-deficiency anemia.
Strong recommendation, low quality evidence.

MR4 ESGE recommends conservative management in those patients with suspected small-bowel bleeding and high quality negative small-bowel capsule endoscopy.
Strong recommendation, moderate quality evidence.

MR5 ESGE recommends device-assisted enteroscopy to confirm and possibly treat lesions identified by small-bowel capsule endoscopy.
Strong recommendation, high quality evidence.

MR6 ESGE recommends the performance of small-bowel capsule endoscopy as a first-line examination in patients with iron-deficiency anemia when small bowel evaluation is indicated.
Strong recommendation, high quality evidence.

MR7 ESGE recommends small-bowel capsule endoscopy in patients with suspected Crohn’s disease and negative ileocolonoscopy findings as the initial diagnostic modality for investigating the small bowel, in the absence of obstructive symptoms or known bowel stenosis.
Strong recommendation, high quality evidence.

MR8 ESGE recommends, in patients with unremarkable or nondiagnostic findings from dedicated small-bowel cross-sectional imaging, small-bowel capsule endoscopy as a subsequent investigation if deemed likely to influence patient management.
Strong recommendation, low quality evidence.

MR9 ESGE recommends, in patients with established Crohn’s disease, the use of a patency capsule before small-bowel capsule endoscopy to decrease the capsule retention rate.
Strong recommendation, moderate quality evidence.

MR10 ESGE recommends device-assisted enteroscopy (DAE) as an alternative to surgery for foreign bodies retained in the small bowel requiring retrieval in patients without acute intestinal obstruction.
Strong recommendation, moderate quality evidence.

MR11 ESGE recommends DAE-endoscopic retrograde cholangiopancreatography (DAE-ERCP) as a first-line endoscopic approach to treat pancreaticobiliary diseases in patients with surgically altered anatomy (except for Billroth II patients).
Strong recommendation, moderate quality evidence.
Conclusion

The introduction of small-bowel capsule endoscopy (SBCE) and device-assisted endoscopy (DAE) over 20 years ago marked the beginning of a new era for investigating the small intestine. There is now more solid scientific evidence on established indications, and more data on new applications of enteroscopy are available. The aim of this Guideline, commissioned by the European Society of Gastrointestinal Endoscopy (ESGE) as an update of the previous 2015 Guideline [1], is to provide guidance for the clinical application of enteroscopy techniques in the management of adult patients with small-bowel (SB) disorders.

Methods

ESGE commissioned this clinical Guideline (ESGE Guideline Committee Chair, K.T.) and appointed a guideline leader (M.P.) who formed a coordinating team (M.P., E.R., P.C.V.). The guideline leader established six task forces, each with its leader (C.S., E.D., M.K., D.S.S., T.M., X.D.). Key questions were prepared by the coordinating team according to the PICO (patients, interventions, controls, outcomes) format and divided among the six task forces (see Table 1s, Key Questions, available online-only in Supplementary Material). Given that this is an update of the 2015 ESGE Clinical Guideline [1], each task force performed a structured, systematic search, using keywords, for available literature (English-language articles) from December 2014 to November 30 2021 in Ovid MEDLINE, EMBASE, Google Scholar, and the Cochrane Database of Systematic Reviews; the literature search was then updated up to April 1 2022, to look for recently released papers. A dedicated manual search was also performed in the same timeframe by checking references of relevant papers. The hierarchy of studies included in this evidence-based guideline was, in decreasing order of evidence level: published systematic reviews/meta-analyses, randomized controlled trials (RCTs), prospective and retrospective observational studies, and case series.
Evidence on each key question was summarized in tables (Table 2, Evidence tables), using the Grading of Recommendations Assessment, Development and Evaluation (GRADE) system, wherever applicable [2]. The evidence grading depends on the balance between any health intervention’s benefits and their risk or burden. Further details on ESGE guideline development are available elsewhere [3].

The literature search results and answers to PICO questions were presented to all guideline group members during an online meeting on October 8, 2021. Subsequently, drafts for each topic were prepared by each task force leader and distributed between the task force members for revision and discussion. In June 2022, a draft prepared by the coordinating team, including all the statements, was sent to all guideline group members. All the statements were discussed and modified in real time, if necessary, during an online meeting on June 24, 2022. After the agreement of all members was obtained, the manuscript was reviewed by two independent external reviewers. The manuscript was then sent to the 51 ESGE member societies and to individual members for further comments. The final revised manuscript, having been agreed upon by all authors, was submitted for publication to the journal Endoscopy.

This ESGE Guideline was issued in 2022 and will be considered for update in 2027. Any interim updates will be noted on the ESGE website: http://www.ESGE.com/esge-guidelines.html.

Evidence statements and Recommendations

Evidence statements and Recommendations are grouped according to the different task force topics: suspected small-bowel bleeding (SSBB) and iron-deficiency anemia (IDA) (task force 1), Crohn’s disease (CD) (task force 2), small-bowel tumors (SBTs) and inherited polyposis syndromes (task force 3), celiac disease (task force 4), other indications (task force 5), and innovations (task force 6). Each statement is followed by the assessment of the strength of evidence, based on GRADE. Table 1 summarizes all recommendations in this updated Guideline.

**Suspected small-bowel bleeding**

**RECOMMENDATION**
ESGE recommends small-bowel capsule endoscopy as the first-line examination, before consideration of other endoscopic and radiological diagnostic tests, for suspected small-bowel bleeding, given the excellent safety profile of capsule endoscopy, its patient tolerability, and its potential to visualize the entire small-bowel mucosa.

Strong recommendation, moderate quality evidence.

Small-bowel (SB) bleeding is defined as bleeding in the gastrointestinal (GI) tract between the ampulla of Vater and the ileocecal valve. SB bleeding is suspected when a patient presents with GI bleeding but has negative upper and lower endoscopy findings; it can present as overt or occult bleeding. The term “obscure gastrointestinal bleeding” (OGIB) should be reserved for patients not found to have a source of bleeding even after the performance of SB evaluation [4].

The diagnostic yield of small-bowel capsule endoscopy (SBCE) in patients with suspected small-bowel bleeding (SSBB) ranges from 55% to 62% [5–7]. Compared with alternative modalities, SBCE has been consistently shown in prospective studies to be significantly superior to push-enteroscopy [8], computed tomography enterography (CTE) [9], CT angiography and standard angiography [10], and intraoperative enteroscopy [11], and to be as good as DAE [6] in evaluating and finding the lesion(s) causing the bleeding in patients with SSBB. Careful patient selection may improve the diagnostic yield of SBCE in patients with SSBB. Diagnostic yield is greatest if the interval between SBCE and the last bleeding episode is as short as possible [12] (see following statements and supporting evidence). Other characteristics associated with an increased yield include a history of an overt bleed, use of antithrombotic agents, inpatient status, male sex, older age, and liver and renal comorbidities [13, 14]. From a technical point of view, a careful and focused review, performed by adequately trained readers, using the latest available technological advances (e.g., chro-moendoscopy [15], and artificial intelligence [AI]) might contribute to further increasing the diagnostic yield of capsule endoscopy.

In patients with SSBB, SBCE showed an excellent safety profile. The rates of capsule retention range from 1.2% [5] to 2.1% [16]. Thus, routine cross-sectional imaging or the use of a patency capsule is not essential before SBCE in these patients.

It is known that cross-sectional techniques may be helpful in SSBB [4]. This updated Guideline can report only a few further studies that have been published on this subject. A meta-analysis, with 9 mainly high quality studies (396 patients), evaluated the diagnostic accuracy of CTE on SSBB detection [17]. The pooled sensitivity and specificity of CTE were 0.724 (95% CI 0.651–0.789) and 0.752 (95% CI 0.691–0.807), respectively. The area under the curve (AUC) was 0.7916 (95%CI 0.723–0.860). A small retrospective cohort study [18] showed that when CTE and SBCE were used in combination within 30 days, the sensitivity was significantly higher at 30/31 (96.8%) than that of SBCE alone at 24/31 (77.4%; P=0.0412).

Although CTE showed only moderate accuracy in the diagnosis of SSBB, it must also be remembered that SBCE can miss solitary protruding lesions in the proximal small bowel, such as small-bowel tumors (SBTs) [19]. CTE may thus be reasonably used as a complementary diagnostic method to SBCE, especially when an SBT is suspected.

DAE is both diagnostic and therapeutic but compared with SBCE, it has a lower rate of complete examination of the small bowel and is more invasive. In addition, the diagnostic yield of double-balloon enteroscopy (DBE) improves from 56% (95% CI 48.9%–62.1%) to 75% (95% CI 60.1%–90.0%) if DBE is preceded by a positive SBCE (odds ratio [OR] for positive DBE 1.79, 95% CI 1.09%–2.96%; P=0.02) [6]. Although the clinical presentation may indicate the preferential endoscopic insertion route for DAE, SBCE is also an effective tool for guiding the selection of the correct DAE approach (oral vs. anal) [20].
**Table 1** Small-bowel capsule endoscopy and device-assisted enteroscopy for diagnosis and treatment of small-bowel disorders. Summary of all ESGE Guideline 2015 and ESGE Guideline 2022 recommendations. Changes from the 2015 Guideline (new or modified recommendations) are shown in bold.

<table>
<thead>
<tr>
<th>ESGE Guideline 2015</th>
<th>ESGE Guideline 2022 (in bold if modified)</th>
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<tbody>
<tr>
<td><strong>Suspected small-bowel bleeding</strong></td>
<td><strong>1. ESGE recommends small-bowel capsule endoscopy as the first-line examination, before consideration of other endoscopic and radiological diagnostic tests for suspected small-bowel bleeding, given the excellent safety profile of capsule endoscopy, its patient tolerability, and its potential to visualize the entire small-bowel mucosa. Strong recommendation, moderate quality evidence.</strong></td>
</tr>
<tr>
<td>1. ESGE recommends small-bowel video capsule endoscopy as the first-line investigation in patients with obscure gastrointestinal bleeding (strong recommendation, moderate quality evidence).</td>
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<td>2. ESGE recommends against push-enteroscopy as the first-line investigation in patients with obscure gastrointestinal bleeding, because of its lower diagnostic yield compared with small-bowel capsule endoscopy (strong recommendation, moderate quality evidence).</td>
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<tr>
<td>3. ESGE recommends performance of small-bowel capsule endoscopy as the first-line examination, before consideration of small bowel radiographic studies or mesenteric angiography, when small-bowel evaluation is indicated for obscure gastrointestinal bleeding (strong recommendation, high quality evidence). Computed tomography enterography/enteroclysis may be a complementary examination to capsule endoscopy in selected patients (weak recommendation, low quality evidence).</td>
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<tr>
<td>4. Because of capsule endoscopy’s excellent safety profile, patient tolerability, and potential for complete enteroscopy, ESGE recommends performance of small-bowel capsule endoscopy as the first-line examination, before consideration of device-assisted enteroscopy, when small-bowel evaluation is indicated for obscure gastrointestinal bleeding (strong recommendation, moderate quality evidence).</td>
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<tr>
<td>5. In patients with overt obscure gastrointestinal bleeding ESGE recommends performing small-bowel capsule endoscopy as soon as possible after the bleeding episode, optimally within 14 days, in order to maximize the diagnostic yield (strong recommendation, moderate quality evidence).</td>
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<td>6. ESGE suggests that emergency small-bowel capsule endoscopy should be considered in patients with ongoing overt obscure gastrointestinal bleeding (weak recommendation, moderate quality evidence). In such patients, ESGE suggests that device-assisted enteroscopy should also be considered as a possible first-line test, given that it allows diagnosis and treatment in the same procedure (weak recommendation, low quality evidence).</td>
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<tr>
<td>7. Given the spectrum of findings usually identified in patients with obscure gastrointestinal bleeding, when small-bowel capsule endoscopy is unavailable or contraindicated, ESGE suggests consideration of device-assisted enteroscopy as the first diagnostic test in these patients (weak recommendation, low quality evidence). ESGE suggests that device-assisted enteroscopy performed with diagnostic intent should be done as soon as possible after the bleeding episode (weak recommendation, low quality evidence).</td>
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<tr>
<td>8. ESGE does not recommend the routine performance of second-look endoscopy prior to small-bowel capsule endoscopy; however whether to perform second-look endoscopy before capsule endoscopy in patients with obscure gastrointestinal bleeding or iron-deficiency anaemia should be decided on a case-by-case basis (strong recommendation, low quality evidence).</td>
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<tr>
<td>9. ESGE recommends conservative management in those patients with obscure gastrointestinal bleeding (OGIB) and a negative small-bowel video capsule endoscopy (VCE) who do not have ongoing bleeding shown by overt bleeding or continued need for blood transfusions, since their prognosis is excellent and the risk of re-bleeding is low (strong recommendation, moderate quality evidence).</td>
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<tr>
<td>6. ESGE does not recommend routine second-look endoscopy prior to small-bowel capsule endoscopy in patients with suspected small-bowel bleeding or iron-deficiency anaemia. Strong recommendation, low quality evidence.</td>
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<tr>
<td>7. ESGE recommends conservative management in those patients with suspected small-bowel bleeding and high quality negative small-bowel capsule endoscopy. Strong recommendation, moderate quality evidence.</td>
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<tr>
<td>10. ESGE recommends further investigation using repeat VCE, device-assisted enteroscopy, or computed tomography-enterography/enteroclysis for patients with OGIB and a negative VCE who have ongoing bleeding shown by overt bleeding or continued need for blood transfusions (strong recommendation, moderate quality evidence).</td>
<td>8. ESGE recommends further investigation using repeat small-bowel capsule endoscopy, device-assisted enteroscopy, or dedicated small-bowel cross-sectional imaging for patients with suspected small-bowel bleeding and high quality negative small-bowel capsule endoscopy who have ongoing overt bleeding or continued need for blood transfusions. Strong recommendation, moderate quality evidence.</td>
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<tr>
<td>11. In patients with positive findings at small-bowel capsule endoscopy, ESGE recommends device-assisted enteroscopy to confirm and possibly treat lesions identified by capsule endoscopy (strong recommendation, high quality evidence).</td>
<td>9. ESGE recommends device-assisted enteroscopy to confirm and possibly treat lesions identified by small-bowel capsule endoscopy. Strong recommendation, high quality evidence.</td>
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**Iron-deficiency anaemia**

| 12. In patients with iron-deficiency anaemia, ESGE recommends that prior to small-bowel capsule endoscopy, all the following are undertaken: acquisition of a complete medical history (including medication use, comorbidities, and gynaecological history in premenopausal females), oesophagogastroduodenoscopy with duodenal and gastric biopsies, and ileocolonoscopy (strong recommendation, low quality evidence). | 10. ESGE recommends that in patients with iron-deficiency anaemia, the following are undertaken prior to small bowel evaluation: acquisition of a complete medical history, esophagogastroduodenoscopy with duodenal and gastric biopsies, and ileocolonoscopy. Strong recommendation, low quality evidence. |
| 13. In patients with iron-deficiency anaemia, ESGE recommends performance of small-bowel capsule endoscopy as a first-line examination, before consideration of other diagnostic modalities, when upper and lower gastrointestinal endoscopies are inconclusive and small-bowel evaluation is indicated (strong recommendation, moderate quality evidence). | 11. ESGE recommends the performance of small-bowel capsule endoscopy as a first-line examination in patients with iron-deficiency anaemia when small bowel evaluation is indicated. Strong recommendation, high quality evidence. |

**Suspected Crohn’s disease**

<p>| 15. In patients with suspected Crohn’s disease and negative ileocolonoscopy findings, ESGE recommends small-bowel capsule endoscopy as the initial diagnostic modality for investigating the small bowel, in the absence of obstructive symptoms or known stenosis (strong recommendation, moderate quality evidence). | 13. ESGE recommends small-bowel capsule endoscopy in patients with suspected Crohn’s disease and negative ileocolonoscopy findings as the initial diagnostic modality for investigating the small bowel, in the absence of obstructive symptoms or known bowel stenosis. Strong recommendation, high quality evidence. |
| 16. ESGE does not recommend routine small-bowel imaging or the use of the PillCam patency capsule prior to capsule endoscopy in these patients (strong recommendation, low quality evidence). | 14. ESGE does not recommend routine cross-sectional small-bowel imaging or the use of a patency capsule prior to capsule endoscopy to prevent the retention of the device in patients with suspected Crohn’s disease. Strong recommendation, high quality evidence. |
| 17. In the presence of obstructive symptoms or known stenosis, ESGE recommends that dedicated small-bowel cross-sectional imaging modalities such as magnetic resonance enterography/enteroclysis or computed tomography enterography/enteroclysis should be used first (strong recommendation, low quality evidence). | 15. ESGE recommends that dedicated small-bowel cross-sectional imaging modalities be used first in patients with suspected Crohn’s disease and obstructive symptoms or known bowel stenosis. Strong recommendation, moderate quality evidence. |</p>
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<td>19. ESGE recommends discontinuation of nonsteroidal anti-inflammatory drugs (NSAIDs) for at least 1 month before capsule endoscopy since these drugs may induce small-bowel mucosal lesions indistinguishable from those caused by Crohn’s disease (strong recommendation, low quality evidence).</td>
<td>18. ESGE recommends discontinuation of both selective and non-selective nonsteroidal anti-inflammatory drugs, including short-term use, as well as of low dose and/or enteric-coated aspirin (if the patient’s condition allows), for at least 4 weeks before capsule endoscopy since these drugs may induce small-bowel mucosal lesions that are indistinguishable from those caused by Crohn’s disease. Strong recommendation, low quality evidence.</td>
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<tr>
<td>Not addressed in the 2015 Guideline</td>
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<tr>
<td>20. ESGE recommends device-assisted enteroscopy with small-bowel biopsy in patients with noncontributory ileocolonoscopy and with suspicion of Crohn’s disease on small-bowel cross-sectional imaging modalities or small-bowel capsule endoscopy. Device-assisted enteroscopy with small-bowel biopsy is more likely to provide definitive evidence of Crohn’s disease than cross-sectional imaging, although the latter offers a useful less invasive alternative that better defines transmural complication (strong recommendation, high quality evidence).</td>
<td>19. ESGE recommends device-assisted enteroscopy with small-bowel biopsies in patients with noncontributory ileocolonoscopy and suspected Crohn’s disease on small-bowel cross-sectional imaging modalities or small-bowel capsule endoscopy. Strong recommendation, high quality evidence.</td>
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<tr>
<td>Established Crohn’s disease</td>
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<tr>
<td>21. In patients with established Crohn’s disease, based on ileocolonoscopy findings, ESGE recommends dedicated cross-sectional imaging for small-bowel evaluation since this has the potential to assess extent and location of any Crohn’s disease lesions, to identify strictures, and to assess for extraluminal disease (strong recommendation, low quality evidence).</td>
<td>20. ESGE recommends, in patients with established Crohn’s disease based on ileocolonoscopy findings, dedicated cross-sectional imaging for small-bowel evaluation since this has the potential to assess the extent and location of any Crohn’s disease lesions, to identify strictures, and to assess for extraluminal disease. Strong recommendation, high quality evidence.</td>
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<tr>
<td>22. In patients with unremarkable or nondiagnostic findings from such cross-sectional imaging of the small bowel, ESGE recommends small-bowel capsule endoscopy as a subsequent investigation, if deemed to influence patient management (strong recommendation, low quality evidence).</td>
<td>21. ESGE recommends, in patients with unremarkable or nondiagnostic findings from dedicated small-bowel cross-sectional imaging, small-bowel capsule endoscopy as a subsequent investigation if deemed likely to influence patient management. Strong recommendation, low quality evidence.</td>
</tr>
<tr>
<td>Not addressed in the 2015 Guideline</td>
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<tr>
<td>23. ESGE suggests the use of activity scores (such as the Lewis score and the Capsule Endoscopy Crohn’s Disease Activity Index) to facilitate prospective small-bowel capsule endoscopy follow-up of patients for longitudinal assessment of the course of small-bowel Crohn’s disease and its response to medical therapy (using mucosal healing as an end point) (weak recommendation, low quality evidence).</td>
<td>23. ESGE recommends the use of activity scores (such as the Lewis score and the Capsule Endoscopy Crohn’s Disease Activity Index [CEDCAI]) to facilitate prospective small-bowel capsule endoscopy follow-up of patients for longitudinal assessment of small-bowel Crohn’s disease and its response to medical therapy (using mucosal healing as an endpoint). Strong recommendation, low quality evidence.</td>
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<tr>
<td>24. When capsule endoscopy is indicated, ESGE recommends use of the PillCam patency capsule to confirm functional patency of the small bowel (strong recommendation, low quality evidence).</td>
<td>24. ESGE recommends, in patients with established Crohn’s disease, the use of a patency capsule before small-bowel capsule endoscopy to decrease the capsule retention rate. Strong recommendation, moderate quality evidence.</td>
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<tr>
<td>25. ESGE recommends initial conservative treatment in the case of capsule retention. ESGE recommends device-assisted enteroscopy if medical therapy has not led to promote spontaneous passage (strong recommendation, low quality evidence).</td>
<td>25. ESGE recommends initial conservative treatment in the case of capsule retention. Strong recommendation, high quality evidence.</td>
</tr>
<tr>
<td>26. ESGE recommends device-assisted enteroscopy if small-bowel endotherapy is indicated (including dilation of Crohn’s disease small-bowel strictures, retrieval of foreign bodies, and treatment of small-bowel bleeding) (strong recommendation, low quality evidence).</td>
<td>26. ESGE recommends device-assisted enteroscopy if medical therapy has not achieved spontaneous capsule passage. Strong recommendation, high quality evidence.</td>
</tr>
<tr>
<td>27. ESGE recognises small-bowel capsule endoscopy/device-assisted enteroscopy and magnetic resonance or computed tomography enterography/enteroclysis as complementary strategies (weak recommendation, low quality evidence). Cost-effectiveness data regarding optimal investigation strategies for diagnosis of small-bowel Crohn’s disease are lacking.</td>
<td>27. ESGE recommends device-assisted enteroscopy if small-bowel endotherapy is indicated (including dilation of Crohn’s disease small-bowel strictures, retrieval of a retained capsule, and/or treatment of small-bowel bleeding). Strong recommendation, high quality evidence.</td>
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See statements 13, 15, 19, 20, 21, 27
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<td>28. ESGE recommends that surveillance of the proximal small bowel in familial adenomatous polyposis is best performed using conventional forward-viewing and side-viewing endoscopes (strong recommendation, moderate quality evidence).</td>
<td>29. ESGE does not recommend small-bowel capsule endoscopy for surveillance of the proximal small bowel in familial adenomatous polyposis. Strong recommendation, moderate quality evidence.</td>
</tr>
<tr>
<td>29. When small-bowel investigation is clinically indicated in familial adenomatous polyposis, ESGE suggests that small-bowel capsule endoscopy and/or cross-sectional imaging techniques may be considered for identifying polyps in the rest of the small bowel, but the clinical relevance of such findings remains to be demonstrated (weak recommendation, moderate quality evidence).</td>
<td>30. ESGE suggests that small-bowel capsule endoscopy and/or cross-sectional imaging techniques may be considered when investigation of the mid-distal small-bowel is clinically indicated in familial adenomatous polyposis. Weak recommendation, moderate quality evidence.</td>
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<tr>
<td><strong>Peutz–Jeghers syndrome</strong></td>
<td>31. ESGE recommends, for small bowel surveillance in patients with Peutz–Jeghers syndrome, small-bowel capsule endoscopy and/or magnetic resonance enterography and enterolysis appear adequate methods for this purpose, depending on local availability and expertise, or patient preference (strong recommendation, moderate quality evidence)</td>
</tr>
<tr>
<td>30. ESGE recommends small-bowel surveillance in patients with Peutz–Jeghers syndrome. Small-bowel capsule endoscopy and/or magnetic resonance enterography and enterolysis appear adequate methods for this purpose, depending on local availability and expertise, or patient preference (strong recommendation, moderate quality evidence)</td>
<td>32. ESGE recommends device-assisted enteroscopy with timely polypectomy when large polyps (&gt;10–15 mm) are discovered by radiological examination or small-bowel capsule endoscopy in patients with Peutz–Jeghers syndrome (strong recommendation, moderate quality evidence).</td>
</tr>
<tr>
<td>31. ESGE recommends device-assisted enteroscopy with timely polypectomy when large polyps (&gt;10–15 mm) are discovered by radiological examination or small-bowel capsule endoscopy in patients with Peutz–Jeghers syndrome (strong recommendation, moderate quality evidence).</td>
<td>33. ESGE recommends that routine evaluation of the small bowel in juvenile polyposis patients should be limited to the duodenum and based on flexible forward-viewing endoscopy. Strong recommendation, low quality evidence.</td>
</tr>
<tr>
<td><strong>Juvenile polyposis</strong></td>
<td>34. ESGE recommends the use of small-bowel capsule endoscopy in patients where there is an increased risk of a small-bowel tumor. Strong recommendation, moderate quality evidence.</td>
</tr>
<tr>
<td>Not addressed in the 2015 Guideline</td>
<td>35. ESGE does not recommend, in the setting of suspected small-bowel tumor, specific investigations before small-bowel capsule endoscopy unless patients are considered to be at risk of capsule retention. Strong recommendation, low quality evidence.</td>
</tr>
<tr>
<td><strong>Small-bowel tumors</strong></td>
<td>36. ESGE recommends consideration of device-assisted enteroscopy in preference to small-bowel capsule endoscopy if imaging tests have already shown suspicion of small-bowel tumor (strong recommendation, low quality evidence).</td>
</tr>
<tr>
<td>32. ESGE recommends early use of small-bowel video capsule endoscopy in the search for a small-bowel tumour when obscure gastrointestinal bleeding and iron-deficiency anaemia are not explained otherwise (strong recommendation, moderate quality evidence).</td>
<td>37. ESGE recommends cross-sectional imaging for staging and ascertaining operability when there is a small-bowel capsule endoscopy finding of a small-bowel tumor with high diagnostic certainty. Strong recommendation, low quality evidence.</td>
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<td>33. In the setting of suspicion of a small-bowel tumour, ESGE does not recommend specific investigations before small-bowel capsule endoscopy in patients without evidence for stenosis or previous small-bowel resection (strong recommendation, low quality evidence).</td>
<td>38. ESGE recommends, when there is an uncertain diagnosis of small-bowel tumor at capsule endoscopy, biopsy sampling and tattooing of its location by device-assisted enteroscopy. Strong recommendation, low quality evidence.</td>
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<td>34. ESGE recommends consideration of device-assisted enteroscopy in preference to small-bowel capsule endoscopy if imaging tests have already shown suspicion of small-bowel tumour (strong recommendation, low quality evidence).</td>
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36. When a submucosal mass is detected by small-bowel capsule endoscopy, ESGE recommends confirmation of the diagnosis by device-assisted enteroscopy (strong recommendation, low quality evidence).

37. When capsule endoscopy shows high suspicion of submucosal mass and there is a negative but incomplete device-assisted enteroscopy, ESGE suggests cross-sectional imaging tests to confirm the diagnosis (weak recommendation, low quality evidence).

38. ESGE recommends against small-bowel capsule endoscopy in the follow-up of treated small-bowel tumours because of lack of data (strong recommendation, low quality evidence).

39. ESGE recommends, when a subepithelial mass is detected by small-bowel capsule endoscopy, confirmation of the diagnosis by device-assisted enteroscopy and/or cross-sectional imaging, depending on local availability and expertise. Strong recommendation, low quality evidence.

40. ESGE does not recommend small-bowel capsule endoscopy in the follow-up of treated small-bowel tumors because of lack of data. Strong recommendation, low quality evidence.

41. ESGE suggests considering enteroscopic placement of self-expanding metal stents in the palliation of malignant small-bowel strictures as an alternative option to surgery. Weak recommendation, low quality evidence.

42. ESGE does not recommend small-bowel capsule endoscopy to diagnose celiac disease. Strong recommendation, low quality evidence.

43. ESGE recommends using small-bowel capsule endoscopy in cases of equivocal diagnosis of celiac disease since it is essential for final diagnosis and therapy. Strong recommendation, low quality evidence.

44. ESGE recommends in nonresponsive or refractory celiac disease, small-bowel capsule endoscopy followed by device-assisted enteroscopy for diagnosis and disease monitoring. Strong recommendation, high quality evidence.

45. ESGE does not recommend small-bowel capsule endoscopy as the first-line investigation for patients with isolated chronic abdominal pain. Strong recommendation, low quality evidence.

46. ESGE recommends device-assisted enteroscopy as an alternative to surgery for foreign bodies retained in the small bowel requiring retrieval in patients without acute intestinal obstruction. Strong recommendation, moderate quality evidence.

47. ESGE suggests that in patients requiring jejunostomy for enteral feeding, DAE-assisted percutaneous endoscopic jejunostomy (PEJ) is a possible alternative to surgical jejunostomy. Weak recommendation, moderate quality evidence.


Table 1 (Continuation)

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Celiac disease

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<td>39. ESGE strongly recommends against the use of small-bowel capsule endoscopy for suspected coeliac disease but suggests that capsule endoscopy could be used in patients unwilling or unable to undergo conventional endoscopy (strong recommendation, low quality evidence).</td>
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Chronic abdominal pain

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Foreign-body retrieval

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DAE-assisted percutaneous endoscopic jejunostomy (PEJ) for enteral feeding

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DAE-ERCP in patients with altered anatomy

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DAE, device-assisted enteroscopy; ERCP, endoscopic retrograde cholangiopancreatography; ESGE, European Society of Gastrointestinal Endoscopy; PEJ, percutaneous endoscopic jejunostomy
As already stated in previous guidelines [1] and on the basis of all the above scientific evidence, SBCE can be recommended as the first-line investigation in patients with SSBB. This agrees with the recommendations of other scientific societies [4, 21, 22].

Fig. 1 presents recommended approaches for diagnosis and treatment of SSBB.

RECOMMENDATION
ESGE recommends small-bowel capsule endoscopy in patients with overt suspected small-bowel bleeding as soon as possible after the bleeding episode, ideally within 48 hours, to maximize the diagnostic and subsequent therapeutic yield. Strong recommendation, high quality evidence.

Despite the unquestionable role of early SB evaluation in patients with SSBB, especially in cases of overt bleeding, the optimal timing is still debated. The 14-day timeframe, suggested in the previous ESGE guideline [1], is somewhat arbitrary and quite broad.

Since the publication of the initial guideline [1], six retrospective studies and two meta-analyses have been published to compare the diagnostic and therapeutic yield of SB endoscopic procedures in the setting of overt SB bleeding according to the timing of SB evaluation (performed with either SBCE or DAE).

Zhao et al. [23] carried out a propensity score-matching study on 997 patients, that supported previous ESGE statements; they found that early SBCE (within 14 days from last bleeding event) was associated with a significantly higher rate of diagnosis (56.4% vs. 45.5%, \( P=0.001 \)), with ORs of 0.648 (95% CI 0.496–0.847, \( P=0.001 \)) and 0.666 (95% CI 0.496–0.894, \( P=0.007 \)) at univariate and multivariate analysis, respectively. In this study, the incidence of rebleeding within 1 year following treatment was significantly lower (24.7% vs. 36.7%, \( P=0.041 \)) for patients who underwent early SBCE. Chao et al. [24] reported a detection rate for the source of bleeding ranging from 70% to 77.6% if SBCE was performed in the first 3 days from the first bleeding episode in patients (\( n=60 \)) with overt bleeding. In contrast, the detection rate decreased to 36.4% if SBCE was performed after the 4th day. Using a 48-hour cut-off, Kim et al. [25] found that among 94 patients, the 30 who underwent SBCE within 2 days from the last bleeding had a greater diagnostic yield (66.7% vs. 40.6%, \( P=0.019 \)), a greater subsequent therapeutic yield (24.7% vs. 9.4%, \( P=0.028 \)) and a shorter hospital stay (5 days, 95% CI 4.8–7.7 vs. 7 days, 95% CI 6.9–10.1, \( P=0.039 \)). A shorter hospital stay, as well as a decrease in resource utilization in the index hospitalization, was also demonstrated by Wood et al. [26] in inpatients receiving an early SBCE. Iio et al. [27] found a lesion detection rate of 80% (12/15) in patients with ongoing overt bleeding who underwent early SBCE (15/127) compared to 47% (53/112) in the “late” group (\( P=0.0174 \)). These data were consistent with the results of Song et al. [28], who showed that early deployment of SBCE results in a significantly higher diagnostic yield (OR for relevant lesion detection was 4.99 for \( <24h \) group vs. 8-day group). On the other hand, in the study of Gomes et al. [29] (\( n=115 \)), where the timing of SBCE was further divided (≤48h, 48h–14d, ≥14d), the overall diagnostic yield was high (about 80%) and similar among the three groups irrespective of SBCE timing (\( P=0.39 \)). However, the three timing-based subgroups were small (about 30 patients in each) and when SBCE was performed within 48 hours, a trend toward an increased diagnostic yield was observed (\( P=0.06 \)). In addition, the early group showed the highest therapeutic yield (66.7% vs. 40% vs. 31.7%, \( P=0.005 \)) and the lowest rebleeding rate (15.4% vs.
34.3 % vs. 46.3 %, P=0.007), with a longer time to rebleed when compared with the > 48-h groups (P=0.03).

Recently, a meta-analysis from Uchida et al. [30], by pooling 19 previous studies (9 prospective, 9 retrospectives, 1 unspecified), confirmed that performing SBCE within 2 days leads to high diagnostic and therapeutic yields (55.9 % and 65.2 %, respectively). However, the metaregression was based on subgroups with small sample size and heterogeneous data [30]. The largest meta-analysis available so far, involving 39 studies, confirmed higher pooled diagnostic yields for SBCE performed in the first 24, 48, and 72 hours, being 83.4 % (95%CI 76.30 %– 90.46 %), 81.3 % (95%CI 75.20 %– 87.43 %) and 63.6 % (95%CI 45.59 %– 81.51 %), respectively. The pooled therapeutic yields for the same timings were 57.56 % (95%CI 36.95 %– 78.16 %), 59.09 % (95%CI 43.66 %– 74.52 %) and 18.90 % (95%CI 11.26 %– 26.54 %), respectively [31].

**RECOMMENDATION**

ESGE suggests that device-assisted enteroscopy be considered as an alternative first-line test in selected cases, given that it allows diagnosis and treatment in the same procedure, depending on the clinical scenario and local availability.

Weak recommendation, low quality evidence.

Two previously mentioned studies [30, 31] not only evaluated the diagnostic yield of SBCE but also dealt with the performance of DAE in the same setting. According to Estevinio et al. [31], the pooled diagnostic and therapeutic yields of early DAE were superior to those of SBCE by 7.97 and 20.89 percentage points, respectively (P<0.05). However, it is not possible to exclude that the DAE results may be influenced both by a selection bias, related to patient features (e.g., patients undergoing direct DAE are likely to have more severe bleeding), and by a detection bias, since several patients may have received another diagnostic test, with a positive result, before DAE. In addition, urgent DAE may raise significant organizational issues; it is not readily available in most centers and requires trained personnel.

Therefore, even in overt SSBB, a sequential approach with a diagnostic examination (e.g., SBCE, CT angiography etc.) followed by a potentially therapeutic one (e.g., DAE) should be preferred. Performance of DAE in the first 72 hours is most often dependent on performance of SBCE in the first 48 hours [31]. A recent retrospective study with a large sample size of patients undergoing both SBCE and DBE [32] also confirmed that a short interval between the two procedures maximizes the effectiveness of the diagnostic/therapeutic process. Although the agreement between SBCE and DBE was generally rated as suboptimal (k=0.059), it markedly improved (k=0.323) when the procedures were performed within 1–5 days of each other. As demonstrated for SBCE, in the overt SB bleeding setting, recent data confirm the importance of keeping the interval between DAE and the bleeding episode as short as possible. In fact, in the pooled analysis of double-arm studies [31], the odds for a positive diagnosis (OR 3.99; P<0.01; I²=45 %) and subsequent therapeutic intervention (OR 3.86; P<0.01; I²=67 %) were significantly superior in the early group, for either DAE or SBCE.

**RECOMMENDATION**

ESGE recommends, in patients with overt suspected small-bowel bleeding, device-assisted enteroscopy to be performed optimally within 48–72 hours after the bleeding episode.

Strong recommendation, high quality evidence.

SBCE has a very limited number of absolute contraindications [33], such as GI obstruction. However, SBCE may also be unavailable, especially in emergency settings, although lately, there is a trend of increasing use outside the endoscopy suite [34]. Overall, there is not enough evidence-based data to recommend a single specific examination as first-line when SBCE is unavailable. A meta-analysis [9] of a total of 18 studies (n = 660 patients) reported the pooled diagnostic yield of CTE in evaluating SSBB as 40 % (95%CI 33 %–49 %). Seven studies (n = 279) compared the yield of CTE with SBCE. The yields for CTE and SBCE for all findings were 34 % and 53 %, respectively (incremental yield –19 %, 95%CI –34 % to –4 %). Therefore, CTE has been described as an effective modality to show the precise location of bleeding and guide subsequent enteroscopy management, especially in patients with bleeding from tumors and overt bleeding [9]. In an emergency setting, DAE has been described as effective as suggested by a recent systematic review and meta-analysis [31], including retrospective studies in which this procedure was performed as first-line for selected patients.

**RECOMMENDATION**

ESGE does not recommend routine second-look endoscopy prior to small-bowel capsule endoscopy in patients with suspected small-bowel bleeding or iron-deficiency anemia.

Strong recommendation, low quality evidence.
Good quality upper and lower GI endoscopy is crucial in the investigation of SSBB. Evidence and recent guidelines propose an acceptable minimal examination time to ensure good quality examination and meeting minimum standards [35, 36]. In patients where bidirectional endoscopy has been negative, with the persistence of symptoms or suspicion of SB bleeding, SBCE is the preferred next diagnostic test. Several studies had investigated routine second-look endoscopy before capsule endoscopy and highlighted this as not being cost-effective, as stated in the 2015 Guideline [1]. Since the publication of the latter, eight further studies have been published on this subject. A study by Innocenti et al. [37] showed non-SB lesions detected in 30% of cases, of which 43% were bleeding. The study was retrospective and without randomization. Similarly, another retrospective study by Cleré-Jehl et al. [38] studied 69 endoscopy-negative patients >65 years, with persistent IDA. Further investigations were performed in 45 patients; 64% of the second-look GI endoscopies led to significant changes in treatment compared with 25% for the capsule endoscopies. Conventional diagnoses of IDA were ultimately established for 19 (27%) patients and included 3 cancer patients suggesting second-look endoscopy is favored for persistent IDA. On the other hand, a prospective study by Riccioni et al. [39] showed that at SBCE, findings in the upper GI tract were found in 21% and the colon in 6.4%. Subsequent studies by Akin et al. [40], Hoedemaker et al. [41], and Juanmartiñena Fernández et al. [42–44] (this last group published three separate studies about esophageal, gastroduodenal, and colonic findings on SBCE), all retrospective in nature, conclude that clinicians should carefully review not just SB images but also those of the esophagus, stomach, and colon.

There have been no further cost-effectiveness studies. Overall, the current literature is inadequate to support routine repetition of standard endoscopy, and this should be reserved on a case-by-case basis. However it highlights the importance of a good standard of baseline endoscopy performance.

**RECOMMENDATION**

ESGE recommends conservative management in those patients with suspected small-bowel bleeding and high quality negative small-bowel capsule endoscopy. Strong recommendation, moderate quality evidence.

Analogously to upper and lower GI endoscopy, for SBCE to be considered a reliable diagnostic tool on which subsequent follow-up is based, it must be rated a high quality examination, according to ESGE quality standards [45], and evaluated by a dedicated and properly trained reader, according to ESGE curriculum criteria [46]. Even more than in upper and lower endoscopy, given the passive nature of capsule endoscopy (e.g., lavage and aspiration cannot be done), the characteristics of the luminal contents (e.g., presence of bubbles, fecal material, or turbid fluid) strongly impact the quality of the examination. Therefore, adequate SB visualization is a crucial element in ensuring a reliable assessment of the small intestine. Although the current ESGE technical guidelines specifically address this issue [47], the evidence is rapidly evolving [48] and remains somewhat controversial [49].

A systematic review and meta-analysis [50], including 26 mostly high quality studies with 3657 individuals, showed that a negative SBCE implies adequate assurance of a subsequently low risk of rebleeding. The pooled rate of rebleeding after negative SBCE was 0.19 (95% CI 0.14–0.25; P < 0.0001). The pooled OR of rebleeding was 0.59 (95% CI 0.37–0.95; P < 0.001), and moreover, the effect was more pronounced in studies with a short follow-up (OR 0.47, 95% CI 0.24–0.94; P < 0.001). On top of that, prospective studies showed a lower OR of rebleeding at 0.24 (95% CI 0.08–0.73; P = 0.01). Lastly, there was no statistically significant difference in rebleeding after SBCE for occult and overt OGIB. Therefore, patients with negative SBCE after an episode of SSBB can be safely managed with watchful waiting, at least in the short term [51, 52].

However, in the long-term, recurrence of bleeding is not uncommon [53–55], and further investigations could be required. In these cases, repeating the diagnostic workup by SBCE appears to have more diagnostic value than DAE; a small study from Japan showed that the rate of positive findings in the repeat SBCE group was significantly higher than in the DBE group [56]. A closer follow-up has been proposed in patients with a higher red blood cell transfusion requirement previous to an SBCE and overt bleeding [55, 57, 58] or severe anemia [59], as they are associated with higher rebleeding rates. Recently, de Sousa Magalhães et al. developed and validated a score (RHE-MITT) that accurately predicts the individual risk of SB rebleeding after initial SBCE [60, 61].

**RECOMMENDATION**

ESGE recommends device-assisted enteroscopy to confirm and possibly treat lesions identified by small-bowel capsule endoscopy. Strong recommendation, high quality evidence.

It is known that the diagnostic yield of DBE significantly improves if DBE is preceded by a positive SBCE [6] and a recent meta-analysis reported that this sequential approach increased the diagnostic yield for vascular lesions by 7% [62]. Moreover, in patients with negative SBCE, a subsequent DBE can identify the source of the bleeding in about one third [6, 56]. In addition
to its therapeutic possibilities, DBE has been reported to help clarify the origin of bleeding when SBCE shows only blood in the lumen or doubtful findings [63]. The correct management of patients with SSBB involves using both techniques.

Although several studies have assessed the diagnostic and therapeutic yield of SBCE and DAE in SB bleeding, the emphasis should be on meaningful results when we consider outcomes in clinical practice. In this clinical setting, a positive patient outcome should be either bleeding cessation or anemia resolution. In addition, other important clinical outcomes for evaluation may include mortality and hemoglobin levels or the reduction in the numbers of endoscopic procedures, hospitalizations, and blood transfusions.

In this regard, both the older literature [1] and the more recent studies evaluating the impact of SB endoscopy on the clinical outcomes of patients with SB bleeding have produced conflicting results [32,64–68]. This is probably because considerable heterogeneity exists across studies in the definition, relevance, and clinical management of vascular lesions and follow-up periods. Furthermore, the studies differ in the severity of the bleeding of the enrolled patients, and, above all, a standardized intervention protocol for the identified bleeding lesions had not always been established a priori. Though a recent meta-analysis [31] assessing the impact of early SB endoscopy in patients with overt SSBB showed a lower recurrent bleeding rate (OR 0.40; \( P < 0.01; I^2 = 0 \% \)) when SBCE/DAE was performed very close to the bleeding episode, further high quality research, including randomized trials, is needed to clarify the open questions and clinical management regarding SB bleeding.

Iron-deficiency anemia

**RECOMMENDATION**

ESGE recommends that in patients with iron-deficiency anemia, the following are undertaken prior to small-bowel evaluation: acquisition of a complete medical history, esophagogastroduodenoscopy with duodenal and gastric biopsies, and ileocolonoscopy.

Strong recommendation, low quality evidence.

**RECOMMENDATION**

ESGE recommends the performance of small-bowel capsule endoscopy as a first-line examination in patients with iron-deficiency anemia when small bowel evaluation is indicated.

Strong recommendation, high quality evidence.

The evidence published since the previous ESGE guideline [1] and the most recent practice guideline on IDA [69] confirm that, before evaluation of the small-bowel, patients with IDA should undergo a thorough anamnestic evaluation and a multi-step diagnostic-therapeutic workup that includes endoscopic evaluation of the upper and lower digestive tract.

Furthermore, the British Society of Gastroenterology (BGS) guideline for the management of IDA in adults [69] recommends that, before the SB evaluation is planned, an empirical iron replacement trial (IRT), should be performed with appropriate dosage and duration. According to the BSG guideline, endoscopic SB examination should be performed only if the target values are not reached in the initial IRT or if anemia recurs at the end of treatment. However, no clinical trials have compared the clinically relevant outcomes (e.g., diagnostic yield and possible diagnostic delay) in patients referred for SB study according to the IRT outcome. This policy may lead to different results in different subgroups of patients. Therefore, the available evidence appears insufficient to recommend using the IRT as a decision-making tool in deciding to perform an SB study.

Considering multiple clinical issues, a comprehensive overall assessment should always be performed when planning SBCE. Several studies pursued the aim of identifying such predictive factors for SB pathology. Male sex, older age, low mean corpuscular volume (MCV), low hemoglobin values, high transfusion requirement, use of nonsteroidal anti-inflammatory drugs (NSAIDs) in the last 2 weeks before SBCE, and antithrombotic therapy have been demonstrated to correlate with diagnostic yield in IDA patients [70–75]. Hypoalbuminemia has also been shown to increase the proportion of positive findings at SBCE in a subgroup of celiac disease patients presenting with persistent IDA despite a gluten-free diet (GFD) [76].

In recent years, new evidence has also emerged concerning the possible role of fecal occult blood testing (FOBT), either guaiac or immunochemical, as a filter test to select IDA patients for SBCE [77–79]. The meta-analysis by Yung et al. [80] found, for all positive FOBT, sensitivity 0.60 (95%CI 0.50–0.69), specificity 0.72 (95%CI 0.52–0.86), and diagnostic OR 3.96 (95%CI 1.50–10.4) for SB findings. Corresponding values for fecal immunochemical testing alone were sensitivity 0.48 (95%CI 0.36–0.61), specificity 0.60 (95%CI 0.42–0.76), and diagnostic OR 1.41 (95%CI 0.72–2.75). Nevertheless, there is still insufficient evidence to recommend FOBT in routine practice as a screening tool for deciding whether to perform SBCE in IDA patients. Larger studies may better clarify its usefulness and lead to future guidance changes.

In recent years, it has also been shown that, although there are some differences in terms of both diagnostic yield and the spectrum of findings between young and elderly patients, age is not a discriminating factor when SB studies are performed in patients with IDA and negative bidirectional endoscopy [74]. Interestingly, two studies [81,82] focused on the subgroup of female IDA patients and showed a lower diagnostic yield in premenopausal women compared to post-menopausal women. Moreover, Silva et al. [82] found that in premenopausal women, only 1.8% required therapeutic endoscopy, whereas in 17.3% of post-menopausal women, SBCE findings led to additional endoscopic treatment. Furthermore, the rebleeding rate at 1, 3 and 5 years was 3.6%, 10.2%, and 10.2% in premenopausal women and 22.0%, 32.3%, and 34.2% in post-menopausal women. These figures might suggest a higher threshold for SBCE in pre-menopausal women. 

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Pennazio Marco et al. Small-bowel capsule endoscopy... Endoscopy | © 2022. European Society of Gastrointestinal Endoscopy. All rights reserved.
menopausal women. However, this evidence is insufficient to make any firm recommendation.

According to previous ESGE guidelines [1], large studies have confirmed that SBCE is the test of choice for evaluating the small intestine in patients with IDA, both because of its high diagnostic yield and favorable safety profile [70, 71, 77, 83, 84]. In contrast, there is conflicting and inconclusive evidence about the role of second-look endoscopy before SBCE in IDA patients [37, 38, 73]. Therefore, repetition of upper and lower endoscopies should be decided on a case-by-case basis, considering the timing and quality of upper and lower endoscopy performed before SBCE.

Furthermore, recent data confirm that negative SBCE provides adequate evidence of a low risk of rebleeding. Such patients can therefore be safely managed with watchful waiting [50, 53, 85, 86]. Nevertheless, SB neoplasia and diverticula are mural-based lesions that can cause IDA but can be missed at SBCE, and for which CTE has been shown to have higher sensitivity [9, 17, 87]. Since the 2015 ESGE clinical guideline [1] there have been no recent large studies that have investigated the diagnostic yield of DAE exclusively in IDA patients. However, performance can be similar to that reported for patients in the SSBB setting.

**Crohn’s disease**

**Suspected Crohn’s disease**

**RECOMMENDATION**

ESGE recommends ileocolonoscopy as the first endoscopic examination for investigating patients with suspected Crohn’s disease.

Strong recommendation, high quality evidence.

**RECOMMENDATION**

ESGE recommends small-bowel capsule endoscopy in patients with suspected Crohn’s disease and negative ileocolonoscopy findings as the initial diagnostic modality for investigating the small bowel, in the absence of obstructive symptoms or known bowel stenosis.

Strong recommendation, high quality evidence.

**RECOMMENDATION**

ESGE does not recommend routine cross-sectional small-bowel imaging or the use of a patency capsule prior to capsule endoscopy to prevent the retention of the device in patients with suspected Crohn’s disease.

Strong recommendation, high quality evidence.

Up to 83% of patients with CD have SB involvement at diagnosis [88], and in approximately 90% of patients with SB CD, the disease involves the terminal ileum [89]. Thus, ileocolonoscopy is considered to be the first-line investigation for CD and is sufficient to establish the diagnosis in most patients [90]. While the addition of capsule assessment may improve specificity, the discriminatory ability of SBCE was shown in a recent study not to be superior to ileocolonoscopy alone as an initial investigation for CD [91].

Skip lesions may result in a false-negative ileocolonoscopy [92], and SBCE should be considered when ileoscopy is not achieved or when proximal SB disease must be excluded.

For patients with suspected CD, two recent meta-analyses have confirmed SBCE has a diagnostic yield for SB disease similar to that of magnetic resonance enterography (MRE), CTE, and abdominal ultrasound, while confirming its superiority to both small-bowel follow-through and enteroclysis [93, 94]. Subgroup analysis of the 2017 meta-analysis of Koplov et al. [93] suggests that for patients with established disease, SBCE is more sensitive for proximal (jejunal) disease compared with MRE (OR 2.79, 95%CI 1.2–6.48; P=0.02). Similarly, Choi et al.’s meta-analysis [94] found that SBCE detected more ileal disease in patients with established CD than ileocolonoscopy (SBCE 60% vs. ileocolonoscopy 48%; weighted incremental yield [lyw] 0.11, 95%CI 0.00–0.22; P=0.004). Two recent studies have confirmed a diagnostic advantage for SBCE in assessing SB disease in established CD, for the entire small bowel versus MRE [95], and for the proximal and mid-small bowel versus MRE and CTE [96]. These studies support SBCE as the appropriate next investigation in patients with suspected CD after failed ileocolonoscopy and as the most sensitive means of mapping SB disease in patients with established CD [95, 96].

SBCE should be seen as complementary to ileocolonoscopy in doubtful cases, to confirm the diagnosis and simultaneously determine disease location, extent, and activity. Even after positive ileocolonoscopy findings, SBCE can add important diagnostic information and support a CD diagnosis.

A retrospective observational study by Freitas et al. [97] investigated 102 patients found to have “isolated terminal ileitis” at ileocolonoscopy, endoscopic abnormalities proximal to the terminal ileum were found in 36.3% of patients; one third (35/102) were finally diagnosed with CD. Similarly, isolated ileitis on SBCE can frequently herald an ultimate diagnosis of CD, even in patients with an initial negative ileocolonoscopy [98, 99].

The risk of capsule retention in patients with suspected CD, without obstructive symptoms or known stenosis, and no history of SB resection is low and similar to that of patients who are being investigated for SB bleeding [100]. A careful clinical history may be the most helpful way to determine the risk of capsule retention in this setting.

In 2017, Rezapour et al. [16] published a meta-analysis showing a slightly higher SBCE retention rate even in suspected CD than previously reported. Retention rates were 8.2% (95%CI 6.0%–11.0%) for established CD and 3.6% (95%CI 1.7%–6.6%) for suspected CD (studies of patients with strictures on CTE/MRE or patency capsule retention were excluded). However, there was significant heterogeneity among the studies ($I^2 = 69$).
A more recent meta-analysis by Pasha et al. [100] evaluated SBCE retention in patients with suspected and established CD. The retention rate in patients with established CD was 4.63% (95% CI 3.42%–6.25%; 32 studies) and in patients with suspected CD it was 2.35% (95% CI 1.31%–4.19%; 16 studies). Patients with established CD were 3.5 times more likely to experience retention than those with suspected CD (95% CI 2.12–5.78; 16 studies).

Several additional observational studies have also reported a low risk of capsule retention in patients with suspected CD [91, 101–103]. These studies have also shown that the use of either cross-sectional imaging [101, 102] or patency capsule tests [102] in high risk patients with suspected CD (suspected stricture) can avoid capsule retention.

**RECOMMENDATION**

ESGE recommends that dedicated small-bowel cross-sectional imaging modalities be used first in patients with suspected Crohn’s disease and obstructive symptoms or known bowel stenosis. 

Strong recommendation, moderate quality evidence.

If patients with suspected CD present with obstructive symptoms or known stenosis, dedicated SB cross-sectional imaging in the form of CTE or MRE (which may also provide an additional evaluation of mural and extramural disease) should be the investigation of choice.

Recent studies have shown a high incidence of SB strictures in patients with newly diagnosed CD, particularly in those with isolated SB rather than ileocolonic disease (OR 3.04, P=0.02 [104]; and 20.5% vs. 9.4%, P=0.002 [105]). The efficacy of MRE to detect SB stenosis has been confirmed in a meta-analysis [106] and a comparative observational study with enteroscopy [107], reporting sensitivities of 65% and 61% and specificities of 93% and 93%, respectively. Moreover, magnetic resonance imaging (MRI) combined with clinical assessment can accurately predict complications (fistulas in 98% and intra-abdominal abscesses in 99%) [108].

The retrospective study by Al-Bawardy et al. [109] revealed that patients with SBCE retention were more likely to have, as identified on pre-SBCE CTE, strictures (63% vs. 23%), partial SB obstruction (63% vs. 38%), or SB anastomosis (88% vs. 23%), as compared with patients who had passed the capsule. SBCE may still be applied in this setting if the use of a patency capsule confirms the functional patency of the small bowel. Dedicated SB cross-sectional imaging can overestimate or have low specificity and low positive predictive value (PPV) for the presence of stenosis [110,111]. Therefore, use of a patency capsule is recommended even in cases of negative findings from cross-sectional modalities in those with suspected CD and obstructive symptoms. A study in 2016 by Rondonotti et al. [110] supports this assertion, with capsule retention occurring in their at-risk cohort with negative CTE findings prior to SBCE. Rozendorn et al. [111] evaluated the ability of MRE to predict retention; because of the low specificity (59%) and low PPV (40%) of MRE for prediction of retention, the authors also recommended patency capsule use prior to SBCE in at-risk patients, regardless of MRE findings.

The corollary is also true; in 2008, Herrerías et al. [112] evaluated 106 patients with stenosis seen on small-bowel follow-through or CT, who were subsequently also given a patency capsule. The patency capsule confirmed functional patency in 59 patients (56%). These patients later underwent SBCE safely, with no cases of capsule retention. González-Suárez et al. reported similar overestimation of stenosis for MRE [95].

It is also important to note that a few case series have reported patency capsule retention in patients with suspected CD [113,114]. In all patients with findings of wall thickening or stenosis, CT was performed before patency capsule use. Patency capsule retention may cause transient obstructive symptoms, which usually resolve spontaneously, albeit resultant SB perforation has been reported [114,115].

**SBCE is indicated for investigating patients with suspected CD, nondiagnostic terminal ileitis, or inflammatory bowel disease, type unclassified (IBD-U) [116]. Symptoms alone are a poor predictor of CD. The International Conference on Capsule Endoscopy (ICCE) [117] recommended a broader definition of suspected CD that includes inflammatory markers, abnormal imaging, and/or extraintestinal manifestations [118,119]. It has also been demonstrated that ICCE criteria can be used as an effective selection tool for SBCE since patients with fewer than two ICCE criteria are not only unlikely to have inflammatory changes in the small bowel but also to be diagnosed with CD in the follow-up [118].**

Recent meta-analyses have consistently demonstrated that fecal calprotectin has significant diagnostic accuracy for detecting SB CD [120–122]. The likelihood of a positive diagnosis is very low in patients with suspected CD with calprotectin <50μg/g. A cutoff of 100μg/g has demonstrated high sensitivity and specificity and appears to be the optimal cutoff value to be used as a screening tool for SB CD [118,121]. Moreover, in a prospective validation study, a combined diagnostic strategy...
based on clinical presentation with Red Flags index score ≥ 8 and/or fecal calprotectin > 250 ng/g showed average values (ranges) of sensitivity 100% (29%–100%), specificity 72% (55%–85%), PPV 21% (5%–51%), and NPV 100% (88%–100%) for the diagnosis of CD [123]. Evidence also shows that a combination of biomarkers can further enhance patient selection.

A diagnostic workflow is proposed for investigation of patients with suspected CD and nondiagnostic ileocolonoscopy (▶Fig. 2).

**RECOMMENDATION**
ESGE recommends discontinuation of both selective and nonselective nonsteroidal anti-inflammatory drugs, including short-term use, as well as of low dose and/or enteric-coated aspirin (if the patient’s condition allows), for at least 4 weeks before capsule endoscopy since these drugs may induce small-bowel mucosal lesions that are indistinguishable from those caused by Crohn’s disease. Strong recommendation, low quality evidence.

NSAIDs, including enteric-coated or low-dose aspirin, are a common cause of SB erosions and ulcerations because of direct toxicity and systemic effects on prostaglandin metabolism. Cyclo-oxygenase 2 (COX 2)-selective agents have also been shown to cause comparable SB damage; therefore, the current ESGE recommendations apply to both selective and nonselective NSAIDs. Severe enteropathy, such as circumferential ulcers with stricturing (diaphragmatic disease), has been described in approximately 2% of patients on long-term NSAID use [124]. Short-term use results in SB injury in most patients, manifesting as multiple petechiae or red spots, erythematous patches, loss of villi, erosions, and ulcers [125]. After only 2 weeks of treatment, up to 71% of patients have some evidence of drug-induced SB lesions [124, 126, 127], and the reported prevalence in long-term low dose aspirin users is 88.5%–100% [128]. Characteristic features of NSAID-induced injury include: (i) multiple superficial lesions; (ii) similar distribution in the jejunum and ileum; (iii) lesions < 1 cm; (iv) uncommon ileocecal valve involvement [129].

The use of proton pump inhibitors (PPIs), histamine H2-receptor antagonists, or enteric-coated aspirin formulations is associated with a higher risk for NSAID-induced enteropathy [130, 131]. Indeed, a prospective SBCE study found that PPI use (OR 2.04, 95%CI 1.05–3.97) and use of enteric-coated aspirin (OR 4.05, 95%CI 1.49–11.0) were the two most important risk factors for the presence of mucosal breaks [132]. Chronic acid suppression could lead to SB bacterial overgrowth, namely of enterobacteria which contribute to the development of NSAID-induced enteropathy, while enteric-coated aspirin formulations dissolve in the small bowel rather than the stomach or duodenum, resulting in localized direct toxicity.

No data are available regarding the interval required for spontaneous healing of NSAID/low dose aspirin and/or enteric-coated aspirin-induced SB mucosal lesions. However, in the setting of suspected CD, the current recommendation to suspend NSAIDs for 4 weeks before SBCE to allow for complete mucosal healing remains generally recommended if the patient’s clinical condition allows. If discontinuation is clinically contraindicated, interpretation of SBCE findings should consider that any lesion identified may have been caused by the ongoing use of these medications.

**RECOMMENDATION**
ESGE recommends device-assisted enteroscopy with small-bowel biopsies in patients with noncontributory ileocolonoscopy and suspected Crohn’s disease on small-bowel cross-sectional imaging modalities or small-bowel capsule endoscopy. Strong recommendation, high quality evidence.
As stated in the previous guideline [1], despite all the recent advances in endoscopic and dedicated SB cross-sectional imaging, CD may still pose a diagnostic challenge, mainly if it is confined to the small bowel [90, 133]. Furthermore, it may be challenging to differentiate inflammatory SB lesions with other etiologies, such as infection (e.g., mycobacterial disease), drugs (e.g., NSAIDs and olmesartan), and malignancy (e.g., lymphoma), from similar lesions caused by CD. In such circumstances, direct endoscopic evaluation and biopsy of lesions at DAE is helpful in ruling out other causes and/or providing corroborative evidence of a diagnosis of SB CD [1,47]. Since 2015 [1], there has been further support for the usefulness of DAE in this context [134,135]. A retrospective series by Tun et al. (n = 100) [134], evaluated the role of DBE in the setting of suspected CD, where a definitive diagnosis through other modalities remained elusive. In this cohort, histopathology of biopsies taken at DBE was helpful to support a diagnosis of CD in 23%. In another similar retrospective series by Holleraan et al., which included 13 adult patients, single-balloon enteroscopy (SBE) contributed to the diagnosis of CD in 39% [135].

Established Crohn’s disease

**RECOMMENDATION**
ESGE recommends, in patients with established Crohn’s disease based on ileocolonoscopy findings, dedicated cross-sectional imaging for small-bowel evaluation since this has the potential to assess the extent and location of any Crohn’s disease lesions, to identify strictures, and to assess for extraluminal disease.

Strong recommendation, high quality evidence.

**RECOMMENDATION**
ESGE recommends, in patients with unremarkable or nondiagnostic findings from dedicated small-bowel cross-sectional imaging, small-bowel capsule endoscopy as a subsequent investigation if deemed likely to influence patient management.

Strong recommendation, low quality evidence.

**RECOMMENDATION**
ESGE suggests that small-bowel capsule endoscopy may be useful for assessment of Crohn’s disease extent and for monitoring and guiding the “treat-to-target” strategy.

Weak recommendation, low quality evidence.

The present ESGE guideline confirms that, in the setting of established CD, when SB evaluation is indicated, SB cross-sectional imaging with CTE or MRE generally takes precedence over SBCE since these modalities can assess the transmural and extraluminal nature of the disease and its anatomical distribution [1,136]. However, as discussed previously, there is growing evidence from published meta-analyses and observational studies to show that SBCE is more sensitive than cross-sectional imaging for mucosal disease throughout the small bowel in patients with established as well as suspected CD [93–96]. SBCE has been shown to be a complementary test, increasing the identification of more diffuse SB disease even in patients with a positive ileocolonoscopy.

Recent studies have evaluated the potential benefit of a panenteric capsule endoscopy for further evaluation of patients with CD. A study by Bruining et al. [137] compared panenteric capsule endoscopy with MRE and ileocolonoscopy. The overall sensitivities for active enteric inflammation (panenteric capsule endoscopy vs. MRE and/or ileocolonoscopy) were 94% vs. 100% (P=0.125) and the specificities were 74% vs. 22%, respectively (P=0.001). The sensitivity of panenteric capsule endoscopy was superior to that of MRE within the proximal small bowel (97% vs. 71%, P=0.021), and similar to that of MRE and/or ileocolonoscopy within the terminal ileum and colon (P=0.500–0.625). The study by Tai et al. [102] showed that the use of panenteric capsule endoscopy resulted in management change in 46.5% of cases. Overall, the presence of active inflammatory findings resulted in a change in medical management in 64.6% of patients with established CD. Proximal SB findings led to an upstaging of disease in 19.7% and predicted escalation of therapy (OR 40.3). Similarly, in a prospective comparative study of panenteric capsule endoscopy and ileocolonoscopy by Leighton et al. [138] in patients with active CD, panenteric capsule endoscopy was shown to have a higher lesion detection rate in all SB segments including the terminal ileum.

Despite recommendation by new guidelines that all patients newly diagnosed with CD undergo SB assessment by ultrasound, MRE, and/or SBCE [90], it is still not clear whether these techniques are alternative or complementary. Evidence is scarce, but Greener et al. [139] compared the changes in disease extent and localization after performing MRE, SBCE, and both modalities. The investigators demonstrated that previously unrecognized disease locations were detected with SBCE and MRE in 51% and 25%, respectively (P<0.01) and by both modalities combined in 44 patients (55%). Using both modalities together may alter the original Montreal classification in 64% of patients [139].

For patients with established CD, the use of SBCE and panenteric capsule endoscopy may lead to changes in management in 50%–60% of patients [102,140], as they allow assessment of mucosal healing [141]. Indeed, in a meta-analysis by Niv [142], mucosal healing detection by capsule was shown to be a good predictor of long-term clinical remission.

Although the Lewis score and the Capsule Endoscopy Crohn’s Disease Activity Index (CECDAI) have shown good correlation with each other [142,143], there seems to be poor correlation between capsule activity index scores and clinical and laboratory parameters. The study by Kopylov et al. [144] emphasizes that SBCE may detect mucosal inflammation even in patients in clinical and biomarker remission. Furthermore, a Lewis score of ≥270 has been identified as a predictor of...
The invention of capsule endoscopy introduced the need for quantitative metrics to assess mucosal inflammation. Furthermore, as treatment targets focus on mucosal healing, this has become even more essential. Several quantitative inflammatory scores for capsule endoscopy have been developed over the years [1, 141–143]. Regarding SBCE reporting, along with the Lewis score and CECDAI, a new activity index, the Eliakim score combining evaluation of SB and colonic findings, has been proposed. When panenteric capsule endoscopy is used to allow for an integrated assessment of the small bowel and the colon, the Eliakim score has shown a good correlation with the Lewis score [150].

**RECOMMENDATION**
ESGE recommends the use of activity scores (such as the Lewis score and the Capsule Endoscopy Crohn’s Disease Activity Index (CECDAI)) to facilitate prospective small-bowel capsule endoscopy follow-up of patients for longitudinal assessment of small-bowel Crohn’s disease and its response to medical therapy (using mucosal healing as an endpoint).
Strong recommendation, low quality evidence.

Capsule retention is the main adverse event of SBCE. As stated in the previous guideline [1], the recommendation is that asymptomatic patients should be managed conservatively/medically in the first instance, with DAE retrieval reserved for cases of persistent retention. Large series published since 2015 [1] have confirmed the validity of this recommended strategy. A multicenter retrospective study by Fernández-Urién...
et al. (n = 5428; different indications for SBCE) [154] showed an overall retention rate of 1.8%; >50% of retained capsules passed with conservative management (37% spontaneously; 20% with concomitant medical therapy). Nemeth et al., 2 years later also demonstrated a favorable outcome with this strategy: medical management resulted in the passage of 24% of retained capsules, while endoscopic retrieval was required in 44% [155]. This recommendation was also supported by the findings of another large retrospective series (n = 5348; all indications) [156] and a retrospective study focused on patients with established CD, which also reported a high rate (70.5%) of passage of retained capsules with conservative measures [157].

The evidence to support specific medical management regimens remains scant, albeit most series reported on the use of glucocorticoids for capsule retention in the context of CD [154, 155, 157], with immunomodulators also used as an alternative [157]. Published egestion rates with medical management range from 10% to 70% [155–157], being higher in patients with established CD. In a multivariate analysis published by Lee et al. [158], the presence of abdominal symptoms after capsule retention was an independent predictive factor for a surgical outcome (OR 18.56, 95% CI 1.87–183.82; P = 0.013).

Endoscopic retrieval has been a safe alternative in asymptomatic patients or in those with slight symptoms. Recently, a systematic review of 12 studies (n = 150) regarding the use of DBE for retrieval of retained capsules [159], demonstrated a pooled retrieval success rate of 86.5% (95% CI 75.6%–95.1%). Factors associated with higher success were the antegrade approach (74.7% vs. 26.3%; P < 0.001) and the presence of malignant strictures (100.0% vs. 78.3%; P = 0.043) [159].

RECOMMENDATION
ESGE recommends device-assisted enteroscopy if small-bowel endotherapy is indicated (including dilation of Crohn’s disease small-bowel strictures, retrieval of a retained capsule, and/or treatment of small-bowel bleeding).
Strong recommendation, high quality evidence.

Since the publication of the 2015 ESGE guideline [1] the evidence favoring the effectiveness and safety of DAE-facilitated endoscopic balloon dilation (EBD) of CD SB strictures has strengthened. This is best summarized in a recent meta-analysis by Bettenworth et al. [160], which evaluated 18 studies including a total of 463 patients and 1189 endoscopic balloon dilations. The pooled per-study analysis demonstrated that the technical success of endoscopic balloon dilation was 95% (95% CI 86.7%–98.1%; 13/18 studies), with clinical efficacy in 82.3% of patients (95% CI 68.1%–91%; 9/18 studies) in the short term. The major complication rate (including bleeding, perforation, and emergency surgery) was 5.3% (95% CI 3.5%–8.1%; 14/18 studies). Longer-term outcomes (as reflected by 20.5 months of follow-up) showed that symptomatic recurrence had occurred in 48.3% of patients (95% CI 33.2%–63.7%; 11/18 studies).

Nonetheless, this was managed by repeat endoscopic balloon dilation in 38.8% of patients (95% CI 27%–52%; 16/18 studies); recourse to surgery was required in 27.4% (95% CI 21.9%–33.8%; 15/18 studies). This meta-analysis [160] further interrogated detailed data from four of the included high volume centers (218 patients; 384 dilations) to identify potential risk factors associated with outcomes. On per-patient-based multivariable analysis, active SB disease was associated with reduced short-term clinical efficacy (OR 0.32, 95% CI 0.14–0.73; P = 0.007). Furthermore, concomitant active disease of the small and/or large bowel increased the risk for surgery (hazard ratio [HR] 1.85, 95% CI 1.09–3.13; P = 0.02; and HR 1.77, 95% CI 1.34–2.34; P < 0.001). Conversely, ongoing anti-TNF-alpha treatment at the time of dilation correlated with reduced re-intervention (HR 0.78, 95% CI 0.63–0.96; P = 0.019).

Based on the current evidence, an algorithm for the endoscopic management of SB strictures is suggested in Fig. 3 [161, 162].
Inherited polyposis syndromes

Familial adenomatous polyposis

RECOMMENDATION

ESGE recommends surveillance of the proximal small bowel in familial adenomatous polyposis, using conventional forward-viewing and side-viewing endoscopes. Strong recommendation, moderate quality evidence.

Peutz–Jeghers syndrome

Most polyps are localized within the small bowel in patients with Peutz–Jeghers syndrome (PJS). Patients have a significant risk of non-neoplastic complications (intussusception, bleeding, anemia) as well as an increased risk of malignancies (intestinal and extraintestinal) [169]. SB surveillance in PJS aims to prevent polypl-related complications (by reduction of the polypl burden) and to detect early premalignant or malignant changes with advancing patient age.

Guidelines from ESGE and the European Society for Paediatric Gastroenterology, Hepatology and Nutrition (ESPGHAN) recommend starting SB surveillance no later than 8 years of age (and earlier in patients with symptoms or complications) [166, 170]. Based on the number and size of SB polyps, a 1–3-yearly surveillance interval is recommended [166]. Cancer risk is significantly increased in PJS [171]. However, the potential for malignant transformation of the SB hamartomas remains unknown.

SB surveillance should be a part of the complex multiorgan screening program for patients with PJS [169]. SBCE is superior at detecting SB polyps in comparison with small-bowel follow-through and standard CT scans [168, 172]. The direct comparison of MRE and SBCE shows at least equivalent sensitivity of both methods in detection of SB hamartomas; there is some risk of missing clinically relevant polyps with both techniques [173, 174]. Some data suggest better localization of polyps and more accurate size estimation with MRE [173, 174], but SBCE superiority for detection of small polyps (<15 mm) [174]. A meta-analysis of 15 comparative studies (821 patients) of DAE and SBCE confirmed high concordance (93%) in the identification of SB polyps and tumors [172]. In a retrospective multicenter study, 25 patients underwent SBCE followed by DBE when treatment was indicated. Authors found a strong agreement for polypl location and size but not for the number of polyps; DAE was more accurate for the former [175]. Two small studies reported high concordance of MRE with DBE, laparoscopic enteroscopy, or surgery (93%). They also showed comparable diagnostic yields from MRE and DBE for SB polyps >15 mm [176, 177].

In summary, MRE, SBCE, and DAE are complementary methods with similar diagnostic yields and a similar risk of missed lesions. The limited data do not allow preference for any one of the methods. Thus, both noninvasive techniques (SBCE or MRE) can be recommended for SB surveillance in patients with PJS, based on local availability and experience.

A patient history of SB resection (and therefore a risk of intra-abdominal adhesions) may mean a higher risk of SBCE retention, especially in patients with obstructive symptoms [178]. The routine use of the patency capsule [179] is not recommended in PJS and should be considered only on a case-by-case basis.
**Juvenile polyposis**

**RECOMMENDATION**
ESGE recommends that routine evaluation of the small bowel in juvenile polyposis patients should be limited to the duodenum and based on flexible forward-viewing enteroscopy.

Strong recommendation, low quality evidence.

Involvement of the small bowel in juvenile polyposis seems infrequent and mainly limited to the duodenum in patients harboring a SMAD4 mutation [192, 193]. No case of SB cancer has been reported at this time in the well-characterized juvenile polyposis family. The ESGE 2019 consensus and the recent pediatric consensus on genetic syndromes do not recommend using SBCE or DAE in juvenile polyposis syndrome [166, 194].

In conclusion, there is no evidence of the usefulness of capsule enteroscopy and no published case of histologically proven juvenile polyposis in the distal small bowel in these patients. According to ESGE and ESPGHAN recommendations, duodenoscopy appears sufficient, specifically in SMAD4 mutation carriers, because of the frequency of duodenal polyps.

**Small-bowel tumors**

**RECOMMENDATION**
ESGE recommends the use of small-bowel capsule endoscopy in patients where there is an increased risk of a small-bowel tumor.

Strong recommendation, moderate quality evidence.

Most SBTs are detected during work-up for SSBB or unexplained IDA but are the cause in only about 3.5%–5% of these patients, making these symptoms weak predictors. Some subsets of patients have an increased risk of SBT, such as those with liver metastases of previously undiagnosed primary neuroendocrine tumor, stage IV malignant melanoma, or stage III malignant melanoma with positive FOBT, or with nonresponsive/complicated celiac disease (see Celiac disease section) [19]. In contrast, recent data do not suggest a significant yield for SBT or polyps in patients with sporadic duodenal adenomas [195], long-standing SB CD [196], or asymptomatic Lynch syndrome [197, 198]. The risk for underlying SBT does not seem to be higher in patients with recurring or ongoing bleeding than in patients with the first bleeding episode [199].

Because of the rarity of SBTs, prospective studies are lacking, and data are primarily retrospective from SSBB and IDA studies. In this setting, SBCE has exhibited good diagnostic performance for identifying SBTs [74, 200]. Although Johnston et al. have reported more frequent detection of SB malignancy at SBCE in younger patients (<55 years) [201], most studies did not consider the duodenum as a target for SBCE in patients with Peutz–Jeghers syndrome.

Strong recommendation, moderate quality evidence.

An SB polyp size >15 mm is the most important risk factor for SB intussusception, which can lead to intestinal obstruction and acute abdomen [180, 181]. On the other hand, in children (because of the smaller intestinal diameter), even polyps smaller than 15 mm may represent a risk, and polyps may result in other complications such as chronic bleeding with IDA [181]. Consequently, large (>15 mm), symptomatic, or rapidly growing polyps should be promptly removed.

Both in adults and children, DAE is clinically useful for diagnosis and relatively safe for therapy of SB polyps [180, 182–184]. In a study of 50 enteroscopies using the antegrade (84%) and retrograde (16%) approach, the therapeutic interventions resulted in complete clearance of polyps >10 mm in 76% of patients [184]. However, considering the safety profile of DAE polypectomy (complication rate in PJS patients: 4%–6% [183–185]), enteroscopy should be used only as a targeted approach after previous noninvasive SB examination (using SBCE or MRE).

Motorized spiral enteroscopy has only recently been used in patients with PJS [186]. The published data on this technique are promising but insufficient for a final recommendation for patients with PJS.

Various technical improvements, including underwater resection [187] and ischemic polypectomy using polyp strangulation with endoclips and/or detachable snare (possibly also with an underwater approach), have been reported [188, 189]. They could represent a safer and faster alternative to conventional polypectomy; however, their benefits need future verification. In some clinical situations (high polyp burden and incomplete polyp clearance during previous DAE), the direct indication for the next DAE (without repeated SBCE or MRE) can be considered in an individualized time frame. A gradual decline in polyp size, numbers, and complication rate can be expected in the course of surveillance and repeated DAE polypectomies [182, 185, 190, 191].

When a polyp is too large for safe removal with DAE or cannot be reached using this modality (because of adhesions), intraoperative enteroscopy as a complementary technique could be considered for SB evaluation and polypectomy [183, 184]. Combined treatment of SB hamartomas with device-assisted and intraoperative enteroscopy significantly increases clearance success by 16% [184]. This approach may reduce the need for future surgery and SB resection in PJS patients.
not reveal any significant differences in the incidence of SBTs depending on the age of the patients, albeit there were variations in the definition of the younger versus older age groups [202–204]. The diagnostic yields of double-balloon enteroscopy for SBTs in the SSBB setting were also similar between patients <65 years old and elderly patients (>65 years), except for cases of incomplete SB obstruction where a higher rate of adenocarcinoma was identified in the elderly group (19.4% vs. 7.1%, \( P=0.038 \)) [205].

In an RCT in the setting of SSBB, SBCE had a higher diagnostic yield for SBTs and polyps than push enteroscopy [206]. Compared to DAE in SSBB, SBCE had detection rates similar to single-balloon enteroscopy for SBTs [207,208]. Also double-balloon enteroscopy and SBCE had comparable diagnostic yields for SBTs [209,210], even in a context of SB re-examination, where double-balloon enteroscopy was compared to repeat SBCE for SSBB [56]. Nevertheless, the concordance between SBCE and single-balloon enteroscopy was not significant regarding SB masses [211], and the agreement between SBCE and double-balloon enteroscopy was lower for SBTs than for other SB pathology in the setting of SSBB [212,213]. Suspected SB neoplasia was related to increased diagnostic and therapeutic yield for both single- and double-balloon enteroscopy. Although previous SB investigations, including SBCE and/or imaging studies, improved the diagnostic yield of enteroscopy, this was not statistically significant [214].

On the other hand, the risk of false-negative SBCE results has been documented for SBTs, especially for lesions located in the proximal SB [168] or subepithelial tumors with minimal endoluminal components, such as GI stromal tumors (GISTs) [215] and neuroendocrine neoplasms (NEs) [216]. Therefore, in the case of a negative SBCE, albeit with a strong suspicion of an SBT, further dedicated SB cross-sectional imaging should be performed for confirmation.

Regarding imaging studies, CTE was accurate in raising the suspicion of SBTs [18], primarily when performed for SSBB [217]. CT angiography had a higher diagnostic yield for bleeding SBTs than for SB bleeding of nontumoral origin [218]. In a retrospective comparison of CTE and MRE, all cases of SBTs were accurately diagnosed by both modalities [219]. Conversely, in a prospective study comparing SBCE and CTE in the context of SSBB, the sensitivity of SBCE for SBTs was 66.67% compared to 100% for CTE [87]. In a retrospective study comparing double-balloon enteroscopy with SBCE and imaging modalities (CTE and MRE) for detecting SBTs, double-balloon enteroscopy was superior to all methods in terms of sensitivity, specificity, accuracy, and negative predictive value (NPV). Only CTE exhibited slightly higher PPV than double-balloon enteroscopy (93.5% vs. 90.0%) with comparable specificity, whereas MRE was outperformed in every aspect [220]. In another retrospective study comparing SBCE, double-balloon enteroscopy, and CTE for SSBB, all three approaches were comparable, complementing each other in detecting SBTs [221]. Thus, a combination of SBCE, dedicated cross-sectional SB imaging (e.g., CTE) and DAE may be required in the setting of suspected SBT since all three modalities are complementary to each other and provide supplementary information to establish the diagnosis of an SBT.

**RECOMMENDATION**

ESGE does not recommend, in the setting of suspected small-bowel tumor, specific investigations before small-bowel capsule endoscopy unless patients are considered to be at risk of capsule retention.

Strong recommendation, low quality evidence.

**RECOMMENDATION**

ESGE recommends consideration of device-assisted enteroscopy in preference to small-bowel capsule endoscopy if imaging tests have already demonstrated suspected small-bowel tumor.

Strong recommendation, low quality evidence.

The ESGE Technical Review on SBCE and DAE recommends that no specific investigations be routinely performed on every patient referred for SBCE unless they are considered at risk for capsule retention. Careful assessment of symptoms such as abdominal pain/distension, nausea/vomiting, a history of previous SB resection, abdominal/pelvic radiation, or chronic use of NSAIDs may be used to distinguish patients at a higher risk of capsule retention [47]. Ultrasound could be a noninvasive initial diagnostic option in these patients, as a sensitivity of >90% for SBTs >2 cm has been reported [222].

The capsule retention rate in the case of SBTs varies among studies [201,203]; nevertheless, in a meta-analysis, the capsule retention rate was 2.1% for patients with SSBB, representing the most common indication for SB investigations in patients with SBTs [16]. In the setting of suspected SBT in imaging studies, DAE should be preferred over SBCE to avoid capsule retention and acquire biopsies for histological diagnosis [1]. Furthermore, in the case of capsule retention, surgery remains the mainstay of treatment when neoplastic disease is unequivocally suggested, allowing both capsule retrieval and tumor resection [47]. If the nature of the SB lesion cannot be determined with certainty, then DAE can be an alternative for capsule retrieval and tissue sampling and/or endoscopic resection if deemed feasible in the case of benign tumors [159,223].

**RECOMMENDATION**

ESGE recommends cross-sectional imaging for staging and ascertaining operability when there is a small-bowel capsule endoscopy finding of a small-bowel tumor with high diagnostic certainty.

Strong recommendation, low quality evidence.
When SBCE findings strongly suggest an SBT (stenotic or protruding, ulcerated, bleeding mass lesion), direct surgical referral without preoperative histological diagnosis would be justifiable. In these cases, preoperative cross-sectional imaging is mandatory to provide further information on disease extent and resectability. If the underlying etiology of the tumor is uncertain (e.g., adenocarcinoma vs. lymphoma), tissue sampling through DAE is indicated to establish a histopathological diagnosis that may guide the course of subsequent management. When subepithelial protrusions or bulges of uncertain nature are identified on SBCE, further investigations (DAE or/and dedicated SB cross-sectional imaging) are warranted to avoid a false-positive diagnosis of subepithelial lesions such as GISTs or NENs. It should be noted that the prominent extraluminal component of GISTs may challenge endoscopic diagnosis, not only with SBCE but with DAE too. The effectiveness of histological confirmation by DAE in this setting has a wide range (46%–88%) [223–225]. Placement of a tattoo during DAE is mandatory to facilitate recognition of an SB mass lesion at subsequent (laparoscopic) surgery [1].

Regarding SB subepithelial lesions, CTE was shown to be superior to abdominopelvic CT for identifying SB GISTs [215] and SB NENs [226]. MRE has exhibited high degrees of sensitivity for the diagnosis of NENs > 10 mm (94%), but for lesions <10 mm, sensitivity was only 45% [227]. In a retrospective study assessing imaging techniques and double-balloon enteroscopy for the management of SB NENs, double-balloon enteroscopy was significantly better at identifying the primary tumor than CT, MRI, or somatostatin receptor imaging, as well as for detection of multifocal lesions when compared to CT and somatostatin receptor imaging but not compared to MRI [228]. Double-balloon enteroscopy also detected additional lesions in 62.2% of patients who underwent an evaluation to exclude multifocal disease in the setting of SB NENs [216].

In patients with treated follicular lymphoma, Nakamura et al. found that SBCE detected lesions at a similar rate to double-balloon enteroscopy; however, identifying residual lymphoma required biopsy, and the authors recommend DBE for follow-up [229]. Only 1 of 11 patients with an SBCE diagnosis of malignant SBT who underwent surgery had recurrent bleeding; in this patient, it was caused by metastasis of gastric and papillary cancer in familial adenomatous polyposis [230]. After complete resection of SB GIST in 32 patients, no intraluminal recurrence was seen during a median follow-up of 30 months (range 3–54 months) [225].

There are no studies that support regular follow-up of asymptomatic patients after resection of SBT in the absence of inherited polyposis syndromes.

Similarly, SBCE seems to have a very limited role in staging SBTs diagnosed with other techniques. SBCE and enteroscopy can help define the extent of GI non-Hodgkin’s lymphoma, although they do not change the stage of follicular lymphoma [231]. Similarly, the number of detected NENs in the small bowel could be increased without demonstrating an impact of multifocality on outcomes [216].

A summary of published reports on self-expanding metal stents (SEMs) placement by endoscopy (n = 69) in malignant SB strictures found the method to be safe and effective [232]. Recent small series confirmed this result. Clinical improvement was observed following SEMS placement but not with medical treatment [233]. DAE can also be applied for ink marking of malignant SB strictures for palliative surgery [234].

Celiac disease

ESGE does not recommend small-bowel capsule endoscopy to diagnose celiac disease. Strong recommendation, low quality evidence.
In studies assessing the utility/efficacy of SBCE in diagnosing celiac disease (i.e., ability to detect histologically proven villous atrophy), the sensitivity, specificity, PPV, and NPV of SBCE were 70%–100%, 64%–100%, 96%–100%, and 71%–93%, respectively [235–239]. All these studies consistently show that, in the presence of antiendomysial antibody (EmA) or significantly elevated antitransglutaminase antibody (tTG), the PPV and specificity for recognizing endoscopic markers of celiac disease are 100%. However, the high pre-test probability of celiac disease in all of these studies may be a potential limitation leading to an overestimation of SBCE performance. A later meta-analysis confirms the previous findings [240], and an RCT has demonstrated that frontal and lateral view capsules are equivalent in detecting villous atrophy [241]. From a clinical point of view, new data suggest that when upper endoscopy is impossible, a diagnostic pathway similar to the pediatric sequence, based upon serology, could also be applied in adults [242], further limiting the potential use of SBCE in this setting.

Consequently, the actual scenario does not support the use of SBCE in this setting (basically, patients with positive serology necessitating a histological confirmation of the diagnosis) and probably, when necessary, the adoption of serological criteria could avoid any endoscopic procedure to diagnose celiac disease. Although currently unproven, the use of computerized image enhancement could modify this situation in future [243].

As with the previous ESGE guideline [1], there is no new evidence supporting the use of SBCE to routinely map the extent of disease. However, two recent studies from Chetcuti Zammit et al. [244, 245] reported that the extent of villous atrophy could be efficiently verified by SBCE and atrophy extent could correlate with clinical parameters in some specific subgroups of patients (e.g., those with nonresponsive celiac disease, or severe bone involvement). The first study analyzed SBCE in 300 celiac patients and demonstrated an acceptable agreement among readers to define the severity of celiac disease [244]; the second analyzed a cohort of 80 celiac patients and showed that, in individuals with a relevant percentage of small bowel involved by villous atrophy, bone mineral density decreased significantly [245]; furthermore, bone mineral density did not correlate with histological severity of atrophy, underlining the potential relevance of atrophy extent. In conclusion, more recent studies suggest that atrophy extent could be efficiently quantified using SBCE and that this finding could correlate with some clinical parameters. However, because of the absence of other than gluten-free diet therapies for celiac disease, this factor is merely descriptive, and SBCE cannot be routinely recommended for this purpose. Nevertheless, this scenario could rapidly change in the near future once pharmacological therapies for celiac disease become available.

**RECOMMENDATION**

ESGE recommends using small-bowel capsule endoscopy in cases of equivocal diagnosis of celiac disease since it is essential for final diagnosis and therapy.

Strong recommendation, low quality evidence.

Celiac disease frequently presents a benign course with an optimal prognosis; however, up to 20% of patients show persistent or recurrent symptoms despite 6–12 months of following a strict gluten-free diet [246, 251]. This “nonresponsive” form of celiac disease requires a careful diagnostic work-up to detect the presence of neoplastic and neoplastic complications, such as refractory celiac disease (RCD), ulcerative jejunoileitis, enteropathy-associated T-cell lymphoma (EATL), and SB adenocarcinoma. RCD is defined by malabsorption and villous atrophy despite a correct gluten-free diet; RCD can be further subtyped into RCD type 1 (RCD-1) and type 2 (RCD-2) depending on the presence of an aberrant T-cell type in the duodenal mucosa, detected using cytology. RCD-2 is less frequent but characterized by a severe prognosis with mortality of up to 50% in 5 years and a higher risk of neoplastic evolution [252].
For these reasons, nonresponsive celiac disease and RCD-1 and RCD-2 warrant surveillance of the small bowel and early detection of neoplastic complications.

Previously, two studies evaluating patients with nonresponsive disease identified a few severe complications with SBCE [248, 253]. Focusing on RCD, Barret et al. [254] used SBCE to investigate disease severity in 29 RCD patients; notably, after tissue sampling with DAE, they diagnosed 3 cases of EATL and 5 cases of ulcerative jejunoileitis requiring specific treatment in the RCD cohort. The sequential approach, SBCE followed by DAE in the case of suspect findings, appears justified by the potentially relevant diagnosis (EATL and ulcerative jejunoileitis) and the importance of the consequent therapies [255, 256].

More recently, different studies have investigated the clinical use of SBCE and DAE in this setting, including a large number of patients in single-center and multicenter patient cohorts [256–261]. Notably, all these studies confirmed a diagnostic yield of SBCE close to 50%, with the detection of SBTs in 3%–10% of cases. SBCE represents the first-line investigation, while DAE is performed to obtain tissue samples that usually reveal an EATL or that can be used in cytofluorimetry to diagnose or monitor RCD.

Furthermore, two studies [257, 259] demonstrated that atrophy extent correlates with mortality more than histology does. In 40% of cases, SBCE findings were beyond the Treitz ligament and thus not accessible at upper endoscopy, underlining the pivotal role of SBCE/DAE in RCD. These findings have been strengthened by a recently published meta-analysis [262] demonstrating a diagnostic yield for malignancies and ulcerative jejunoileitis of 13% in the case of SBCE and 30% for DAE. Given the scenario described above, in the case of nonresponsive celiac disease or RCD, upper endoscopy and SBCE are mandatory; the first to take biopsies to perform routine histology, the second to detect other lesions to be targeted by DAE [263].

**Other indications**

**Chronic abdominal pain**

**RECOMMENDATION**

ESGE does not recommend small-bowel capsule endoscopy as the first-line investigation for patients with isolated chronic abdominal pain.

Strong recommendation, low quality evidence.

Chronic abdominal pain is usually defined as a constant or recurrent pain that lasts 3 months or more. Chronic abdominal pain without pathological findings in upper endoscopy, colonoscopy and/or imaging techniques is a prevalent condition [264].

Interestingly, many case reports and case series have described diagnosis by SBCE of significant pathologies in patients with chronic abdominal pain (e.g., Meckel’s diverticulum [265], eosinophilic enteritis [266], and SBTs [220]). However, the available evidence highlights that the probability of detecting significant findings at SBCE is very low (below 20%) when isolated chronic abdominal pain is the indication for SBCE. At the same time, this rises significantly when associated with signs/symptoms or altered laboratory findings.

Shim et al. [267] retrospectively analyzed 110 patients with unexplained chronic abdominal pain: diagnostic yield was 17.3%, and in multivariate analysis weight loss was a significant risk factor for positive findings at SBCE (OR 18.6, 95% CI 1.6–222.4; \( P = 0.02 \)). Katsinelos et al. [268] conducted an open-label prospective nonrandomized multicenter clinical trial. In this study, diagnostic yield was 44.4%, and in multivariate regression analysis positive findings from SBCE were associated with elevated erythrocyte sedimentation rate (ESR) (OR 67.9, 95% CI 9.3–310.6, \( P < 0.001 \)) and C-reactive protein (CRP) (OR 41.5, 95% CI 6.2–213.4, \( P < 0.001 \)). Huang et al. [269] conducted a retrospective study which included 341 patients with chronic abdominal pain. In this study, the diagnostic yield was 28.15%, and these features were positively associated with SBCE diagnosis: weight loss (OR 2.827, 95% CI 1.938–28.15%, and these features were positively associated with elevated erythrocyte sedimentation rate (ESR) (OR 4.025, 95% CI 3.178–6.892, \( P = 0.016 \)), and increased CRP (OR 7.539, 95% CI 5.365–11.723, \( P = 0.002 \)). More recently, Kim et al. [270] performed a meta-analysis showing that the presence of elevated CRP (OR 14.09, 95% CI 2.81–70.60, \( P = 0.001 \)) and ESR (OR 14.45, 95% CI 9.92–227.33, \( P = 0.06 \)) significantly increased the diagnostic yield of SBCE in patients with unexplained abdominal pain.

These data underscore how, on the one hand, the SB endoscopic evaluation plays a very limited role in cases of isolated abdominal pain and, on the other, how relevant it is in this subset of patients to plan a comprehensive diagnostic workup (including laboratory tests, imaging tests, and accurate collection of clinical history), since when abdominal pain is associated with other clinical features, SBCE may lead to establishing a definite diagnosis.

**Foreign body retrieval**

**RECOMMENDATION**

ESGE recommends device-assisted enteroscopy as an alternative to surgery for foreign bodies retained in the small bowel requiring retrieval in patients without acute intestinal obstruction.

Strong recommendation, moderate quality evidence.

SB foreign-body retention that needs intervention is a rare event. Most frequently the foreign bodies involved are endoscopy capsules or other medical devices (e.g., migrated plastic or metallic stents). Capsule retention is defined as a capsule remaining in the digestive tract for at least 2 weeks, and retention rates vary between 2.1% and 8.2% [16]. Previous abdominal surgery or SB disease (e.g., strictureing CD or SBT) may contribute to retention. A systematic review has shown that DAE is a...
reliable alternative to surgery, with a retrieval rate of 74.7% when the capsule is retained in the jejunum and can be reached via the antegrade approach [158]. However, when the capsule is retained in the ileum, the retrograde approach often necessitates endoscopic balloon dilation of the stricture before the capsule can be reached and is, therefore, less effective, as illustrated by a retrieval rate of only 26.3%. The serious adverse event rate is low (1.3% SB perforation risk) and associated with balloon dilation or neoplasia. One multicenter study reported that symptoms were the only independent predictor of successful retrieval using DAE (OR 13.40, 95% CI 1.10–162.56; P = 0.042) [271]. In addition to retrieving the retained capsule, DAE can also facilitate the diagnosis and treatment of the underlying intestinal disease, by endoscopic biopsy, endoscopic balloon dilation, and preoperative tattooing. However, the indication for endoscopic or surgical intervention should be evaluated on a case-by-case basis and depends on local availability and expertise.

DAE-assisted percutaneous endoscopic jejunostomy (PEJ) for enteral feeding

**RECOMMENDATION**

ESGE suggests that in patients requiring jejunostomy for enteral feeding, DAE-assisted percutaneous endoscopic jejunostomy (PEJ) is a possible alternative to surgical jejunostomy.

Weak recommendation, moderate quality evidence.

Direct percutaneous endoscopic jejunostomy (DPEJ) is an accepted alternative to nasojejunal or surgical jejunal feeding in patients who require long-term post-pyloric feeding [272].

DPEJ using an enteroscope has a technical success rate of up to 90%. Technical failures are reported mostly because of limited enteroscope advancement in patients with a history of abdominal surgery and adhesions. DPEJ by DAE has a significant adverse event rate of 3.5% [273–276]; these include bleeding and SB perforation. DAE-assisted PEJ can represent an alternative to surgical jejunostomy according to local availability and expertise.

**DAE-ERCP in patients with altered anatomy**

**RECOMMENDATION**

ESGE recommends DAE-ERCP as a first-line endoscopic approach to treat pancreaticobiliary diseases in patients with surgically altered anatomy (except for Billroth II patients).

Strong recommendation, moderate quality evidence.

Since the advent of DAE, multiple retrospective studies have been published on DAE-endoscopic retrograde cholangiopancreatography (ERCP) in patients with surgically altered anatomy. Biliary indications are more frequent than pancreatic indications. The most frequently met surgical reconstructions are Billroth II partial gastrectomy, Roux-en-Y total gastrectomy, Roux-en-Y gastric bypass (RYGB), Whipple’s pancreaticoduodenectomy (also with Roux-en-Y), and Roux-en-Y hepatopancreatjejunostomy [277]. According to ESGE guidelines [278], use of a side-viewing duodenoscope is the first option for performing ERCP in Billroth II patients. However, DAE-ERCP is equally effective [279].

Several recent meta-analyses on using long and short DBE, SBE, and manual spiral enteroscopy for performing ERCP in patients with altered anatomy, are based on multiple retrospective case series [280–284] (see Table 3s). They show that procedural success has seemed to increase over time, reaching >75% in the most recent meta-analysis, and even much higher success rates in individual retrospective series. DBE and SBE are equally effective. Short versions of both DBE and SBE have been developed, allowing the use of conventional ERCP accessories. Studies have shown equal procedural success when using short-type DAE, except in the cases of Roux-en-Y surgery without gastrectomy and long limb Roux-en-Y surgery such as RYGB, where the short-type DAE device may be too short to reach the biliopancreatic system [283, 285, 286]. Except for a single preliminary case report, there are currently no data available on the use of motorized spiral enteroscopy to perform ERCP in patients with surgically altered anatomy [287]. Overall, adverse events show low rates (up to 8% in meta-analysis reviews) and are mild with little indication for surgical intervention (mainly due to intestinal perforation), and mortality related to DAE-ERCP is close to 0%.

DAE-ERCP in patients with surgically altered anatomy can be considered a first-line technique to treat biliopancreatic pathology thanks to the good overall procedural success rate and the low adverse event rate. However, since the overall procedural success rate is good but not excellent, alternative, more invasive techniques have emerged, showing higher technical success and adverse event rates. Thanks to the excluded stomach in RYGB, multiple alternative approaches currently exist, including laparoscopy-assisted ERCP, endoscopic ultrasound (EUS)-directed transgastric ERCP, EUS-guided intrahepatic puncture with antegrade clearance, and percutaneous transhepatic biliary drainage [288, 289]. Both laparoscopy-assisted ERCP and EUS-directed transgastric ERCP have high (>90%) procedural success rates but also higher adverse event rates (12%–24%) [290]. Also, in patients with Whipple’s pancreaticoduodenectomy, transgastric EUS-guided drainage of the pancreatic duct is feasible with a good technical success rate of more than 70%, but with an adverse event rate of 20%–35% [291, 292]. ERCP in patients with surgically altered anatomy is challenging and should be referred to expert centers. The technique of choice depends on local availability and expertise, as previously suggested by ESGE [293].
Innovations

SBCE

Since their inception at the dawn of this millennium, SBCE and DAE have continually evolved. For the former, two main paths lead to further development. First, technological advances are expected to lead to paradigm shifts. Second, patient- and society-related outcomes may drastically change SBCE practice in the coming years.

The latest generation of commercially available SBCE devices and software currently provides high resolution images captured by powerful central processing units, an adaptive frame rate, post-processing chromoendoscopy options, long-life batteries (enabling gastroenteric or enterocolonic examinations) and expert systems (allowing faster reading) [294]. Implementation of AI in software is a significant step [295]. These solutions allow a drastic reduction (of around 90%) in image selection and reading time, while maintaining very high sensitivity (above 98%) for lesion detection [296, 297]. Further high level and expert systems (allowing faster reading) [294]. Implementation of AI in software is a significant step [295]. These solutions allow a drastic reduction (of around 90%) in image selection and reading time, while maintaining very high sensitivity (above 98%) for lesion detection [296, 297]. Further high level clinical assessment and discussions with scientific societies and regulatory authorities are required before AI can routinely be used in clinical practice. This allows the triage of normal videos and/or images within videos. Additionally, some AI software also enables characterization of abnormalities [297]. Researchers in AI are working to address the challenges of automated evaluation of anatomical landmarks, of completion, and of cleanliness [295]. In addition, progress in miniaturization and energy-saving may provide more room for batteries within the capsule and thereby longer battery life.

Consequently, it is expected that a genuinely “panenteric” (mouth-to-anus) capsule endoscope will be available in the near future [298]. In addition, magnetically guided capsule endoscopy has been developed and clinically assessed for examination of the stomach or combined stomach and small bowel [299, 300]. However, active capsules with embedded AI, microbiota or tissue sampling, or therapeutic options, are still in the early stages of development [300].

Furthermore, emerging healthcare and societal trends may profoundly modify how we practice SBCE. For example, some capsule endoscopy manufacturers have recently obtained approval from the US Food and Drug Administration for capsule home delivery, provided that a healthcare provider accompanies patients for the procedure [300]. As a result, patients’ comfort and reporting times would be significantly improved. In addition, there is growing concern regarding the ecological impact of endoscopy. Capsule endoscopy is expected not to escape the debate around avoiding the yearly release of tens of thousands of batteries and electronic material into the environment [300]. Overall, such developments may widen the indications for capsule endoscopy and how we practice SBCE in the future.

DAE

Motorized spiral enteroscopy

A novel motorized spiral enteroscopy device (Olympus, Tokyo, Japan) has recently been introduced. The activation of an integrated electric motor permits the rotational movement of a spiral overtube, achieving advancement by pleating the SB. Since its introduction, several case reports have been published, showing the potential abilities of this new endoscopy device. The first prospective trial was conducted in 132 patients from two European tertiary referral centers. It showed diagnostic and therapeutic yields for antegrade explorations similar to those from previous studies with balloon enteroscopy. However, longer insertion length (mean 450 cm, range 0–600 cm) in a shorter procedural time (mean 25 min, range 3–122 min) was achieved [301]. Two other clinical studies from Europe and Asia reported similar results; moreover, total enteroscopy rates were 61% and 70% [302, 303]. Nonetheless, some issues regarding this technique are still unclear, such as the need for general anesthesia for antegrade procedures, the learning curve, and the target population. Furthermore, only minimal information exists on the impact of prior major abdominal surgery on the feasibility and the safety of motorized spiral enteroscopy [304, 305].

Water-aided enteroscopy

The water-exchange intubation technique has been proposed to achieve higher total enteroscopy rates. The method is the same as when applied for the exploration of the colon, with warm saline (37 °C) infused into the intestinal lumen to maintain the endoscopic view and mostly suctioned during the insertion phase. During the antegrade procedure, saline is infused once the ligament of Treitz is reached, while during the retrograde procedure, water exchange begins from insertion at the anus [306]. Of note, an adaptor connecting the water pump tube to the accessory channel of the enteroscope is needed.

The two studies available so far have produced conflicting results. One randomized, nonblinded, single-center study compared the total enteroscopy rates between patients undergoing water-exchange-assisted (n = 55) and CO2-insufflated (n = 55) SBE [306]. The total enteroscopy rate was significantly higher in the water-exchange group (58.2% vs. 36.4%), as well as the overall and antegrade approach insertion depths, the overall insertion time, and the insertion time for the oral route. Diagnostic yields and adverse event rates were similar between groups. In a prospective, comparative and observational study, 46 patients were randomly allocated to water exchange-assisted (n = 23 patients) and CO2-insufflated (n = 23 patients) DBE. The median insertion depth was greater in the CO2 group, at 260 cm vs. 160 cm (P = 0.048). Multiple logistic regression showed a statistically significant difference in the insertion depth using CO2 insufflation (OR 1.009, 95%CI 1.001–1.017; P = 0.034). Adverse event rates were similar between groups [307]. Other larger RCTs comparing the water-exchange technique with CO2 are awaited.
Interventional enteroscopy

Snare and ischemic polyectomy, and conventional and underwater mucosectomy by DAE, have become the first-line treatments for SB polyps, especially in the setting of PJS. These techniques are efficient, safe and cost-effective. Complete resection rates are over 60%, with infrequent adverse events (mostly in the form of immediate or delayed bleeding and pancreatitis) [183, 184]. The outcomes of DAE dilation of benign SB strictures are mentioned in a previous section.

Disclaimer


Acknowledgments

ESGE wishes to thank external reviewers Prof. Ian M Gralnek of the Rappaport Faculty of Medicine Technion Israel Institute of Technology, Haifa, Israel, and Prof. Owen Epstein of the Royal Free Hospital, London, UK, for their critical review and appraisal of this Guideline. We would like also to thank the following ESGE members who reviewed the Guideline and made interesting suggestions for improvement: Abdulbaqi Al Toma, Rafael Barreto Zuñiga, Gerardo Blanco Velasco, Rosamaria Bozzi, Alessandro Rimondi, Stylianos Stylianidis, Tony Tham, Olga Bednarska on behalf of the Swedish Society of Gastroenterology (SGF), and Rodica Cincul on behalf of the French Society of Digestive Endoscopy (SFED).

Competing interests

C. Carretero provides consultancy and receives speaker fees from Medtronic (ongoing). X. Dray is a founder of and shareholder in Augmented Endoscopy (May 2019 to present); he is a member of the International Capsule Endoscopy REnseignements (iCARE) group (December 2021 to present); he holds four patents (shared with his institutions) related to artificial intelligence in endoscopy. E. J. Despot has received educational grants in support of conference organization, and honoraria, from Fujifilm, Pentax, and Olympus (2017–2021), and honoraria from Ambu (2021). L. Elli has held a lecture/consultancy role for Medtronic (2018-2020) and Capsocam (2016). L. Fuccio is a Co-Editor of Endoscopy journal. M. Keuchel has received speaker's fees and travel support from and provided consultancy to Medtronic, and received speaker’s fees from Olympus (both from 2021 to present); his department has received study support from AnX Robotics (from 2021 to present). A. Koulouzidis is a co-founder and shareholder of AJM MED-i-Caps (from 2017, ongoing) and iCERV (from 2022, ongoing), and has received consultancy fees from Chi and Jinshan Science & Technology and lecture honoraria from Medtronic (all from 2020, ongoing), travel assistance fees from Aquilant (2019), material support for clinical research from SynMed and Intromedic (2016–2020), and lecture honoraria and AB meeting fees from Dr Falk Pharma UK (2016–2020); and has participated in an advisory board for Ankon (2019); his department has received a grant from Medtronic (2016–2020); he is a founding and board member of iCARE; he or his department holds a patent related to this Guideline. D. McNamara received an iCloud Capsule Platform introductory fee waiver from Medtronic (2021–2022). T. Moreels received speaker’s fees from Olympus (2019–2022). H. Neumann is a consultant to Fujifilm, Medtronic, and Jinsha (from 2020, ongoing); his department receives study support from Fujifilm (from 2020, ongoing). M. Pennazio received speaker’s fees from Medtronic, Olympus, and Alfascigma (2015–2019). E. Perez-Cuadrado-Robles provided consultancy to Boston Scientific (2020–2021). E. Rondonotti has been an expert group member and speaker for Fujifilm (January 2021 to December 2021) and provided consultancy to Medtronic (2021); his department received a research grant from Fujifilm (January 2021 to December 2021). B. Rosa provided consultancy to Medtronic (2020–2021). C. Spada provided consultancy to Medtronic (2017–2022) and AnX Robotics (2020–2022). J. C. Saurin provided consultancy to Intromedic, Capsocam, Medtronic, and Povepharm (2021–2024), and teaching for Medtronic (2021–2024). I. Tacheci is Scientific Secretary to the Czech Society of Gastroenterology and responsible for dissemination of guidelines (2022). P. Cortegoso Valdivia, B. Gonzalez Suarez, L. Kuvovsky, E. Perez-Cuadrado-Martinez, S. Piccirelli, D.S. Sanders, R. Siddhu, K. Triantafyllou, and E. Vlachou have no competing interests.

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