Endoscopic diagnosis of sessile serrated polyp: A systematic review

Hiroshi Kashida

Department of Gastroenterology and Hepatology, Kindai University, Osaka, Japan

The aim of the present review was to clarify how we should detect and diagnose sessile serrated polyps (SSP) endoscopically. A systematic search was conducted of MEDLINE from January 2004 through March 2018. Nine findings: (i) proximal location; (ii) size >10 mm; (iii) irregular shape; (iv) indistinctive border; (v) cloud-like surface; (vi) mucus cap; (vii) rim of debris in white-light endoscopy; (viii) dilated vessels; and (ix) dilated crypts (pits) in image-enhanced endoscopy were considered to be candidate discriminators of SSP from hyperplastic polyps. Prospective studies in a general setting are warranted to validate the above-mentioned endoscopic features of SSP during real-time colonoscopy and to determine whether these features are useful for the differential diagnosis of SSP.

Key words: differential diagnosis, endoscopy, sessile serrated adenoma, sessile serrated polyp, sessile serrated polyp/adenoma

INTRODUCTION

More than 20 years have passed since Longacre et al., Torlakovic et al. and Snover et al. reported on serrated lesions in the colorectum which are now divided into hyperplastic polyp, traditional serrated adenoma, and sessile serrated adenoma/polyp (SSA/P) with or without dysplasia by the WHO classification system. Sessile serrated adenoma/polyp is considered to be an important precursor of colorectal cancer which remains one of the leading causes of cancer deaths worldwide. However, Payne et al. described that the term “serrated” never appeared in pathology reports at 10 of 32 centers. There is significant interobserver variation among pathologists in identifying and classifying serrated lesions. Even the terminology for serrated lesions may be subject to change. In this article, the terms conventional adenomatous polyp (AP), traditional serrated adenoma (TSA), sessile serrated polyp (SSP) and hyperplastic polyp (HP) will be used.

There is a recent tendency that the detection rate of SSP is increasing. Reasons for this phenomenon are attributed not only to the use of new endoscopic techniques but also to the increased awareness of endoscopists and pathologists concerning the significance and appearance of SSP. Detection rates of SSP are also known to be endoscopist-dependent. The true prevalence of SSP is not evident due to lack of knowledge, difficulty in detection and unestablished diagnostic criteria. Among others, endoscopic discrimination between SSP and HP remains a problem. The role of advanced imaging techniques (IEE) such as narrow-band imaging (NBI) in recognition of these lesions is also not well established. Use of an appropriate classification system including IEE would facilitate the recognition of SSP. The aim of the present review was to clarify specific characteristics and defining features that can help distinguish SSP from HP.
original papers were selected and reviewed in the present study (Table 1).18–32

FINDINGS IN WHITE-LIGHT ENDOSCOPY

HASEGAWA et al.18 conducted an investigation on 107 sessile serrated lesions (31 32 TSA, 22 SSP and 54 HP) in order to clarify endoscopic features of sessile serrated lesions and to determine whether SSP can be distinguished from other serrated lesions. Mean lesion size of SSP was larger than that of HP (14.2 vs 6.2 mm; P < 0.01). SSP was located more often in the proximal colon than in the distal colon (81.8% vs 18.2%), whereas HP was located less often in the proximal colon than in the distal colon (44.4% vs 55.6%). Limitation of this study is the small number of subjects and the retrospective nature.

Kashida et al.19 found 673 (6.0%) serrated lesions out of 11 253 colorectal localized lesions that were resected in their institute. Among them, those >5 mm (232 TSA, 35 SSP, and 95 HP) were evaluated. Average size of lesion was substantially bigger in SSP than in HP (14.0 ± 7.1 vs 7.9 ± 3.6 mm, P < 0.001). SSP were mostly located in the proximal colon, whereas the majority of HP were located in the distal colon.

Yamada et al.20 conducted a multivariate analysis among 242 lesions (118 SSP and 124 HP) on the clinical features of SSP compared with HP. They found that proximal location, size ≥10 mm, and sessile morphology were independent features of SSP, with odds ratio (OR) (95% CI) of 18.3 (7.2–46.2), 26.6 (10.6–66.4) and 4.3 (1.8–10.2), respectively.

Yamada et al.21 retrospectively examined 202 serrated lesions of the colorectum (41 TSA, 57 SSP and 104 HP). SSP were more predominantly located in the proximal colon and significantly larger than HP (P < 0.01). Polyp surface was granular or nodular in 40% of the SSP, more frequently than in HP (P < 0.01). Margins were more irregular or vague in SSP than in HP (P = 0.02). “Mucus-coated

Table 1 Study design of the articles included in the present study

| Reference/First author | Study design | No. of SSP | Modality | Evidence level |
|------------------------|--------------|------------|----------|----------------|
| Hasegawa18              | Retrospective cohort, single center | 22 | WLE, chromo, zoom | Low |
| Kashida19               | Retrospective cohort, single center | 35 | HR-WLE, chromo, zoom | Moderate |
| Yamada20                | Retrospective cohort, single center | 118 | NBI, zoom | Moderate |
| Yamada21                | Retrospective cohort, single center | 19 | WLE, chromo | Moderate |
| Ishigooka22             | Retrospective cohort, single center | 57 | WLE, chromo, zoom | Moderate |
| Tadepalli23             | Retrospective cohort, single center | 158 | WLE, NBI | Moderate |
| Bouwens24               | Retrospective cohort, single center | 170 | HR-WLE, i-scan, chromo | Moderate |
| Lee26                   | Retrospective cohort, single center | 634 | HR-WLE | Low |
| Kawasaki26              | Retrospective cohort, single center | 80 | HR-WLE, NBI, chromo, zoom | Low |
| Hazewinkel27            | Retrospective cohort, single center | 50 | HR-WLE, NBI | Moderate |
| IJspeert28              | Validation study, 10 experts | First 15 | HR-WLE, NBI | High |
|                        |                        | Second 11 | HR-WLE, NBI | High |
| Nakao29                 | Retrospective cohort, single center | 46 | HR-WLE, NBI, zoom | Low |
| Uraoka30                | Prospective, multicenter | 38 | HR-WLE, NBI, chromo, zoom | High |
| Yamashina31             | Retrospective cohort, single center | 39 | NBI, zoom | High |
|                        | Validation study, 12 experts | 51 | HR-WLE, chromo, zoom | High |
| Kimura32                | Retrospective and prospective, single center | Training 69 | Validation 55 | High |

HR-WLE, high-resolution white-light endoscopy; NBI, narrow-band imaging; SSP, sessile serrated polyp.
appearance” was observed in nine of 57 SSP, but only in one of 104 HP \((P < 0.01)\).

Ishigooka et al.\(^{31}\) investigated 112 lesions including 39 TSA, 50 SSP and 23 HP. SSP were larger in diameter than HP (13.62 ± 8.62 \(\text{mm} \) vs 7.74 ± 3.24 \(\text{mm} \)), more commonly located in the proximal colon (84.0% vs 30.4%), flat-elevated (90.0% vs 30.4%), orthocolor or pale (80% vs 34.8%), and with a larger amount of mucus (72.0% vs 21.7%).

Tadepalli et al.\(^{23}\) retrospectively analyzed 158 proximal SSP in 124 patients, 47.1% of which were larger than 10 mm in diameter. By Paris Classification,\(^{32}\) all lesions were type 0; 1.9% were protruded 0-Ias, 96% were slightly elevated (0-Ila), and 4% were truly flat (0-IIb). They evaluated the prevalence of seven morphological findings in 158 SSP and in 40 AP. Prevalent characteristics of SSP were mucus cap (63.9%), rim of debris/bubbles (51.9%), alteration of fold contour (37.3%), and obscured blood vessels (32.3%). In the review of 69 videos with 69 SSP and 31 AP, the most frequent keys to the detection of SSP were mucus cap (24.6%), alteration of fold contour (24.6%), and rim of debris/bubbles (21.7%). Some of the most frequent findings such as alteration of fold contour were witnessed in both SSP and AP; therefore, these findings cannot be used as discriminators between SSP and AP. Moreover, it would be distracting to discuss this point further when the main purpose of the present study is discrimination between SSP and HP.

Bouwens et al.\(^{24}\) retrospectively examined 170 SSP based on Tadepalli’s criteria.\(^{23}\) The lesions showed obscured blood vessels in 54.7%, alteration of fold contour in 5.3%, rim of debris/bubbles in 40.5%, mucus cap in 32.6% and pale color in 19.4%. The major aim of the study was to clarify endoscopic characteristics of SSP with dysplasia but there was no significant difference of endoscopic appearance between those with and without dysplasia except that dome-shaped elevation was more often witnessed in the former.

SSERRATED LESIONS MIMICKING DIVERTICULA

Lee et al.\(^{25}\) reported a group of SSP with a depressed surface. They accounted for seven out of 634 SSP. Kawasaki et al.\(^{26}\) found similar lesions in seven out of 80 SSP and in two out of 35 HP. The present author has also encountered many such cases of SSP. Whether or not these lesions with depressed surface are a subgroup of SSP is yet to be established. However, we should be aware of this type as there is a potential risk of misdiagnosing them as colonic diverticula.

These lesions look similar to so-called “inverted hyperplastic polyps” in previous reports,\(^{24–38}\) most of which had been published before the concept of SSP became popular and well-recognized. Presence of colonic crypts in submucosa has been also termed as herniation of crypts, pseudoinvasion, or epithelial-misplacement. Hu et al.\(^{39}\) conducted a prospective pathological investigation. Among 2560 colorectal polyps, frequencies of herniation of crypts were 1.79% (10/559) in SSP, 0.2% (3/1487) in HP and 0% (0/514) in polyoid normal tissue. Although the specificity of herniation of crypts for diagnosing SSP (vs HP and polypoid normal tissue) was 99.85% (1998/2001), its sensitivity was only 1.79% (10/559). Their multivariate analyses identified an independent association between herniation of crypts and diagnosis of SSP (OR = 11.47, \(P = 0.009\) for vs HP). Note that diverticulum-mimicking SSP may not necessarily be associated with the presence of crypts in the submucosal layer.

The present author has also encountered many such cases of SSP. There is no convincing evidence to conclude that “diverticulum-mimicking appearance” is a feature of SSP versus HP because the number of study subjects was small and the overall incidence of this finding was low. However, the incidence might be underestimated as a result of insufficient awareness of the phenomenon by most endoscopists.

NARROW-BAND IMAGING WITHOUT MAGNIFICATION

In the study conducted by Rex et al.,\(^{40}\) 800 patients were randomized to the NBI arm and to the white light (WL) arm. The authors concluded that NBI may increase the detection of proximal serrated lesions, although the result in this study was not statistically significant.

Boparai et al.\(^{41}\) examined seven patients with serrated polyposis syndrome (SPS) using high-resolution (HR) WL and NBI. In total, subjects had 15 AP, 32 SSP and 19 HP. Endoscopic differentiation between AP and HP was possible with the surface pattern and vessel pattern in NBI, but that between SSP and HP was unsatisfactory.

In the study by Tadepalli et al.,\(^{23}\) NBI was carried out in 62 SSP. In 43 (69.4%) SSP, NBI enhanced the visibility of SSP compared with WL. The enhancing effect of NBI was most frequently noticed in the SSP with mucus cap (38/45, 84.4%), those with rim of debris (31/37, 83.8%) and those with obscured blood vessels (17/22, 77.3%).

In the study by Hazewinkel et al.,\(^{27}\) HR-WLE and NBI pictures of 150 polyps (50 AP, 50 SSP and 50 HP) in 45 patients with SPS were systematically assessed using seven potential endoscopic features of SSP as follows: indistinctive border, cloud-like surface, dark spots inside the crypts, irregular shape, absence of tiny microvessels crossing the surface, pit pattern II SSA/P (mixture of open crypts and small elongated star-shaped pits), and vascular pattern intensity (VPI).
Multivariate analysis showed that indistinctive borders (OR, 3.11; 95% CI, 1.57–6.15) and cloud-like surface (OR, 2.65; 95% CI, 1.21–5.78) were independent predictive findings of SSP in HR-WLE. In NBI cloud-like surface (OR, 4.91; 95% CI, 2.42–9.97), irregular shape (OR, 3.17; 95% CI, 1.59–6.29), indistinctive borders (OR, 2.38; 95% CI, 1.14–4.96), and dark spots inside the crypts (OR, 2.05; 95% CI, 1.02–4.11) were independent predictive factors for SSP.

A lesion was presumed to be SSP if all of six SSP features were present, whereas it was considered to be HP if all of these features were absent. Using these conditions, sensitivity and specificity were 75% and 79% in HR-WLE and 89% and 96% in NBI, respectively. In their study, only serrated polyps that showed either all or no independent features were compared.

Ijspeert et al. developed the Workgroup serrAted polyps and Polyposis (WASP) classification. The polyps were first evaluated using the NBI International Colorectal Endoscopic (NICE) criteria in order to divide them into HP-like lesions (Type 1 polyps) and those resembling AP (Type 2 polyps). Secondly Hazewinkel’s criteria for SSP were used to differentiate between SSP and HP for Type 1 polyps, and between SSP and AP for Type 2 polyps. Presence of at least two out of four SSP features was considered sufficient for the diagnosis of SSP.

Ten consultant gastroenterologists were asked to participate in the training. Location and size of polyps were not presented. Overall accuracy was 0.63 before training, which increased significantly to 0.79 after training (16% improvement; 95% CI: 9–22%; P < 0.001). In the polyps diagnosed with high confidence, overall accuracy significantly increased from 0.73 to 0.87 (14% improvement; 95% CI 4–24%; P < 0.01).

**NARROW-BAND IMAGING WITH MAGNIFICATION**

Nakao et al. studied 46 SSP and 25 HP with magnifying endoscopy with NBI (M-NBI). Frequency of the findings with M-NBI was as follows: capillary dilatation: 11% in SSP and 4% in HP; mucus cap: 94% in SSP and 60% in HP; and pit dilatation: 80% in SSP and 28% in HP. In the distinction of SSP from HP, sensitivity and specificity were 10% and 96% for capillary dilatation, 94% and 40% for mucus cap, and 80% and 72% for pit dilatation, respectively.

Uraoka et al. investigated 89 lesions including 38 SSP (median size: 17 mm), and 41 HP (median size: 11 mm). They evaluated: (i) location; (ii) size; (iii) macroscopic type; (iv) adherent mucus; (v) brownish area in non-magnified NBI (N-NBI); (vi) “varicose microvascular vessel (VMV)” in magnified NBI (M-NBI); (vii) depression in chromoendoscopy (indigocarmine); (viii) “expanded Type II (E-II) pit pattern”; and (ix) “IIHI pit pattern”, using high-magnification chromoendoscopy. Multivariate analysis showed that three factors: “VMV”, size ≥10 mm, and proximal location, were independent diagnostic factors for SSP with OR of 8.2 (P = 0.001), 7.2 (P = 0.0017), and 6.1 (P = 0.0041), respectively. Sensitivity and specificity of “VMV” were 87.8% and 57.9%, respectively. If a lesion was considered SSP when two or more criteria were positive, sensitivity was 89.5% and overall accuracy was 82.3%.

Yamashina et al. conducted a pilot study with M-NBI and detected brownish, oval, “expanded crypt openings (ECO)” and “thick and branched vessels (TBV)” as diagnostic features of SSP. They then prospectively validated the criteria in 125 polyps with NICE Type 1 (51 SSP and 74 others) with an average size of 4–7 mm. Sensitivity and specificity of ECO for SSP were 84.3% and 81.1%, whereas those of TBV were 45.1% and 68.9%, respectively. If a lesion presumed to be SSP when ECO or TBV or both were present, the sensitivity and specificity were 98% and 59.5%, respectively.

Yamada et al. conducted a study using M-NBI and selected five features: dilated and branching vessels (DBV), irregular dark spots (iDS), regular network pattern, disorganized network pattern, and dense pattern as potential M-NBI features of SSP. Multivariate analysis among 242 lesions estimated that DBV only was the criterion for discriminating SSP from HP. They then systematically validated the diagnostic accuracies by using the pictures of 40 lesions (19 SSP and 21 HP). Sensitivity and specificity of DBV were 79% and 67%, respectively, whereas those of iDS were 42% and 81%, respectively. Assuming that a lesion is SSP when both DBV and iDS were present, the sensitivity, specificity and OR (95% CI) by univariate analysis were 26%, 95% and 7.14 (0.75–68.0), respectively. Assuming that a lesion is SSP when either of DBV or iDS was present, the sensitivity, specificity and OR (95% CI) were 95%, 52% and 19.8 (2.22–176.6), respectively.

**CHROMOEENDOSCOPY WITH MAGNIFICATION**

Kimura et al. identified a pit pattern that was specific to SSP and named this “Type II-open (Type II-O)”. Pits with this type were similar to conventional Type II pits, but were larger and more rounded in shape, reflecting the dilated crypts. The authors hypothesized that an overproduction of mucin was the cause of this phenotype. Next, they prospectively analyzed an independent validation set (n = 116; 61 AP and 55 serrated lesions). The results showed that “Type II-O pattern” was highly predictive of SSP (sensitivity: 65.5% and specificity: 97.3%).

In the study by Ishigooka et al., magnified colonoscopy showed “Type II-O pit pattern” as characteristic of SSP.

© 2018 The Authors. Digestive Endoscopy published by John Wiley & Sons Australia, Ltd on behalf of Japan Gastroenterological Endoscopy Society.
(sensitivity: 83.7% and specificity: 85.7%). The pinecone-like pit pattern was characteristic of TSA (sensitivity: 96.7% and specificity: 89.9%). Note that Uraoka et al. expressed this pit pattern as “expanded Type II (E-II) pit pattern”.

**DISCUSSION**

It would be better to discuss the detection of SSP and its diagnosis separately; however, this was impossible because almost none of the reviewed articles did so. Actually, it is very difficult to completely separate these two processes in a real-world clinical setting: for example, interruption of mucosal capillary network can be a key to detect a flat lesion, but it is not a good discriminator among AP, HP and SSP. Mucus cap is a key to detect a serrated lesion and, at the same time, it is a key to diagnose an SSP.

**Figure 2** White light images of sessile serrated polyp. (a) An irregular-shaped lesion with cloud-like surface and mucus cap. (b) A flat lesion with a rim of debris. (c) A sessile lesion with an irregular shape, pale in color.

**Figure 3** Narrow-band imaging (NBI) of sessile serrated polyp (SSP). (a) Non-magnified NBI. Color of the flat-elevated lesion is the same as the surrounding mucosa. Several dilated vessels can be identified. (b) Magnified NBI. Several dilated crypts are evident.
Other endoscopic technologies such as Full Spectrum Endoscopy (FUSE) (EndoChoice, Inc, Alpharetta, GA, US) or ENDOCUFF™ (Arc Medical Design Ltd, Leeds, UK) have been attempted to improve the detection rate of SSP, but these are out of the range of the present article. I excluded the literature that discussed this point as it is related to improvement of polyp detection in general and is not specific for SSP.

Endoscopic characteristics of HP and SSP are summarized in Table 2. Endoscopic findings: (i) proximal location; (ii) size $> 10$ mm; (iii) irregular shape; (iv) indistinctive border; (v) cloud-like surface; (vi) mucus cap; (vii) rim of debris; (viii) dilated vessels; and (ix) dilated crypts (pits) are candidate discriminators of SSP from HP (Figs 2–5).

SSP can be small or diminutive. However, when a flat and pale-color lesion is more than 10 mm in diameter, it is more likely to be an SSP than a HP. A cloud-like surface is also expressed as a granular, nodular, or bumpy surface and looks like the surface of a cumulus cloud. A dilated vessel is also expressed as a TBV, DBV or VMV. Vessels are not prominent in SSP, but when a non-pericryptal, dilated tortuous vessel is present, the lesion is more likely to be an SSP than a HP. HP can rarely present with some vessels on the surface, but they are usually only lacy.

Dilated crypts seen with NBI are also expressed as dark spots inside the crypts, pit dilatation, ECO, or iDS. Dilated round pits seen with magnified chromoendoscopy are expressed as Type II-O or Type E-II. It is considered that dilated crypts seen with NBI and dilated round pits seen with magnified chromoendoscopy reflect the same feature of SSP and can be used almost as synonyms. Even in SSP it is not that all the pits are dilated, but only some of the pits are round and open. They are usually interspersed with star-like (Type II) pits or fern-like pits. This finding with endoscopy
Table 2 Endoscopic characteristics of HP and SSP

|                  | HP                                      | SSP                                      |
|------------------|-----------------------------------------|------------------------------------------|
| **White light**  |                                        |                                          |
| Location         | Distal                                  | Proximal^a                               |
| Size             | Usually <5 mm                           | Can be >10 mm^b                          |
| Shape            | Flat-elevated oval, circular             | Flat-elevated or sessile irregular^c     |
| Border           | Indistinctive-distinctive               | Indistinctive^d                          |
| Surface          | Smooth                                  | Cloud-like^e                             |
|                  |                                         | Diverticulum-like                        |
|                  |                                         | Mucus cap^f                             |
|                  |                                         | Rim of debris^g                          |
| Vessel           | None or lacy                            | None or dilated^h                        |
| Surface pattern  | Dark or white spots                     | Dilated crypts^i                         |
| Chromoendoscopy  | Star-shaped (Type II)                   | Dilated round pits^j or fern-like        |

^

-A Candidate discriminators of sessile serrated polyps (SSP) from hyperplastic polyps (HP).^  
-A cloud-like surface is also expressed as a granular, nodular, or bumpy surface and looks like the surface of a cumulus cloud.  
-A dilated vessel is also expressed as a thick and branched vessel (TBV), dilated and branching vessel (DBV) or varicose microvascular vessel (VMV).  
-Dilated vessels seen with narrow-band imaging (NBI) are also expressed as dark spots inside the crypts, pit dilatation, expanded crypt openings (ECO), or irregular dark spots (IDS).  
-Dilated round pits seen with magnified chromoendoscopy are expressed as Type II-open (Type II-O) or expanded Type II (Type E-II).  
-It is considered that dilated crypts seen with NBI and dilated round pits seen with magnified chromoendoscopy reflect the same feature of SSP and can be used as synonyms.

Corresponds to the pathological finding of SSP; most of the crypts are serrated with some of the crypts filled with mucus and dilated.

A meta-analysis of 13 studies carried out by Parikh et al. concluded that image-enhanced endoscopy cannot be recommended as a diagnostic tool for SSP. However, this meta-analysis encompassed different modalities of image-enhanced endoscopy including auto-fluorescence imaging (AFI). They described that NBI studies showed promise. More studies are needed to validate the NBI criteria for diagnosing SSP. Likewise, more investigations are required to prove the usefulness of new modalities such as Blue Laser Imaging, i-scan.

Recognition of dysplasia within SSP^4^ is another important concern, but it is out of the range of the present study. It requires another review article.

Prospective studies during real-time colonoscopy in an ordinary clinical setting among a large number of subjects are warranted to confirm the above-mentioned endoscopic features of SSP and to validate whether these features are useful for the detection and differential diagnosis of SSP.

**ACKNOWLEDGMENTS**

I thank DRS Masashi Komeda and Sz-Juan Shiu for their assistance in the selection and assessment of the references.

**CONFLICTS OF INTEREST**

Authors declare no conflicts of interest for this article.

**REFERENCES**

1. Longacre TA, Fenoglio-Preiser CM. Mixed hyperplastic adenomatous polyps/serrated adenomas: a distinct form of colorectal neoplasia. *Am. J. Surg. Pathol.* 1990; 14: 524–37.
2. Torlakovic E, Skovland E, Snover DC et al. Morphologic reappraisal of serrated colorectal polyps. *Am. J. Surg. Pathol.* 2003; 27: 65–81.
3. Snover DC, Jass JR, Fenoglio-Preiser C. Serrated polyps of the large intestine. A morphologic and molecular review of an evolving concept. *Am. J. Clin. Pathol.* 2005; 124: 380–91.
4. Snover DC, Ahnen DJ, Burt RW, Odze RD. Serrated polyps of the colon and rectum and serrated (“hyperplastic”) polyps. In: Bozeman FT, Carneiro F, Hruban RH, Theise N (eds). *WHO Classification of Tumours. Pathology and Genetics. Tumours of the Digestive System*, 4th edn. Berlin: Springer, 2010.
5. Leggett B, Whitehall V. Role of serrated pathway in colorectal cancer pathogenesis. *Gastroenterology* 2010; 138: 2088–100.
6. Payne SR, Church TR, Wandell M et al. Endoscopic detection of proximal serrated lesions and pathologic identification of sessile serrated adenomas/polyps vary on the basis of center. *Clin. Gastroenterol. Hepatol.* 2014; 12: 1119–26.
7. Glatz K, Pritt B, Glatz D et al. A multinational, internet-based assessment of observer variability in the diagnosis of serrated colorectal polyps. *Am. J. Clin. Pathol.* 2007; 127: 938–45.
8. Sandmeier D, Seelentag W, Bouzourene H. Serrated polyps of the colorectum: is sessile serrated adenoma distinguishable from hyperplastic polyp in a daily practice? *Virchows Arch.* 2007; 450: 613–8.
9. Farris AB, Misraji J, Srivastava A et al. Sessile serrated adenoma: challenging discrimination from other serrated colonic polyps. *Am. J. Surg. Pathol.* 2008; 32: 30–5.
10. Wong NA, Hunt LP, Novelli MR et al. Observer agreement in the diagnosis of serrated polyps of the large bowel. *Histopathology* 2009; 55: 63–6.
11. Kim SW, Cha JM, Lee Ji et al. A significant number of sessile serrated adenomas might not be accurately diagnosed in daily practice. *Gut, Liv.* 2010; 4: 498–502.
12. Gill P, Wang LM, Bailey A et al. Reporting trends of right-sided hyperplastic and sessile serrated polyps in a large

© 2018 The Authors. *Digestive Endoscopy* published by John Wiley & Sons Australia, Ltd on behalf of Japan Gastroenterological Endoscopy Society.
teaching hospital over a 4-year period (2009–2012). J. Clin. Pathol. 2013; 66: 655–8.

13 Chen Y, Yu J, Liu Y et al. Increasing detection rate of proximal serrated polyps in a large hospital of China over a 10-year period. Int. J. Clin. Exp. Med. 2016; 9: 12745–50.

14 Hetzel J, Huang C, Coukos J et al. Variation in the detection of serrated polyps in an average risk colorectal cancer screening cohort. Am. J. Gastroenterol. 2010; 105: 2656–64.

15 Kahi CJ, Hewett DG, Norton DL et al. Prevalence and variable detection of proximal colon serrated polyps during screening colonoscopy. Clin. Gastroenterol. Hepatol. 2011; 9: 42–6.

16 Anderson JC, Butler LF, Goodrich M et al. Differences in detection rates of adenomas and serrated polyps in screening versus surveillance colonoscopies, based on the New Hampshire Colonoscopy Registry. Clin. Gastroenterol. Hepatol. 2013; 11: 1308–12.

17 Abdeljawad K, Vemulapalli KC, Kahi CJ et al. Sessile serrated polyp prevalence determined by a colonoscopist with a high lesion detection rate and an experienced pathologist. Gastrointest. Endosc. 2015; 81: 517–24.

18 Hasegawa S, Mitsuyama K, Kawano H et al. Endoscopic discrimination of sessile serrated adenomas from other serrated lesions. Oncol. Lett. 2011; 2: 785–9.

19 Kashida H, Ikehara N, Hamatani S et al. Endoscopic characteristics of colorectal serrated lesions. Hepatogastroenterology 2011; 58: 1163–7.

20 Yamada M, Sakamoto T, Otake Y et al. Investigating endoscopic features of sessile serrated adenomas/polyps by using narrow-band imaging with optical magnification. Gastrointest. Endosc. 2015; 82: 108–17.

21 Yamada A, Notohara K, Aoyama I et al. Endoscopic features of sessile serrated adenoma and other serrated colorectal polyps. Hepatogastroenterology 2011; 58: 45–51.

22 Ishigooka S, Nomoto M, Obinata N et al. Evaluation of magnifying colonoscopy in the diagnosis of serrated polyps. World J. Gastroenterol. 2012; 18: 4308–16.

23 Tadepalli US, Feihel D, Miller KM et al. A morphologic analysis of sessile serrated polyps observed during routine colonoscopy (with video). Gastrointest. Endosc. 2011; 74: 1360–8.

24 Bouwens MWE, van Herwaarden YJ, Winkens B et al. Endoscopic characterization of sessile serrated adenomas/polyps with and without dysplasia. Endoscopy 2014; 46: 225–35.

25 Lee EJ, Kim MJ, Chun SM et al. Sessile serrated adenoma/polyps with a depressed surface: A rare form of sessile serrated adenoma/polyp. Diagn. Pathol. 2015; 10: 75.

26 Kawasaki K, Kurahara K, Oshiro Y et al. Clinicopathologic features of inverted serrated lesions of the large bowel. Digestion 2016; 93: 280–7.

27 Hazewinkel Y, López-Cerón M, East JE et al. Endoscopic features of sessile serrated adenomas: validation by international experts using high-resolution white-light endoscopy and narrow-band imaging. Gastrointest. Endosc. 2013; 77: 916–24.

28 Uspeert JEG, Bastiaansen BAJ, van Leerdam ME et al. Development and validation of the WASP classification system for optical diagnosis of adenomas, hyperplastic polyps and sessile serrated adenomas/polyps. Gut 2016; 65: 963–70.

29 Nakao Y, Saito S, Ohy T et al. Endoscopic features of colorectal serrated lesions using image-enhanced endoscopy with pathological analysis. Eur. J. Gastroenterol. Hepatol. 2013; 25: 981–8.

30 Unoka T, Higashi R, Horii J et al. Prospective evaluation of endoscopic criteria characteristic of sessile serrated adenomas/polyps. J. Gastroenterol. 2015; 50: 555–63.

31 Yamashina T, Takeuchi Y, Uedo N et al. Diagnostic features of sessile serrated adenoma/polyps on magnifying narrow band imaging: a prospective study of diagnostic accuracy. J. Gastroenterol. Hepatol. 2015; 30: 117–23.

32 Kimura T, Yamamoto E, Yamano H et al. A novel pit pattern identifies the precursor of colorectal cancer derived from sessile serrated adenoma. Am. J. Gastroenterol. 2012; 107: 460–9.

33 Lambert R, Kudo SE, Vieth M et al. Pragmatic classification of superficial neoplastic colorectal lesions. Gastrointest. Endosc. 2009; 70: 1182–99.

34 Suzuki Y, Kobayashi M, Ishizuka K et al. Inverted hyperplastic polyp diagnosed accurately by magnifying colonoscopy. Gastrointest. Endosc. 2000; 52: 115–18.

35 Kuribayashi K, Ishii T, Ishidate T et al. Two cases of inverted hyperplastic polyps of the colon and association with adenoma. Eur. J. Gastroenterol. Hepatol. 2004; 16: 107–12.

36 Hirasaki S, Kanzaki H, Suzuki S, Shirakawa A. Pedunculated inverted hyperplastic polyp of the sigmoid colon treated with endoscopic polypectomy. Dig. Endosc. 2009; 21: 275–6.

37 Fu K, Fujii T, Kuwayama H et al. Invasive cancer arising in a colonic inverted hyperplastic polyp. Endoscopy 2010; 42(S2): E29–30.

38 Choi HH, Bae SH, Jang EC et al. Intramucosal colon cancer developing in an inverted hyperplastic polyp. Gut. Liver 2013; 7: 126–7.

39 Hu K, Shen S, Zhang L. Herniation of crypts in hyperplastic polyp and sessile serrated adenoma: a prospective study. Am. J. Cancer Res. 2018; 8: 144–53.

40 Rex DK, Clodfelter R, Rahmani F et al. Narrow-band imaging versus white light for the detection of proximal colon serrated lesions: a randomized, controlled trial. Gastroendosc. Endosc. 2016; 83: 166–71.

41 Boparai KS, van den Broek FJ, van Eeden S et al. Hyperplastic polyposis syndrome: a pilot study for the differentiation of polyps by using high-resolution endoscopy, autofluorescence imaging, and narrow-band imaging. Gastrointest. Endosc. 2009; 70: 947–55.

42 Parikh ND, Chaptini L, Njci B, Laine L. Diagnosis of sessile serrated adenomas/polyps with image-enhanced endoscopy: a systematic review and meta-analysis. Endoscopy 2016; 48: 731–9.

43 Tate DJ, Jayanna M, Awadie H et al. A standardized imaging protocol for the endoscopic prediction of dysplasia within sessile serrated polyps. Gastrointest. Endosc. 2018; 87: 222–31.