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**Sessile serrated adenoma/polyps: Where are we at in 2016?**

Singh R *et al.* SSA/P: Where are we at in 2016?

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**Abstract**

It is currently known that colorectal cancers (CRC) arise from 3 different pathways: the adenoma to carcinoma chromosomal instability pathway (50%-70%); the mutator “Lynch syndrome” route (3%-5%); and the serrated pathway (30%-35%). The World Health Organization (WHO) has classified serrated polyps into three types of lesions: Hyperplastic Polyps (HP), Sessile Serrated Adenomas/Polyps (SSA/P) and Traditional Serrated Adenomas (TSA), the latter two strongly associated with development of CRCs. HPs do not cause cancer and TSAs are rare. SSA/P appear to be the responsible precursor lesion for the development of cancers through the serrated pathway. Both HPs and SSA/Ps appear morphologically similar. SSA/P are difficult to detect. The margins are normally inconspicuous. En bloc resection of these polyps can hence be troublesome. A careful examination of borders, submucosal injection of a dye solution (for larger lesions) and resection of a rim of normal tissue around the lesion may ensure total eradication of these lesions.

**Key words:** Colonoscopy; Sessile Serrated adenoma/polyp; Serrated lesion; Colorectal polyps; Colorectal cancer; Polypectomy; Image enhancing endoscopy; Narrow Band Imaging, Endocytoscopy

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**Core tip:** Colorectal cancers (CRC) arise from 3 pathways: adenoma to carcinoma; “Lynch syndrome”; and serrated. There are 3 types of serrated lesions namely: Hyperplastic Polyps, Sessile Serrated Adenomas/Polyps and Traditional Serrated Adenomas, the latter two are associated with CRC. A careful examination of borders, submucosal injection with dye and ensuring that a rim of normal tissue is removed is paramount.

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# INTRODUCTION

Colorectal cancer (CRC) is a major health concern, especially in western countries. According to the American Cancer Society’s estimates, CRC accounts for almost 50000 deaths in the United States with almost 130000 new cases diagnosed in 2016. It is the third commonest type of cancer. Effective screening programs for identification of malignant and premalignant colorectal lesions are thus of utmost importance. In the last few decades the adenoma to adenocarcinoma pathway has been well recognized. For some time it was believed to be the only pathway apart from the “Lynch syndrome” route that results in the development of CRC. The effort to detect and eradicate adenoma have been the main goal in preventive colorectal programs, leading to improved outcomes. Zauber *et al*[1] showed that colonoscopic removal of adenomatous polyps led to a 53% reduction in mortality from CRC during the first 10 years after polypectomy.

It is currently believed that CRC arises from 3 different pathways: the adenoma to carcinoma pathway which accounts for about 50%-70% of cancers; through the mutator ‘Lynch syndrome’ route (3%-5%); and more recently the serrated pathway (30%-35%). The latter have become increasingly recognized as a separate route which could lead to the development of CRC[2].

This triplet division is based on the combined clinical-molecular characteristics of the lesions. A deeper understanding of the molecular pathways in CRC have been described by Jass in 2007[3] and updated by Phipps *et al*[4] in 2015. They described 5 molecular subtypes and associated genetic distortions to describe each one. Subtypes 1, 2 and 3 are related to the serrated pathway. Subtypes 1 and 2 are either MicroSatellite Instable (MSI)-high or MicroSatellite Stable (MSS)/MSI-low cancers which have the CpG Island Methylator Phenotype (CIMP) and *BRAF*-mutation but are *KRAS* negative. The third subtype represents an alternative pathway which originates in *KRAS* mutation with no CIMP, *BRAF* or MSI association. Subtypes 2 and 3 have a higher association with mortality[4]. Subtype 4 reflects CRC arising from the traditional adenoma-carcinoma sequence, and are MSS/MSI-low, CIMP, *BRAF* and *KRAS* negative. Subtype 5 indicates Lynch Syndrome and is associated with high prevalence of a family history of CRC. They are MSI-high but CIMP, *BRAF* and *KRAS* negative.

The serrated pathway is much less well understood. Systematic resection of premalignant serrated lesions could further improve the outcomes of CRC screening programs. One of the main problems with this protocol is the difficulty in identifying these lesions. Unlike adenomas, not all serrated lesions are linked to colorectal cancer. According to the World Health Organization (WHO), there are three types of serrated lesions: Hyperplastic Polyps (HP), Sessile Serrated Adenomas/Polyps (SSA/P) and Traditional Serrated Adenomas (TSA). TSA is usually easy to identify due to its protuberant pine cone-shape. While SSA/P is also associated with cancer, HP is not and their discrimination is troublesome as they look morphologically similar at colonoscopy, even with Image Enhancing Endoscopy (IEE) techniques. Despite the adoption of numerous different classifications, the ability to predict HP from SSA/P has unfortunately been overlooked[5,6]. More recently, a newly proposed approach known as Workgroup Serrated polypS and Polyposis WASP classification has allowed the distinction between HP and SSA/P with reasonable accuracy[7]. It consists of cloud-like surface, indistinctive borders, irregular margins and open pit patterns, features described as being associated with SSA/P in another previous study[8]. The need to adequately identify SSA/P from HP arises from evidence supporting SSA/P as the major malignant source amongst serrated lesions[2,9,10,11,12].

**IMAGE ENHANCED ENDOSCOPY**

Detecting and characterizing colorectal lesions by image enhanced endoscopy (IEE) has been reported in several articles[13-17] and has been found to have 92.7% sensitivity and 87.3% specificity in differentiating adenomas/adenocarcinomas from “non-neoplastic” lesions[18]. Differentiating serrated lesions, specifically SSA/P from HP is more challenging. The incidence of serrated lesions in the overall population is 5%-8% (contrasting with 30%-40% for adenomas), and they are more difficult to see due to their colour and shape[8,19,20]. Their rarity and discreet morphology could be why there is a longer learning curve compared to that for adenomas[21-24].

The evaluation of dysplasia within the SSA/Ps could also be of value. It has been described by Chino *et al*[25] 2016 that the evaluation of crypts and submucosal vessels with NBI (Narrow Band Imaging) and magnification might be useful in evaluating dysplasia in SSA/P, which leads to poorer outcomes.

Although there is certainly enthusiasm for IEE techniques, histopathology remains the gold standard for evaluating colorectal lesions. Nonetheless, improving technology that could be used by the endoscopist in real time would definitely be beneficial for serrated lesions as it has been for adenomas[26]. This technology will need to provide immediate feedback and accurately predict the final histopathology (Figure1).

**SSA/P AND HP DIFFERENTIATION**

A conceptual way to define each serrated lesion is based on differences in the proliferation zones within the serrated crypts in each group[27]. In HP, the expanded proliferation zone is located at the base of the crypts and cells mature towards the surface symmetrically. In SSA/P, the proliferation zone is to the side of the crypts instead of the base, resulting in maturation of epithelial cells laterally, towards the surface and the base, leading to crypt base dilatation (pattern II-open). Within SSA/P, the presence of dysplasia is usually evident and must be accompanied by SSA/P component adjacent to it once its histopathology is similar to adenomas. Unfortunately, this theoretical classification may be misleading. Confounded even by expert pathologists, the poor agreement for the diagnosis of villous features or high grade dysplasia has a 10-fold variability[28-30].

New techniques for real-time *in vivo* optical diagnosis using IEE have been developed to potentially predict histology and perhaps permit a more practical and economical approach for low-risk polyps; for example the “resect and discard” approach[31-34]. There is evidence from several original articles and meta-analyses that *in vivo* optical diagnosis using either Narrow-Band Imaging (NBI) or Fujinon Intelligent ChromoEndoscopy (FICE) would be more cost-effective compared to histology without significant changes in follow-up decision, especially for diminutive polyps[34-37]. The American Society for Gastrointestinal Endoscopy statement of 2011 (Preservation and Incorporation of Valuable endoscopic Innovations - PIVI) describes the standards that new technologies have to achieve in order to be implemented. For the ‘resect and discard’ strategy, it asks for ≥ 90% agreement in the assignment of post-polypectomy surveillance intervals when compared with decisions based on histopathology. With regards to the policy of leaving suspected rectosigmoid hyperplastic polyps measuring ≤ 5mm in place, a ≥ 90% negative predictive value for adenomatous histology is mandated[31]. Abu Dayyeh *et al*[38] on behalf of the ASGE Technology Committee in 2015 reported in a meta-analysis that the diagnostic value of IEE for diminutive colorectal polyps achieved a pooled NPV 91% and pooled follow-up agreement of 89%. Despite the pooled analysis for agreement in the assignment of surveillance intervals which did not reach the 90% threshold for NBI; experienced endoscopists were able to exceed this (93%) when the diagnosis was made with high confidence.

**PREDICTORS OF MALIGNANCY AMONG SSA/P**

The most common group of lesions are the diminutive polyps (≤ 5mm in size), which represent approximately 60% of all polyps detected at primary screening colonoscopy. Their overall association with advanced pathology is low but not negligible[39,40]. On the contrary, Burgess *et al*[41] have demonstrated that size matters in terms of SSA/P. For every 10mm increase in lesion size, the Odds Ratios (OR) is 1.90 for cytological dysplasia. SSA/P with cytological dysplasia (SSA/P-D) is also associated with presence of 0-Is component of the Paris’ Classification (OR = 3.1), Kudo’s pit pattern III, IV or V (OR = 3.98) and increasing age (OR = 1.69 per decade).

Yamada *et al*[32] recently described the presence of dilated branch vessels (DBV) as an aspect of SSA/Ps with dysplasia. Apart from their characteristics at chromoendoscopy and magnification[42,43], there are some aspects that we can use to distinguish SSA/Ps with and without malignancy potential.

Endocytoscopy is an emerging modality with diagnostic potential for SSA/P. It allows *in vivo* visualization of cells and nuclei facilitating precise real-time pathological prediction. Oval gland lumens with small round nuclei has a sensitivity of 83.3% and specificity of 97.8% for the diagnosis of SSA/P. It is also a promising tool for diagnosing SSA/P-D due to its ability to detect morphological changes in the nuclei as described by Kutsukawa *et al*[44] and Mori *et al*[45].

**OPTIMAL RESECTION OF A SSA/P**

Numerous studies have grim numbers in regards to SSA/P complete resection rates[46-48]. Against these odds, a more recent study from our group[49] studied the resection of 2000 lateral spreading tumors (LST) and attributed the high recurrence to the inconspicuous margins of the SSA/P, which was overcome with IEE techniques. Submucosal instillation of a dye based solution (for larger lesions), a careful examination of borders and a rim of normal tissue resected together with the lesion may have affected the high rate of complete removal of the SSA/P. It is evident the contribution that advanced endoscopy apparel and endoscopist’s expertise is essential[50] in order to keep the recurrence of resection as low as 7%, as described by Pellise *et al*[49] (Figure2).

**FOLLOW-UP**

The current guidelines from the American and European Societies for Gastrointestinal Endoscopy (ASGE and ESGE) advocates the standard 5-10 years surveillance period for low risk lesions (SSA/*P* < 10 mm and without dysplasia), in patients without serrated polyposis syndrome. Patients with larger SSA/Ps or with dysplasia should have their colonoscopy repeated in 3 years-time[51,52]. The serrated polyposis syndrome is defined if any number of serrated polyps occurring proximal to the sigmoid colon in an individual who has a first-degree relative with serrated polyposis; if at least five serrated polyps are found proximal to the sigmoid colon (at least 2 with ≥ 10 mm); and if more than 20 serrated polyps of any size distributed throughout the colon. In these cases, the follow-up should be at 1 year[51]. The major problem is that these guidelines rely upon the assumption that the serrated lesions are detected and resected adequately, which is not always the case.

**CONCLUSION**

SSA/P is an important pre-malignant lesion that can easily be missed. Efforts must be made in order to alter the nomenclature of “non-neoplastic lesions” to non-adenomatous lesions as the role of serrated lesions in the development of colorectal cancer is now well established. A longer training must be pursued and cutting-edge IEE technologies developed and studied in order to diminish the miss rate for serrated lesions. The implementation of a “serrated polyps detection rate” could be implemented alongside the “adenoma detection rate” as a quality indicator for colonoscopy.

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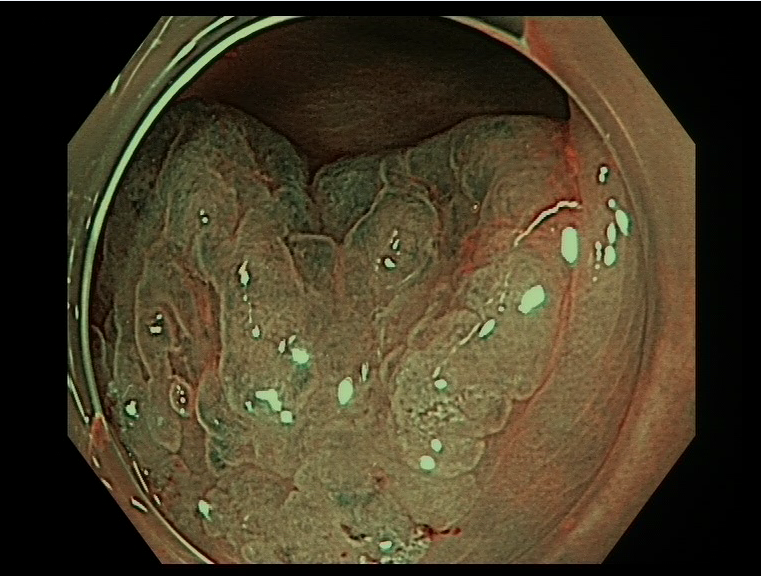
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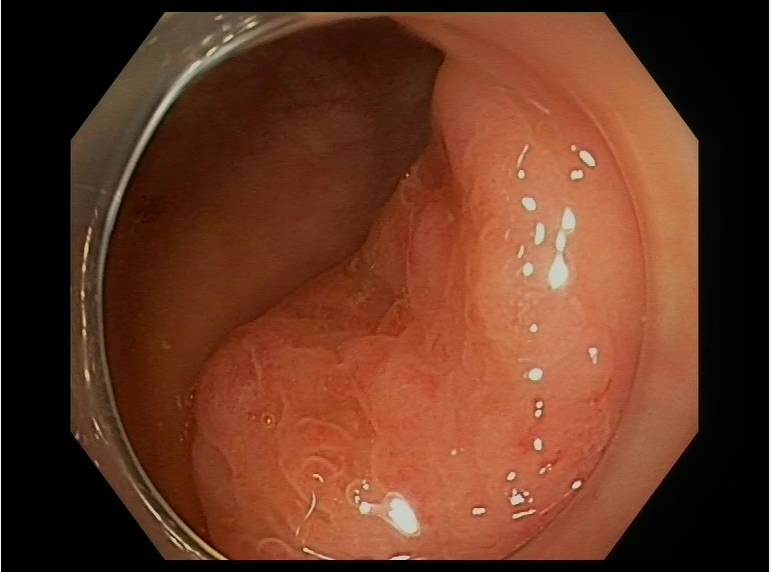
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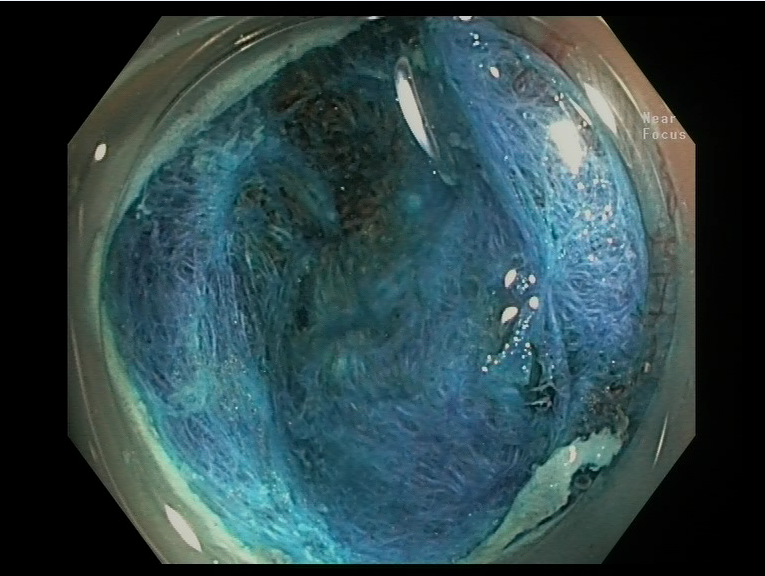
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**Figure 1 Inconspicuous margins of a sessile serrated adenomas/polyps with and without narrow-band imaging.**

 **Figure 2 Resection of a sessile serrated adenomas/polyps with dye of submucosal layer with indigo carmine – no residual lesion.**