Interendoscopist variability in proximal colon polyp detection is twice higher for serrated polyps than adenomas

Jean-François Bretagne, Stéphanie Hamonic, Christine Piette, Jean-François Viel, Guillaume Bouguen

AIM
To assess the interendoscopist variability in the detection of colorectal polyps according to their location and histological type.

METHODS
This study was a retrospective analysis of prospectively collected data from a regional colorectal cancer (CRC) screening program; 2979 complete colonoscopies from 18 endoscopists were included. Variability in performance between endoscopists for detection of at least one adenoma (A), one proximal adenoma (PA), one distal adenoma (DA), and one proximal serrated polyp (PSP) was assessed by using multilevel logistic regression models.

RESULTS
The observed detection rates among the 18 endoscopists ranged from 24.6% to 47.6% (mean = 35.7%) for A, from 19.1% to 39.0% (mean = 29.4%) for DA, from 6.0% to 22.9% (mean = 12.4%) for PA, and from 1.3% to 19.3% (mean = 6.9%) for PSP.

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After adjusting for patient-level variables (sex, age), the interendoscopist detection rates variability achieved a significant level for A, PA, and PSP but not for DA ($P = 0.03$, $P = 0.02$, $P = 0.02$ and $P = 0.08$, respectively). This heterogeneity, as measured by the variance partition coefficient, was approximately threefold higher for PA (6.6%) compared with A (2.1%), and twofold higher for PSP (12.3%) compared with PA.

**CONCLUSION**

These results demonstrate significant interendoscopist variability for proximal polyp particularly for serrated polyps, but not for distal adenoma detection. These findings contribute to explain the decreased effectiveness of complete colonoscopies at preventing proximal CRCs and the need to carefully assess the proximal colon during scope procedure.

**Key words:** Colonoscopy; Colorectal cancer; Adenoma; Serrated polyp; Proximal polyp; Detection rate; Quality performance

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**Core tip:** The present study demonstrates high interendoscopist variability in adenoma, proximal adenoma, and proximal serrated polyp detection rates but not in distal adenoma detection rates. The magnitude of interendoscopist variation was wider for proximal serrated polyps as compared to proximal adenoma detection. Altogether, these findings might explain why complete colonoscopies are less effective at preventing proximal than distal colorectal cancers.

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**INTRODUCTION**

Adenoma detection and removal is the basis for the reduction in colorectal cancer (CRC) incidence and mortality achieved by colonoscopy [1-3]. However, recent studies raised concerns that screening colonoscopies may not decrease CRC incidence and mortality in the proximal colon to the same extent as in the distal colon [4-8]. Although there are multiple plausible explanations for the decreased effectiveness in the proximal colon, the quality of the colonoscopy is a critical issue. Recent studies demonstrated that surrogate indicators for colonoscopy quality performance, such as adenoma detection rates and cecal intubation rates, were predictors of interval CRCs that occur after screening colonoscopies [9-11]. A higher miss rate of proximal adenomas compared to distal adenomas could explain the decreased protective effect of colonoscopy for proximal colon cancer. However, no data are available on the interendoscopist variability of adenoma detection according to the polyp location in the colon, particularly in population-based studies.

Serrated polyps might be another significant contributor to the decreased protective effect of colonoscopies for proximal colon cancer. Serrated lesions can be challenging to visualize because of their morphologic characteristics and could be more likely overlooked as compared to conventional adenomas. Cohort studies demonstrated a wide variation rate among endoscopists of the proximal serrated polyps detection rates [12-14], but no study aimed to compare adenomas detection and serrated polyps detection variability amongst endoscopists, especially for proximal colon location.

This population-based study aimed to test the hypothesis that the variations in adenoma detection rates between colonoscopists are wider for the proximal colon compared with the distal colon, and to compare interendoscopist variability in polyp detection rates in the proximal colon between serrated polyps and adenomas.

**MATERIALS AND METHODS**

**Study population**

The study was conducted in "Ille et Vilaine", which has a population of approximately one million and was one of the first French districts to implement a national screening program at the end of 2002. The mass screening is based on biennial guaiac fecal occult blood tests. The target population for the screening includes asymptomatic men and women between 50 and 74 years of age with no other CRC risk factors. Individuals with a family history of CRC or a personal history of CRC or adenomas, those with inflammatory bowel disease, and those who had undergone a total colonoscopy in the previous five years were excluded from the screening program.

There were 96054 (51.8%) and 89309 (46.7%) participants in the first and second rounds, respectively. The proportion of positive tests amongst the participants was 2.58% and 2.26%, respectively. Positive testing was followed by a colonoscopy in 92.6% and 91.4% of the subjects, respectively. Finally, 2295 and 1848 colonoscopies were performed from 2003 to 2007 in the first and second rounds, respectively.

The 18 endoscopists who had performed at least 30 colonoscopies following a positive test in each of the first two rounds of the screening program were included. Fourteen of the 18 endoscopists were in private practice, and 4 worked in public hospitals. Overall, the 18 endoscopists performed 3487 (84.2%) of the 4143 white-light colonoscopies of the 2
screening rounds. Although high cecal intubation rates were recorded for rounds 1 and 2 (96.3% and 95.9%, respectively), we included only complete examinations of the colon in this study. The data from both rounds were pooled because no difference in the colonoscopy findings was noted between the two rounds. We previously reported that individual endoscopists who had participated in the CRC screening program as a factor was not a significant predictor of CRC detection but was a significant predictor of adenoma detection\[15\].

In the present study, a secondary analysis of the colonoscopy data was done to explore variations in the detection rate of at least one adenoma according to location in the colon and to compare interendoscopist variability in polyp detection rates in the proximal colon according to histological subtype (serrated polyps and adenomas). The CRC screening program was declared and approved by the CNIL “Commission Nationale de l’Informatique et des Libertés” on August 30th 2002 (n° 812571). Research was approved by the CCTIRS “Comité Consultatif pour le Traitement de l’Information en matière de Recherche dans le domaine de la santé”.

**Study design and outcomes**

This was a cross-sectional study that used data retrieved from a prospectively collected database. Three adenoma detection rates, which were expressed as the proportion of complete colonoscopies with at least one adenoma, were calculated for each endoscopist as follows: The distal adenoma detection rate (DA.DR) for at least one adenoma detected in the distal colon (i.e., below the splenic flexure including the flexure), the proximal adenoma detection rate (PA.DR) for at least one adenoma in the proximal colon (i.e., proximal to the splenic flexure) and the A.DR for at least one colorectal adenoma regardless of its location in the colon. Colonoscopies with CRC, including carcinoma, were not included in the analysis because additional polyps in these patients were not recorded in the database. The individual detection rates for serrated polyps in the proximal colon (PSP.DR) were also calculated for each endoscopist. Serrated polyps were defined as an entire group of polyps that included traditional hyperplastic polyps, sessile serrated adenomas/polyps and traditional serrated adenomas.

The observed and adjusted (i.e., according to patient age and gender) adenoma detection rates were calculated for each endoscopist at each site. Similarly, the observed and adjusted (i.e., according to patient age and gender) proximal serrated polyp detection rates were calculated for each endoscopist. The variability between endoscopists in the probability to detect one adenoma/one adenoma in the distal colon/one adenoma in the proximal colon/one serrated polyp in the proximal colon was assessed by multilevel logistic regression models.

Additional analyses were performed after defining a proximal polyp as proximal to the hepatic flexure instead of proximal to the splenic flexure. Furthermore, we assessed the interendoscopist variability for polyps of size $\geq 10$ mm.

**Statistical analysis**

Continuous variables were expressed as the mean, standard error, median and interquartile range, and extremes values; categorical variables were expressed as numbers and percentages. The observed detection rates were compared between males and females and then between age classes using the Wilcoxon test; the use of the Cochran-Mantel Haenszel test permitted endoscopists to be adjustment variables. The patient age- and gender-adjusted adenoma detection rates for each endoscopist (and the corresponding 95%CI) were defined as the observed proportion of colonoscopies with at least one adenoma detected amongst all subjects multiplied by the ratio of the observed to the predicted number of detected adenomas for one endoscopist. The predicted number of detected adenomas for each endoscopist was assessed using logistic regression. Multilevel logistic regression models were used given the hierarchical structure of the sample (i.e., patients are aggregated at the endoscopist level) and binary outcomes (i.e., at least one adenoma/at least one distal adenoma/at least one proximal adenoma/at least one proximal serrated polyp). Each model was a two-level model in which age and gender were included as fixed effects (first or patient level) and in which the endoscopist was introduced as a random effect (second or endoscopist level). The fixed effect results are presented as odds ratios with 95%CI. To determine the proportion of total variance of the outcome that is explained at the endoscopist level, the variance partition coefficient was calculated using the Snijders and Bosker approximation\[16,17\].

$$VPC = \frac{\sigma^2_u}{\sigma^2_u + \pi^2/3}$$

where $\sigma^2_u$ is the variance of the endoscopist-level random effect representing the between-endoscopist variability in terms of the outcome. The correlations between adjusted values of polyp detection rates were tested using the Spearman rank test. For all tests, the significance threshold was $\alpha = 5\%$. The analyses were performed using SAS version 9.2 (SAS Institute Inc., Cary, NC, United States).

**RESULTS**

**Distribution of the adenomas and serrated polyps within the population**

After excluding incomplete examinations ($n = 210$) and colonoscopies harboring CRC ($n = 298$), 2979 colonoscopies were included for the analysis. Of these, 1531 (51.4%) were performed in males and
higher detection rate in males compared with females (7.0% vs 4.3%, respectively, \( P = 0.06 \)).

Factors associated to the adenoma and serrated polyp detection rates from multilevel logistic regression

The results of the multilevel logistic regressions are presented in Table 4. Age and gender were significant factors for polyp detection regardless of the indicator used. After adjusting for patient-level variables, the interendoscopist variability achieved a significant level for A.DR, PA.DR, and PSP.DR but not for DA.DR (\( P = 0.03 \), \( P = 0.02 \), \( P = 0.02 \) and \( P = 0.08 \), respectively). The corresponding variance partition coefficients were as follows: 2.1%, 6.6%, 12.3%, and 1.3%. The heterogeneity between endoscopists was approximately threefold higher for PA.DR compared with A.DR, and twofold higher for PSP.DR compared with PA.DR.

Complementary analyses

The abovementioned results were not affected when the proximal colon was defined as proximal to the hepatic flexure (data not presented).

Table 1 shows the observed detection rates per endoscopist for each of the following four indicators: Adenoma detection rate (A.DR), DA.DR, PA.DR, and proximal serrated polyp detection rate (PSP.DR). The mean detection rates were 35.7%, 29.4%, 12.4% and 6.9%, respectively.

The mean and median values of these four indicators according to gender and age are provided in Tables 2 and 3. For each of the measures related to adenoma detection, the median values were significantly higher in males compared with females, and the values increased with increasing age. The median PSP.DR values did not differ significantly according to age group, but there was a trend for a higher detection rate in males compared with females (7.0% vs 4.3%, respectively, \( P = 0.06 \)).

Individual endoscopists’ neoplasia detection rates

Amongst the 18 colonoscopists, the median gender- and age-adjusted values for the detection of polyps \( \geq 10 \) mm were 17.5%, 16.2%, 2.6% and 0.6% for A.DR, DA.DR, PA.DR and PSP.DR, respectively, without significant statistical interendoscopist variability. The interendoscopist variability amongst the 18 colonoscopists remains significant for the detection rate of proximal polyps of any histological subtype (i.e., proximal serrated polyp and/or proximal polyp detection rate).

### Table 1  Observed polyp detection rates amongst the 18 colonoscopists

| Endoscopist (No.) | Colonoscopies (n) | A.DR | DA.DR | PA.DR | PSP.DR |
|-------------------|------------------|------|-------|-------|--------|
| A                 | 273              | 33.33% | 29.67% | 7.33% | 4.40%  |
| B                 | 272              | 24.63% | 19.49% | 6.25% | 5.51%  |
| C                 | 213              | 46.48% | 38.97% | 17.84% | 19.25% |
| D                 | 210              | 47.62% | 38.10% | 18.57% | 6.19%  |
| E                 | 207              | 28.99% | 25.12% | 7.25% | 5.31%  |
| F                 | 185              | 30.81% | 27.57% | 5.95% | 2.70%  |
| G                 | 172              | 32.56% | 29.07% | 9.88% | 7.56%  |
| H                 | 164              | 31.71% | 28.05% | 8.54% | 8.54%  |
| I                 | 160              | 35.63% | 30.00% | 11.25% | 3.13%  |
| J                 | 157              | 47.13% | 35.67% | 22.93% | 15.92% |
| K                 | 148              | 33.11% | 29.05% | 11.49% | 2.03%  |
| L                 | 148              | 36.49% | 31.76% | 11.49% | 10.14% |
| M                 | 135              | 37.04% | 25.19% | 17.78% | 4.44%  |
| N                 | 135              | 40.74% | 31.11% | 14.07% | 5.93%  |
| O                 | 132              | 34.85% | 33.33% | 6.82% | 2.27%  |
| P                 | 105              | 30.48% | 19.05% | 21.90% | 15.24% |
| Q                 | 85               | 38.82% | 31.76% | 12.94% | 4.71%  |
| R                 | 78               | 32.05% | 25.64% | 11.54% | 1.28%  |
| N mean            | 165.50          | 35.69% | 29.37% | 12.43% | 6.92%  |
| SD                | 54.65           | 6.41% | 5.41% | 5.36% | 5.14%  |
| Median            | 158.50          | 34.09% | 29.37% | 11.49% | 5.41%  |
| Q1-Q3             | 135-207         | 31.71%-38.82% | 25.64%-31.76% | 7.33%-17.78% | 3.13%-8.54% |
| Min-Max           | 78-273          | 24.63%-47.62% | 19.05%-38.97% | 5.95%-22.93% | 1.28%-19.25% |

A.DR: Adenoma detection rate; DA.DR: Distal adenoma detection rate; PA.DR: Proximal adenoma detection rate; PSP.DR: Proximal serrated polyp detection rate.
adenoma) (data not presented). The corresponding variance partition coefficient was 9.6%, which was an intermediary value between the PSP.DR and PA.DR values.

Correlation studies between adjusted values of polyp detection rates

PSP.DR values were not correlated with A.DR values ($\rho = 0.19, P = 0.45$), but were significantly correlated with PA.DR values ($\rho = 0.55, P < 0.002$). PA.DR values were highly significantly correlated with A.DR values ($\rho = 0.83, P < 0.0001$).

**DISCUSSION**

Colonoscopies are known to display great variability in A.DRs between endoscopists in various settings, including in academic[18], mixed community-academic[19], community practices[20,21] and population-based studies[15,22]. However, no study focused on the variability in A.DRs according to the proximal or distal location of the adenomas in the colon. Although

| Table 2 | Comparison of polyp detection rates (%) between males and females amongst the 18 colonoscopists using the Wilcoxon test |
|---------|--------------------------------------------------|
| A.DR    | Males | Females | Total | $P$ value |
| n       | 18    | 18      | 18    |           |
| mean    | 46.3  | 24.5    | 35.7  |           |
| Median  | 44.9  | 23.7    | 34.1  | < 0.0001  |
| Q1-Q3   | 40.7-50.0 | 20.0-27.7 | 31.7-38.8 | |
| Min-Max | 35.8-67.3 | 12.5-39.3 | 24.6-47.6 | |
| DA.DR   | n     | 18      | 18    |           |
| mean    | 38.2  | 20      | 29.4  | < 0.0001  |
| Median  | 37.7  | 20.6    | 29.4  |           |
| Q1-Q3   | 32.4-40.0 | 17.9-22.7 | 25.6-31.8 | |
| Min-Max | 27.9-54.8 | 11.0-29.2 | 19.0-39.0 | |
| PA.DR   | n     | 18      | 18    |           |
| mean    | 17.8  | 7.1     | 12.4  |           |
| Median  | 15.3  | 6.6     | 11.5  |           |
| Q1-Q3   | 11.0-22.4 | 3.6-10.8 | 7.3-17.8 | |
| Min-Max | 8.6-40.0 | 13.0-18.0 | 15.2-22.9 | |
| PSP.DR  | n     | 18      | 18    |           |
| mean    | 8.3   | 5.3     | 6.9   |           |
| Median  | 7.0   | 4.3     | 5.4   | 0.06      |
| Q1-Q3   | 4.5-10.8 | 1.5-6.6 | 3.1-8.5 | |
| Min-Max | 2.2-25.6 | 0.0-16.9 | 1.3-19.2 | |

A.DR: Adenoma detection rate; DA.DR: Distal adenoma detection rate; PA.DR: Proximal adenoma detection rate; PSP.DR: Proximal serrated polyp detection rate.

| Table 3 | Comparison of polyp detection rates (%) according to age groups amongst the 18 colonoscopists using the Wilcoxon test |
|---------|--------------------------------------------------|
| < 55    | 55-59 | 60-64 | 65-69 | $\geq$ 70 | Total | $P$ value |
| A.DR    | n     | 18    | 18    | 18    | 18    | 18    |           |
| mean    | 29.5  | 28.6  | 36.7  | 42.3  | 42.2  | 35.7  |           |
| Median  | 28.5  | 29.6  | 38.8  | 43.5  | 43.0  | 34.1  | < 0.0001  |
| Q1-Q3   | 24.4-34.8 | 21.4-33.3 | 29.4-41.4 | 28.6-50.0 | 37.5-45.5 | 31.7-38.8 | |
| Min-Max | 16.7-42.9 | 12.5-46.2 | 18.4-52.9 | 21.7-73.0 | 29.0-53.8 | 24.6-47.6 | |
| DA.DR   | n     | 18    | 18    | 18    | 18    | 18    |           |
| mean    | 23.9  | 23.6  | 31.9  | 35.3  | 33.4  | 29.4  |           |
| Median  | 22.3  | 23.8  | 33.0  | 39.0  | 33.3  | 29.4  | 0.0003    |
| Q1-Q3   | 17.6-31.8 | 18.5-27.8 | 29.4-37.5 | 22.7-44.4 | 26.5-37.5 | 25.6-31.8 | |
| Min-Max | 11.1-35.3 | 12.5-36.8 | 14.3-42.9 | 13.6-54.1 | 23.1-51.6 | 19.0-39.0 | |
| PA.DR   | n     | 18    | 18    | 18    | 18    | 18    |           |
| mean    | 9.1   | 8.8   | 11.7  | 15.9  | 17.4  | 12.4  |           |
| Median  | 8.3   | 9.5   | 9.1   | 17.5  | 15.9  | 11.5  | 0.0006    |
| Q1-Q3   | 5.9-9.3 | 4.3-11.8 | 5.0-18.2 | 11.9-20.0 | 12.5-20.6 | 7.3-17.8 | |
| Min-Max | 0.250  | 0.179  | 0.310  | 2.5-32.4 | 0.462  | 5.9-22.9 | |
| PSP.DR  | n     | 18    | 18    | 18    | 18    | 18    |           |
| mean    | 6.1   | 6.5   | 6.2   | 8.1   | 8.6   | 6.9   |           |
| Median  | 5.0   | 4.7   | 1.8   | 5.8   | 8.0   | 5.4   | 0.37      |
| Q1-Q3   | 0.7-7.4 | 3.2-10.0 | 0.0-10.0 | 2.5-10.2 | 3.2-12.9 | 3.1-8.5 | |
| Min-Max | 0.0-20.9 | 0.0-21.4 | 0.0-23.1 | 0.0-26.5 | 0.0-20.6 | 1.3-19.2 | |

A.DR: Adenoma detection rate; DA.DR: Distal adenoma detection rate; PA.DR: Proximal adenoma detection rate; PSP.DR: Proximal serrated polyp detection rate.
the present study confirms significant variability for adenoma detection amongst colonoscopists, these data indicate that interendoscopist variability achieves a significant level for proximal adenomas but not distal adenomas detection. Serrated polyps were included to demonstrate that interendoscopist variability was also significant for proximal serrated polyp detection, even higher. These findings which resulted from in-depth statistical analyses using multilevel logistic regressions, demonstrate a higher heterogeneity for proximal serrated polyp than proximal adenoma detection amongst endoscopists. Detection rates for distal serrated polyps were not assessed because we hypothesised that variability between colonoscopists could be related to other factors than the quality of performance by colonoscopists. Some endoscopists might intentionally avoid performing a biopsy or discard small rectal polyps that have the appearance of hyperplastic polyps in the rectosigmoid.

All of these findings contribute to underline that the proximal colon is a difficult issue for colonoscopists. Otherwise, the performances of colonoscopists are more variable for proximal adenomas compared with distal adenomas and within the proximal colon for serrated polyps compared with adenomas. A correlation between adenomas and serrated polyps detection rates is debatable in the literature. No significant correlation between both rates was found similarly to some studies[23,24]. On the opposite, other studies reported a significant correlation between adenoma detection rate and detection rate of polyps with other histological type (sessile serrated polyps, serrated polyps and proximal serrated polyps)[12,14,25-29], but all underlined the poor correlation between A.DR and PSP.DR. Moreover, the significant correlation we found between both proximal polyps detection rates is in accordance with results from one large cohort study[23].

The mean detection rate for proximal adenomas (12.4%) was significantly lower than the 38% rate reported by Kah et al.[20] in a recent series of 6681 screening colonoscopies. The anatomical distribution of adenomas in the large bowel is debatable. A rightsided dominance of adenomas has been reported in some studies for both sexes[12,31], or for women only[31]. Of note, we did not observe such distribution for adenomas. But, our findings are in line with data of colonoscopies following a positive FOBT in France[32]. Interestingly, data of 2821392 nationwide screening colonoscopies in Germany indicated that only 28.7% of adenomas detected were proximal to the sigmoid colon[33].

The ranges of proximal serrated polyp detection rates amongst endoscopists were 1.3%-19.3% in the present study. Two other studies by Kahi et al.[13] (1%-18%) and Ijspeert et al.[30] (2.9%-18.6%) reported similar rates but one study of 7215 screening colonoscopies including 32 endoscopy centers observed lower detection rate of proximal serrated polyps (mean 2.8%, range 0-9.8%)[14]. This discrepancy may be secondary to the selected population, bowel preparation quality, endoscopists’ technique or skill. While the rate of clinically relevant serrated polyps was recently reported to be similar in FOBT-based screening cohorts and in primary colonoscopy screening cohorts[12], the strengths of the current cohort remains its homogeneity due to the population selection by a single indication. With regard to the population, our study underlines the fact that the prevalence of proximal serrated polyps does not differ according to age and gender[13,14,24]. The trend for a higher detection rate in males compared with females that we found in the present study is in accordance with recent findings from post-FOBT colonoscopies[31].

These results point out the substantial numbers of undetected lesions in the proximal colon in clinical practice. The wider variability observed for serrated polyps compared to adenomas amongst endosco-
pists support a more subtle appearance of serrated polyps,[25,36] particularly of small and diminutive polyps because we did not find any significant variation in the detection rates for proximal polyps of a size greater than 10 mm for either adenomas or serrated polyps. We speculate that education and training are important not solely for adequate mucosal exposure, but also for identification of subtle lesions such as serrated polyps.

Individual endoscopists’ adenoma detection rates have been demonstrated to be associated with interval cancer risk[9,11]. The hypothesis that the proximal serrated polyp detection rate or a composite measure for proximal polyp (i.e., adenoma and/or serrated polyp) detection could predict interval CRCs even more accurately than the adenoma detection rate remains to be established.

The present study has several limitations. The preparation quality was not considered in our study. In a recent prospective study, sessile serrated polyps were detected in a significantly smaller proportion of patients with intermediate-quality preparation than high-quality preparation for the whole colon and the right colon[37]. However, it seems unlikely that it could explain the wider magnitude of variation for proximal serrated polyp detection compared with proximal adenoma detection. The withdrawal time was not considered in our study. It has been recently reported that it could affect serrated polyp detection[23,24,38]. Thus, we cannot exclude that a given withdrawal time could affect differently serrated polyp and adenoma detection. Furthermore, patient-related factors known to modify adenoma prevalence, such as smoking, obesity, or aspirin use, were not considered. By contrast, one strength of our study was the exclusion of subjects with a family history of CRC because of their particular distribution of adenomas in the colon[39]. Overall, it seems unlikely that patient-related factors could have explained the magnitude of the variability observed in our study. Nevertheless, as suggested by the variance partition coefficient values found in the current study, other factors than endoscopists could explain interendoscopist variability. Another limitation might be related to the absence of distinction of subtypes of serrated polyps. However, pooling the different histopathological types of serrated polyps is justified when considering the considerable interobserver variation in the differentiation of serrated polyps[12,40,41] and the significant correlation between both detection rates of proximal serrated polyps and clinically relevant serrated polyps[29]. Lastly, we have no information regarding the endoscopes used by the endoscopists during the study period. We hypothesize that all colonoscopies were performed with standard definition endoscopes. Thus, it remains to demonstrate that high definition endoscopes could reduce the variability for serrated polyp detection amongst endoscopists and the gap between proximal serrated polyp and proximal adenoma detection amongst endoscopists.

In conclusion, our findings demonstrate significant variability in the detection of proximal polyps, which included both adenomas and serrated polyps, amongst endoscopists. The heterogeneity was approximately twofold higher for proximal serrated polyps than for proximal adenomas detection. These findings might explain why complete colonoscopies are less effective at preventing proximal CRCs than distal ones. Furthermore, our results question the potential of using a proximal serrated polyp detection rate as a surrogate indicator for interval CRC risk.

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COMMENTS

Background

The variability of detection rate for proximal serrated polyps amongst endoscopists has been reported in the literature. However, no study aimed to compare adenomas and serrated polyps detection variability amongst endoscopists, especially for proximal colon location.

Research frontiers

The authors indicate that interendoscopist variability achieves a significant level for proximal adenomas but not for distal adenomas detection. Moreover, the heterogeneity was approximately twofold higher for proximal serrated polyps than for proximal adenomas detection.

Innovations and breakthroughs

The interendoscopist variability in detection of proximal polyps (serrated polyps and adenomas) had never been compared in previous studies.

Applications

The authors findings might explain why complete colonoscopies are less effective at preventing proximal colorectal cancer (CRCs) than distal ones. Furthermore, our results question the potential of using a proximal serrated polyp detection rate as a surrogate indicator for interval CRC risk.

Terminology

Proximal polyps are defined as polyps located above the splenic flexure. Serrated polyps were defined as an entire group of polyps that included traditional hyperplastic polyps, sessile serrated adenomas/polyps and traditional serrated adenomas.

Peer-review

The study was appropriately designed and analysed. Its result has thoroughly strong impact on routine practice, especially on screening colonoscopy program. The manuscript is clearly constructed and written in appropriate English. There is no major issues to be revised in their paper.

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