Taller Haustral Folds in the Proximal Colon: A Potential Factor Contributing to Interval Colorectal Cancer

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ABSTRACT
Missed cancers have been reported at higher frequencies in the right colon despite optical colonoscopy screening. The purpose of this study was to determine if there are regional differences in haustral fold height between the ascending, transverse, and descending colon using CT colonography (CTC). 50 supine CTC datasets from 50 asymptomatic, adult patients were analyzed (NCI-CBIIT instance of the National Biomedical Imaging Archive). At least 5 consecutive, pairs of unobscured haustral folds in each colonic segment were necessary to be included in this study. Of an initial 201 patients, 151 were excluded due to suboptimal colonic distension, retained fluid, tortuosity, and diverticulosis. For each dataset, the heights of the non-dependent haustral folds were measured in the ascending, transverse, and descending colon on 2D multiplanar reformations. Differences in mean HFHs were assessed using a hierarchical generalized linear mixed model. A total of 2079 colonic folds were measured: 625 in the ascending colon (including the cecum), 687 in the transverse colon, and 767 in the descending colon. The mean number of folds measured per segment was 6.87 ± 2.11. Mean HFHs were significantly taller in the ascending colon (14.62 ± 5.47 mm) than in the transverse (9.49 ± 3.65mm) or descending (6.53 ± 3.12mm) colon; mean HFHs were also significantly taller in the transverse than the descending colon, (P<0.0001, for all comparisons). In conclusion, taller colonic haustral folds are present in the proximal colon and may contribute to more frequently missed lesions (e.g. polyps) in the right colon by conventional, optical colonoscopy.

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Introduction

Colorectal cancer (CRC) is the second leading cause of cancer death in the United States today, with approximately 150,000 incident cases and 50,000 deaths reported each year [1, 2]. The five-year survival rate approaches 90% if the cancer is discovered at an early stage, but is 12% if distant metastases are present [2]. Since optical colonoscopy (OC) allows for the early detection and removal of precancerous adenomas, it is widely considered the best modality for colon cancer screening and prevention [3]. Several longitudinal studies have shown significant reductions in the incidence of CRC following colonoscopic polypectomy [4-7]. Nevertheless, colon cancer remains the third leading cause of cancer nationwide despite increasing rates of screening [2, 8].

Colonoscopy offers incomplete protection from colon cancer, and interval cancers unfortunately do develop in patients despite their having undergone optical colonoscopy. This may occur for a variety of reasons, including aggressive tumor biology resulting in a newly developed fast growing cancer or incomplete resection of a visualized adenomatous polyp, with recurrence progressing to cancer. Interval cancers may also arise if adenomatous polyps are missed at colonoscopy. Tandem colonoscopy studies have confirmed relatively high adenoma miss rates of 6-27% at OC, with miss rates for small polyps less than 6 mm in size at the upper end of this range [9-12]. For advanced adenomas the miss rate in one study was as high as 11% [9].

Polyps may be missed at colonoscopy for a variety of reasons, including inadequate bowel preparation, insufficient colonic distention, and poor colonoscopy technique including rapid withdrawal of the colonoscope in less than 6 minutes [13, 14]. Small or flat polyps may not be recognized by less experienced endoscopists. Performance of colonoscopy by a non-gastroenterologist, such as a primary care physician or surgeon, increases the chance that a potential cancer will be overlooked [15-18].

Recent evidence suggests that interval cancers are more likely to develop in the right (i.e. cecum and ascending colon) and transverse colon than in the left colon [15, 19-24]. The postulated reasons for this differential distribution are varied. A proportion of colonoscopies are incomplete where the cecum cannot be reached for technical reasons. In addition, less experienced endoscopists may occasionally mistake anatomic landmarks such as the hepatic flexure for the cecum. Incomplete colonoscopies with failure of cecal intubation are more likely when performed by endoscopists with less training, and/or lower procedural volumes [25], and when colonoscopy is performed in office settings [15, 25, 26]. A delay between completion of the bowel preparation and colonoscopy, can result in the accumulation of bile and thick mucus in the right colon [27], potentially obscuring polyps, particularly those that are flat. A disproportionate number of interval CRC with microsatellite instability arise in flat serrated lesions [28] that are more common in the proximal colon [29-31]. These adenomas tend to carry aggressive mutations like BRAF V600E [32, 33]. Because they are flatter and have ambiguous color and borders, serrated lesions are also harder to identify during optical colonoscopy [9, 34], especially if withdrawal times are rapid [35].

Adequate visualization of the mucosa on the proximal side of haustral folds remains a challenge given current limitations in standard colonoscope technology and endoscopic technique [36, 37]. Polyps on the proximal aspect of taller colonic folds may be more difficult to visualize at colonoscopy, and studies have indicated that many polyps missed at colonoscopy are on the proximal aspects of colonic folds [37]. We therefore hypothesized that the higher incidence of interval cancers in the proximal compared to the distal colon may stem in part from the existence of taller
hastral folds proximally. To our knowledge, this is the first study to investigate whether the anatomic heights of haustral folds may differ between the ascending (including the cecum), transverse, and descending colon using CT virtual colonoscopy and thus potentially help explain regional differences in interval CRC development.

**Materials and Methods**

The National Biomedical Imaging Archive (NBIA) is a free online national repository of PHI de-identified Digital Imaging and Communications in Medicine (DICOM) images sponsored by the National Cancer Institute Center for Biomedical Informatics and Information Technology. We reviewed 201 supine Computed Tomographic Colonography (CTC) patient datasets selected from the beginning of the NBIA’s Virtual Colonoscopy database [https://imaging.nci.nih.gov/ncia/login.jsf] made publicly available by the (Virtual) CT Colonography clinical trial [38]. Because the datasets were completely de-identified, this study was not considered human subjects research by our Institutional Review Board; thus we were granted permission to conduct our investigation.

A more complete description of the protocol has been previously published in the New England Journal of Medicine [39]. A total of 1233 asymptomatic adults patients age 40 to 79 from three clinical sites underwent same-day screening CTC followed by OC. Twenty-four hours prior to the procedure, patients underwent standard bowel preparation with 90 ml of oral sodium phosphate (Fleet 1 preparation, Fleet Pharmaceuticals) and 10 mg of bisacodyl. Tagging of solid stool with 500 ml of barium (2.1% by weight; Scan C, Lafayette Pharmaceuticals) and of luminal fluid with 120 ml of diatrizoate meglumine and diatrizoate sodium (Gastrografin, Braco Diagnostics) was performed. A single reviewer evaluated all data sets using a DICOM image viewer (OsiriX v4.0) for inclusion into this study. Inclusion criteria were: (1) adequate colonic distension and (2) presence of measurable folds. Having a minimum of at least five consecutive measurable haustral fold pairs in a given colonic segment was established as a marker of adequate bowel distention for optimal measurement of folds. Measurable folds were defined as folds that were not obscured or distorted by retained fluid, luminal collapse, tortuosity or diverticulosis. As haustral folds meet at the teniae coli, two of the three folds were each measured with the most dependent fold excluded; the most dependent folds were most often obscured by fluid if present (Figure 1A). Measurements of fold heights were made on 2D multiplanar reformations; the imaging plane demonstrating the folds in cross-section was established for optimal measurement of fold heights (Figure 1B). Due to the high frequency of tortuosity, suboptimal distention relative to other colonic segments, distortion of the folds and diverticulosis relative to other colonic segments, the sigmoid colon was not included in this study.

![Figure 1A](image-url)
Fifty patient datasets were included that met the minimum criterion for fold measurement in the ascending (including the cecum), transverse, and descending colonic segments and were of sufficient diagnostic quality. This methodology allowed us to achieve an optimal dataset with complete measurements in all three colonic segments for each of the included patients. The segments could thus be compared directly to each other within a patient, so that each patient served as their own internal control for potential demographic or clinical confounders. Differences in average fold height were assessed by statistical analysis using a hierarchical generalized linear mixed model with log link function, in which colon region served as a fixed factor and the patient served as a random factor. Both 95% Confidence Intervals (CI) and P-values were generated using STATA (Release 9.2; StataCorp LP, College Station, TX).

Results

We measured hastral fold heights (HFHs) of the colon in 50 patients. The mean age of the study patients was 60.5 ± 6.8 years. Thirty-five of the patients were male and 15 were female. A total of 2079 colonic folds were measured: 625 in the ascending colon (including the cecum), 687 in the transverse colon, and 767 in the descending colon. The mean number of folds measured per segment was 6.87 ± 2.11. This minimum number of five fold pairs was within one standard deviation of the mean number of fold pairs measured per segment (6.87 ± 2.11). Figure 2 shows histograms of the distribution of colon fold measurements in each of the three colonic segments. Mean hastral fold heights in the ascending colon (including the cecum) (14.62 ± 5.47 mm) were significantly taller than in the transverse (9.49 ± 3.65 mm, P<0.0001) or descending colon (6.53 ± 3.12 mm, P<0.0001) (Table 1). Folds in the transverse colon were also significantly taller than in the descending colon.
(P<0.0001). The ascending colon (including the cecum) folds were 7.60mm (95% CI: 7.09-8.11 mm) and 4.61mm (95% CI: 4.12-5.11 mm) taller than the descending or transverse haustral folds, respectively (Table 2).

<table>
<thead>
<tr>
<th>Region of Colon</th>
<th>Mean height ± standard deviation in millimeters</th>
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<tbody>
<tr>
<td>Ascending (with cecum)</td>
<td>14.62 ± 5.47</td>
</tr>
<tr>
<td>Transverse</td>
<td>9.49 ± 3.65</td>
</tr>
<tr>
<td>Descending</td>
<td>6.53 ± 3.12</td>
</tr>
</tbody>
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Table 1: Regional Mean Heights of Colon Haustral Folds

Discussion

Our study demonstrates that the height of haustral folds increases as one moves proximally in the colon. Folds in the ascending colon (including the cecum) were significantly taller than those in the transverse and descending colon (P<0.0001). Transverse colonic folds were likewise taller than those in the descending colon (P<0.0001). Such differences in regional anatomy may potentially impact the sensitivity of optical colonoscopy in detecting polyps and cancers in different parts of the colon. Polyps hidden on the proximal aspects of the taller folds in the ascending (including the cecum) and transverse colon may be harder to detect at optical colonoscopy. This may be one additional factor contributing to the lower protection offered by optical colonoscopy in the right and transverse colon.

Optical colonoscopy has long been considered the gold standard for colorectal cancer screening because of its therapeutic capability for resecting adenomatous polyps identified during mucosal inspection [6, 7]. However, colonoscopy offers incomplete protection against colon cancer, particularly in the right colon (i.e. cecum and ascending colon). Many factors may explain the higher rates of right-sided interval cancers [15, 17, 19-24, 40]. For example, the endoscopist may not be able to reach the cecum. In large academic gastroenterologist run endoscopy units in the U.S. and Europe, the incomplete colonoscopy rate is around 1.5-3.5% [22, 41, 42]. The rate of incomplete colonoscopy is likely to be higher among less experienced endoscopists, in community settings, and at low volume centers. Despite bowel prep, adherent bile in the right colon may impair visualization of flat polypse. In addition there is a higher prevalence of serrated flat lesions with aggressive tumor markers located in the proximal colon [29-31]. Such flat lesions are difficult to identify at colonoscopy [35]. The limited view offered behind prominent mucosal folds by OC is a potential reason that traditional colonoscopy misses some polyps [36, 37]. An early study published in 1978 by Miller et al. found that endoscopists tended to neglect lesions situated on the proximal side of valves and haustral folds that were subsequently identified by barium enema [36]. Pickhardt et al. also found that

<table>
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<tr>
<th>Region of Colon</th>
<th>Difference of mean heights (millimeters)</th>
<th>95% Confidence Interval</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ascending vs. Transverse</td>
<td>4.61</td>
<td>4.12-5.11</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Ascending vs. Descending</td>
<td>7.6</td>
<td>7.09-8.11</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Transverse vs. Descending</td>
<td>2.98</td>
<td>2.63-3.34</td>
<td>&lt;0.0001</td>
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</table>
71.4% of lesions missed at colonoscopy but found on CTC were located on the proximal aspects of colonic folds [37]. Retroflexion of the colonoscope does allow examination of the proximal aspect of colonic folds and in one study, retroflexion of the colonoscope during withdrawal through the right colon improved detection of adenomas in the right colon in 5.8% of patients [43]. Unfortunately, retroflexion of standard colonoscopes in the proximal colon is difficult due to the large turn radius of the tip of the colonoscope, and the majority of endoscopists do not practice this technique.

Since colonic folds can obscure the view of polyps, and more interval cancers develop in the right colon despite colonoscopy, we postulated that taller folds might exist in the cecum, ascending and transverse colon where more interval cancers develop. Our study has confirmed this hypothesis. To our knowledge, our study is the first to document higher colonic folds in the ascending colon compared with the transverse and descending colon. We were able to show that colonic haustral folds become increasingly taller as one moves proximally along the colon, with folds of the transverse colon taller than descending colon, and ascending colonic folds taller than transverse colonic folds. This additional height of right colonic folds may further impede thorough visualization of lesions located on the proximal aspect of these taller folds [36, 37], which may be an additional important factor, explaining why colonoscopy offers less protection from interval colon cancer in the right colon. Moreover, unlike colonoscopy, the area behind colonic folds is not a blind spot for CTC. CT Colonography may thus prove advantageous for detecting polyps located on the proximal aspect of colonic folds [37]. In one meta-analysis of CTC, only 6 of 16 cancers located proximal to the splenic flexure were missed compared with 10 cancers missed in the rectosigmoid [47]. This would suggest that haustral fold heights may contribute to discrepant regional rates of colon cancer; however, future, longer-term research is needed to determine if relative rates of right-sided colon cancers are decreased with CTC screening or other advanced endoscopic techniques that employ 360° or bidirectional endoscopy.

Our study has several limitations. First, haustral fold morphology can vary depending on luminal distention. Colonic fold thickening has been described with insufficient luminal distention [48], while overdistention can lead to fold effacement. While only patients with well-distended colonic segments were included in this study, the effects of subtle differences in regional bowel distention on fold heights could not be adjusted for. Second, our ability to identify measurable colonic folds was markedly diminished in the sigmoid colon compared to other colonic segments as previously described, leading us to exclude the sigmoid colon from CRC in the splenic flexure and descending colon (PR 0.36, 95% CI 0.16-0.82) when compared to patients without colonoscopy [22]. Such findings suggest that OC may be a less sensitive modality in the cecum, ascending and transverse colon, which may be partially explained by the taller folds we have documented in those regions of the colon.

Several studies have suggested that CTC may have comparable sensitivity and specificity to colonoscopy, especially in detecting advanced neoplasms [44, 45]. Addition of 3-D “fly-through” interpretation [46] and preparation with both cathartic and tagging agents [47] may further improve the screening capability of CTC. Moreover, unlike colonoscopy, the area behind colonic folds is not a blind spot for CTC. CT Colonography may thus prove advantageous for detecting polyps located on the proximal aspect of colonic folds [37].
the analysis. Thus, our inference is limited to the descending colon, and does not necessarily generalize to other parts of the left colon. In addition we measured fold heights only in datasets that included at least five consecutive measurable folds in all three colonic segments. It is possible that taller folds may have existed elsewhere in these segments but were not measured because they were isolated and thus did not fit our minimum criterion. We also excluded patient datasets if measurable folds could not be identified in one of the colonic segments. While this inclusion criterion allowed us to achieve a dataset where every patient had complete comparable measurements in the colonic segments, it also limited our patient sample size and led to imbalance on the potential effect modifier of sex. Future studies would need to reproduce these results in larger populations with a balanced proportion of males and females to ensure there is no interaction by gender.

**Conclusion**

In conclusion, our study indicates that regional differences in haustral fold height exist, with significantly taller folds found in the proximal colon. The potential structural obstruction of view caused by taller haustral folds in the proximal colon, may possibly offer an additional factor explaining the lower protection against interval colon cancer offered by standard optical colonoscopy in the right and transverse colon compared to the left colon. CT Colonography and advanced colonoscopy techniques (e.g. bidirectional or 360° viewing) offer advantages to evaluate and detect precancerous adenomatous polyps and lesions located behind folds, which may be most valuable in the proximal colon as it may translate into lower rates of right-sided colon cancers. Future trials are needed to compare the protection offered by these techniques relative to traditional OC in the prevention of interval colorectal cancer.

**Acknowledgements**

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**Abbreviations**

Colorectal cancer (CRC)
Computed Tomographic Colonography (CTC)
Confidence intervals (CIs)
Haustral fold heights (HFHs)
Optical Colonoscopy (OC)
National Biomedical Imaging Archive (NBIA)
Digital Imaging and Communications in Medicine (DICOM)

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