Proximal versus distal hyperplastic polyps of the colorectum: different lesions or a biological spectrum?

K Baker, Y Zhang, C Jin, J R Jass

Background: Because of their suggested link with microsatellite instability high colorectal cancers, right sided hyperplastic polyps (HPs) may differ from their distally located counterparts. This is highlighted by the recognition of a variant HP, termed sessile serrated adenoma (SSA), which predominates in the proximal colon. HPs displaying the morphological features now associated with SSAs have been shown to have altered expression of “cancer associated” markers, but no studies have investigated whether this is dependent on anatomical location of the polyps.

Aims: To evaluate morphological and functional features in right versus left sided HPs from patients without colorectal cancer with the aim of identifying distinguishing characteristics.

Methods: HPs originating in the proximal and distal colorectum were histochemically and immunohistochemically stained to evaluate a panel of markers related to proliferation and differentiation. In addition, a series of morphological features was evaluated for each polyp.

Results: Crypt serration, crypt dilatation, and horizontal crypt growth were more common among HPs from the right side, whereas histochemical factors including mucin changes, global methylation status, and expression of carcinoembryonic antigen were not significantly different. An age disparity was also seen between patients with right versus left sided lesions, with patients with right sided lesions being an average of more than 10 years younger than those with left sided lesions.

Conclusions: These findings suggest that right and left sided HPs differ mainly in terms of growth regulation rather than cellular differentiation, implying that these lesions belong to a continuous spectrum of serrated polyps that differ quantitatively rather than qualitatively.

Although most instances of sporadic colorectal cancer (CRC) seem to arise through the “adenoma–cancer” sequence,1 a subset of cases that includes microsatellite instability high (MSI-H) CRCs cannot be accounted for by this model.2–4 Instead, most MSI-H carcinomas appear to progress along a “serrated pathway”,5–8 implicating the inactivation of the DNA mismatch repair gene hMLH1,9 as a result of methylation of its promoter region.10–11

“There is a need to identify markers for high risk hyperplastic polyps, in addition to mechanisms that would explain the suggested rapid evolution of the serrated pathway”

Although the nature of the antecedent lesion for sporadic MSI-H cancers is still the subject of debate, there is increasing evidence for the precursor being either a hyperplastic polyp (HP) or a closely related lesion. MSI-H CRCs are thought to arise within a subset of HPs that tend to be large and proximally located. Because MSI-H cancers are predominantly proximal lesions, whereas HPs are uncommon in this location, it appears that right sided HPs have a higher potential for progression to malignancy compared with their left sided counterparts.9–17 However, it is clear that only a small proportion of even these HPs will undergo malignant transformation.18 There is a need to identify markers for high risk HPs, in addition to mechanisms that would explain the suggested rapid evolution of the serrated pathway.15

Recent studies have highlighted a variant HP termed sessile serrated adenoma (SSA), which occurs more frequently in the proximal colon.8–14 An earlier study showed that HPs with the morphological features now associated with SSAs were also characterised by altered expression of “cancer associated” markers, including carcinoembryonic antigen (CEA) and sialic acid variants.19 However, it was not shown whether these features were dependent upon the anatomical location of polyps. Our present study aimed to investigate whether there are particular morphological or functional characteristics that differentiate right from left sided HPs in patients without CRC. The present findings indicate that left and right sided HPs are distinguishable on the basis of morphological features that are identifiable through the examination of routine diagnostic slides but do not differ with respect to functional markers.

MATERIALS AND METHODS

Tissues

HPs were obtained from a total of 23 patients (16 men and seven women) who underwent full colonoscopy at the Royal Victoria Hospital (Montreal, Canada) during a six month period. The patients had either right sided or left sided HPs, but not both. A randomly selected subset of the more numerous left sided HPs was included for study. Polyps were considered right sided if they occurred proximal to the splenic flexure and left sided if they occurred distal to the splenic flexure. Fourteen of the patients had a single HP, five had two HPs, three had three HPs, and a single patient had five HPs. In all, 19 right and 19 left sided polyps were obtained from 11 and 12 patients, respectively. Our study received approval from the institutional review board of the faculty of medicine of McGill University, Canada.

Abbreviations: CEA, carcinoembryonic antigen; CRC, colorectal cancer; HP, hyperplastic polyp; IEL, intraepithelial lymphocyte; 5-MeC, 5-methylcytosine; mPAS, mild periodic acid Schiff; MSI-H, microsatellite instability high; Ste*, sialyl Lewis*; SSA, sessile serrated adenoma
Scoring of morphological and patient features

Using the haematoxylin and eosin stained slide, the following morphological features were evaluated in up to 20 appropriately oriented crypts for each polyp: (1) ratio of intraepithelial lymphocytes (IELs) to the total number of epithelial cell nuclei counted, (2) number of horizontal crypts (extending parallel to the muscularis mucosae) of the total number of crypts counted, (3) number of branched crypts of the total number of crypts counted, (4) number of dilated crypts of the total number of longitudinally oriented crypts counted, (5) number of dilated crypts of the total number of transversely oriented crypts counted, (6) longitudinal serration index (number of “teeth” of the total number of longitudinally oriented crypts counted), (7) transverse serration index (number of teeth of the total number of transversely oriented crypts counted), (8) number of crypts showing serration in the lower third of the total number of longitudinally oriented crypts counted, and (9) number of crypts showing serration in the lower third of the total number of transversely oriented crypts counted. Data from transversely and longitudinally oriented crypts were added together to give a global assessment of polyps. Patient age, sex, and site of lesion (right versus left colon) were recorded for each HP.

Antibodies and staining

Sections (4 μm) of formalin fixed, paraffin wax embedded tissues were cut, mounted on to coated slides, and dried overnight at 37°C. Immunohistochemical staining was performed using recommended dilutions of mouse monoclonal antibodies against Ki-67 (clone MIB-1; Dako, Carpinteria, California, USA), sialyl Lewis ε (SLεε; clone 2Q539; US Biological, Sampscott, Massachusetts, USA), CEA (clone C6G9; Sigma, St Louis, Missouri, USA), and 5-methylcytosine (5-MeC; clone 162 33 D3; Oncogene, San Diego, California, USA). For all antibodies except anti-5-MeC, sections were dewaxed by routine techniques and heat induced antigen retrieval was performed in 0.001M EDTA buffer (pH 8.0) in a microwave oven for 15 minutes. Sections were subsequently incubated at room temperature for one hour with primary antibodies and then with a peroxidase labelled polymer (EnVision + System/HRP, K4005; Dako) for 30 minutes. Sections were subsequently incubated with the chromogen 3-amino-9-ethylcarbazole, counterstained with haematoxylin, and mounted using an aqueous medium. For staining with anti-5-MeC, antigen retrieval and staining were carried out as described previously. Detection was performed using a Histostain Plus kit (85–9643; Zymed, San Francisco, California, USA) with 3,3-diaminobenzidine as the chromogen. Appropriate positive and negative controls were carried out for each stain. For each patient, one slide was also stained using a routine haematoxylin and eosin stain and a mild periodic acid Schiff (mPAS) stain, as described previously.

Scoring of immunohistochemical features

Scoring was carried out without knowledge of the site from which each polyp had been excised. For staining by mPAS, anti-CEA monoclonal antibody, and anti-SLe ε monoclonal antibody, lesions were scored for both intensity (0 for no staining, 1 for pale staining, 2 for moderate staining, 3 for dark staining) and distribution (0 for no staining, 1 for heterogeneous/patchy staining, 2 for diffuse staining). Ki-67 staining was evaluated by counting the number of stained cells/total number of epithelial cells in the lower, middle, and upper third of three well oriented, longitudinal crypts from each polyp. 5-MeC staining was evaluated by counting the number of unstained nuclei/total number of epithelial nuclei in the lower, middle, and upper third of five well oriented, longitudinal crypts from each polyp. Overall intensity of 5-MeC staining was also evaluated through comparison with the intensity of staining of stromal nuclei in the same tissue section (0 for identical intensity as stromal nuclei, 1 for greater intensity than stromal nuclei, 2 for lower intensity than stromal nuclei).

Analysis

HPs were divided into two groups based upon site of origin and the differences between these two groups were subsequently analysed. Age, sex, size of largest fragment, Ki-67 staining, 5-MeC staining, and all morphological features were analysed using the Student’s t test. mPAS staining, in
addition to anti-SLe\textsuperscript{x}, anti-CEA, and anti-5MeC intensity were analysed by the Wilcoxon test. In all cases, p values < 0.05 were considered significant.

RESULTS
Patient features
No difference was detected in the sex distribution between patients with left and right sided polyps. In contrast, age difference proved to be significant (p = 0.004), with patients having right sided lesions being, on average, more than 10 years younger (mean age, 50.37 years) than those with left sided lesions (mean age, 61.84 years).

Morphological features
Crypt branching (p = 1.000) showed no significant predilection for either side, and neither did the number of IELs (p = 0.970). Significant differences between proximal and distal HPs were noted for global crypt dilatation (p = 0.010), global serration index (p = 0.007), and global basal serration (p = 0.002) (fig 1). p Values indicated greater significance for well oriented longitudinal crypts, but significance was maintained even after factoring in non-significant transversely cut crypts to obtain a global score. However, it should be noted that perfectly oriented crypts were poorly represented in larger polyps because of the convoluted crypt orientation in larger lesions. The presence of horizontal crypts also proved to be different (p = 0.020) between polyps from each side of the colorectum.

Monoclonal antibodies and mPAS staining
No differences emerged between left and right sided polyps for global SLe\textsuperscript{x} staining (p = 0.270), although the value for SLe\textsuperscript{x} intensity did approach significance (p = 0.087) (fig 2). Results for global mPAS staining fell just short of significance (p = 0.059), as did those for mPAS intensity (p = 0.057). Although both of these stains detect changes in acetylation patterns of colorectal sialic acid, a lack of correlation between these staining results was noted for many individual polyps (data not shown). No differences were noted for either intensity or distribution when staining for CEA. Staining for Ki-67 and 5-MeC was not significantly different between left and right sided HPs in the three crypt regions evaluated (fig 3).

DISCUSSION
In our study, morphological characteristics were found to be distributed differently among right and left sided HPs. In particular, the presence of crypt dilatation, horizontal crypts, serration in the lower third of crypts, and a high serration index were associated with a proximal HP location. Recent studies have identified an HP variant that is distinguished from classic HPs on the basis of multiple morphological features. These lesions, termed sessile serrated adenomas (SSAs), show a predilection for the proximal colon and have been linked with MSI-H CRC. A high proportion of right sided HPs show the features of SSAs. However, it is unclear whether SSAs and HPs are fundamentally different lesions, or represent a continuous spectrum of “serrated polyps”. In an earlier study, increased expression of CEA and altered sialic acid composition co-segregated with certain morphological features that have now been associated with SSAs. The aim of our present study was to determine whether functional markers expressed by right and left sided HPs highlight biological differences that are not only more diagnostically discriminating than the subtle morphological changes described by others, but may also serve as evidence that proximal and distal HPs are biologically distinct lesions.

Early studies indicated that HPs are age related lesions, but these dealt only with HPs in the rectum. The finding of a full decade of difference in age between patients with proximally versus distally located HPs was unexpected, given the predominance of right sided bowel cancers in older patients. It has been estimated that only one in 25 right sided HPs will progress to an MSI-H CRC. The relatively young age of subjects with right sided HPs in our study suggests that the lesions may remain unchanged for many years before malignant transformation.
No significant differences were found between left and right sided polyps in terms of staining with either mPAS or antibodies against SLeα. It is known that changes occur in the structure of sialic acid (loss of O-acetylation) secreted by malignant colorectal epithelium,20–23 and that these changes may be revealed by mPAS staining. SLeα is detectable in cancerous and precancerous lesions but not in normal colorectal mucosa. The basis for this observation is the same change in sialic acid structure (loss of O-acetylation) that renders SLeα antigenic.24 Because no significant difference was detected between the right and left sided HPs, which nevertheless differed from the normal colorectal mucosa (results not shown), it appears that these mucin histochromatological alterations are not specific for the anatomical site of origin of HPs. The trend towards increased sialic acid abnormality in right sided HPs might assume significance in a larger series. However, the findings suggest that proximal and distal HPs are related lesions that differ quantitatively rather than qualitatively. It would be of interest to examine the mucin core proteins present in right versus left sided HPs because a previous study by Biemer-Hüttmann et al found that serrated polyps were associated with upregulation of intestinal mucin MUC2 and gastric mucin MUC5AC, both of which are known to be upregulated in mucinous MSI-H cancers.5

"The increased serration seen in right sided hyperplastic polyps in our present study may be explained by a greater antiapoptotic effect in this subgroup"

Staining with anti-CEA antibody yielded no significantly different results for either intensity or distribution between the two groups of polyps. Previous studies have found increased CEA expression in HPs,16 19 a finding that parallels the commonly observed increase in CRCs.18 20 24 26 This increased expression of CEA, a GPI linked transmembrane protein implicated in cell signalling,27 could conceivably serve as an indicator of potentially neoplastic progression, even though it does not seem to be differentially expressed in right versus left sided HPs. The lack of a difference in numbers of IELs present in polyps from opposite sides of the colon suggests that lymphocytes are not recruited in abnormally large numbers to areas of hyperplasia, but only with the advent of dysplasia.

The lack of a significant difference in proliferation zone length between distal and proximal HPs, as indicated by staining for the proliferation inducible Ki-67 antigen, implies that expansion of this zone may be common to all subsets of HPs. An increase in cell proliferation, without an accompanying increase in the rate of anoikis (exfoliation induced apoptosis at mucosal surfaces), would represent a possible mechanism for the development of glandular serration.4 The increased serration seen in right sided HPs in our present study may be explained by a greater antiapoptotic effect in this subgroup. It is possible that the observed morphological disparity results from different underlying mechanisms for inhibiting apoptosis. This is supported by the fact that apoptosis inhibition by mutated K-ras is linked to the Akt/ protein kinase B mediated inhibition of BAD and caspase 9, whereas BRAF mutation blocks caspase activation downstream of BAD mediated cytochrome c release.30 31 BRAF mutation occurs more frequently in large, right sided HPs occurring in the condition hyperplastic polyposis.12

With respect to staining for 5-MeC, the lack of a significant difference in the upper, middle, or lower areas of the crypts from right versus left sided polyps indicates a more or less common DNA methylation status in polyps from either side of the colorectum. This particular feature was evaluated because, on the one hand, MSI-H colorectal cancers are associated with hypermethylation in the promoters of certain genes10–12 whereas, on the other hand, global hypomethylation has been associated with many cancers.20 24 26 28 32 Although no consistent methylation differences were apparent, it is notable that only epithelial nuclei from HPs located in the distal colorectum were found to be globally methylated to a greater extent than those of their stromal counterparts. All right sided HPs contained either equally or less methylated epithelial nuclei. A possible explanation for this could be the previous suggestion that global hypermethylation may result from increased methylation at sites of DNA damage, which decreases the general availability of methyltransferases.34 In conclusion, this investigation has shown a significant association between proximal HPs and the presence of horizontal crypts, crypt dilatation, serration in lower crypt thirds, and a high serration index. This is consistent with findings by others.2 24 26 Because the main differences in our study proved to be morphological, rather than functional, it appears that proximal HPs show an exaggerated imbalance of cellular proliferation rather than a dysregulation of cellular differentiation compared with their distal counterparts. The structural differences found in the right sided polyps may be attributable to pressure from increased cell numbers within individual crypts, in turn reflecting a more profound impairment of apoptosis.

ACKNOWLEDGEMENTS

The authors would like to thank Dr B Case from the Department of Pathology, McGill University (Montreal, QC, Canada) for his advice on statistical analysis.

Authors’ affiliations

K Baker, Y Zhang, C Jin, J R Jass, Department of Pathology, McGill University, Montreal, Quebec Canada, H3A 2B4

REFERENCES

33 Arnold CN, Goel A, Boland CR. Role of MLH1 promoter hypermethylation in drug resistance to 5-fluorouracil in colorectal cancer cell lines. Int J Cancer 2003;106:66–73.
Proximal versus distal hyperplastic polyps of the colorectum: different lesions or a biological spectrum?

K Baker, Y Zhang, C Jin and J R Jass

J Clin Pathol 2004 57: 1089-1093
doi: 10.1136/jcp.2004.016600

Updated information and services can be found at:
http://jcp.bmj.com/content/57/10/1089

These include:

References
This article cites 43 articles, 17 of which you can access for free at:
http://jcp.bmj.com/content/57/10/1089#BIBL

Email alerting service
Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

Topic Collections
Articles on similar topics can be found in the following collections

Colon cancer (231)
Immunology (including allergy) (1664)

Notes

To request permissions go to:
http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/