Innominate Grooves of the Colon: Histological Reappraisal

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Abstract. Background: Innominate grooves (IGs) in the colon are ubiquitous structures limiting anhemic folds. The histology of the crypts in anhemic folds is well known but the histology of IGs has remained unattended. In this study, IG crypts in the normal mucosa and in ulcerative colitis (UC) were analyzed. Patients and Methods: IGs displayed a single central-lumen with merging tributary crypts. The number of merging tributary crypts, at both sides and at the bottom of the central lumen, were quantified in 60 IGs: 30 in normal colons and 30 in UC. Results: A total of 152 (median=5) tributary crypts were found merging in the 30 IGs in the normal colon, and 170 (median=5) in the 30 IGs in UC. The difference was non-significant (p=0.08012). Conclusion: Chronic inflammation did not influence the frequency of tributary crypts in IGs. The shape and the length varied considerably in the 60 IGs. Given that the chief putative physiological function of IGs is the expansion of the colon when submitted to increased intraluminal pressure, it is not inconceivable that following that demand, IGs act as hinges that permit grooves to level with the rest of the surrounding mucosa.

In 1965, William observed a fine network of lines in about 25% of routine colon barium examinations (1). The barium lines or innominate grooves (IGs), as they were named, were arranged linearly or in a reticular pattern, 0.5 mm apart. In 1977, Frank et al. identified marginal spiculations and punctate collections of barium in patients having a normal colon (2). The punctate pattern corresponded to the crypts of Lieberkühn which filled with barium, and the marginal spiculations to the barium retained in the IGs (2). In 1977, Matsuura et al. found innominate lines in about 90% of double-contrast examinations, when the colon was partially collapsed (3). Those lines were described as superficial indentations of the mucosa with depths of 0.23 mm. In 1978, Cole described the gross characteristics of innominate lines and grooves in the colonic mucosa with the aid of a dissecting microscope (4). Three basic mucosal patterns emerged: i) Linear circumferential, ii) network, and iii) mixed. Light microscopy showed innominate grooves running transversely across the mucosa every few millimeters, between the anhemic folds (4). Subsequently, in radiological images, Treugut et al. found that the most common localizations of IGs were the transverse colon near the hepatic flexure, and the descending and sigmoid colon (5). Rubesin et al. found a flat, featureless surface when the colonic mucosa was stretched in barium radiography (6). The colonic grooves re-appeared when the muscularis spontaneously contracted, creating a finely nodular surface. It was concluded that innominate grooves were more often seen when the colon was partially collapsed either during colonic spasm or on post-evacuation radiographs. Dissecting microphotography revealed that the grooves were clefts in the mucosa that disappeared with colonic distention. Once the colonic mucosa was stretched, a flat, featureless surface was seen. The caption in one of their illustrations reads: “The presence and visibility of the innominate grooves of the colon are a function of colonic distention” (6). Levine and Haggitt were the first to report IGs as cloverleaf-like crypts connect to a single lumen using light microscopy (7). These grooves limit the anhemic folds, built of regular crypts organized as regular ‘test tubes’ of similar size and shape (7).

A more recent review of the literature revealed that IGs are mentioned in many publications or described as cloverleaf-like crypts connect to a single lumen (7), giving the false impression that all IGs have a similar microscopic pattern. The possibility that IGs might vary in histological appearance in colonic mucosa with chronic inflammation has not been entertained.

Recently, we reported non-dysplastic colonic crypts in symmetric and asymmetric fission in colectomy specimens from patients with UC (8, 9). During those studies, we noticed that IGs adopted various architectural modalities. The question arose as to whether IGs in UC differ from those in the normal colonic mucosa.

The purpose of this survey was to study the microscopic architectural anatomy of IGs in detail, aiming to explore possible differences between normal colonic mucosa (10) and UC.

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Patients and Methods

The study material consisted of 60 consecutive IGs: 30 found in seven colectomy specimens from patients with UC (9) and 30 found in 22 normal colon segments, proximal or distal to surgically removed colonic adenocarcinoma (10).

Definitions. Histology of IGs. The microscopic anatomy of IGs was forensically analyzed. IGs were often found in a slightly depressed mucosal area with a luminal orifice connected to a single central luminal axis, to which horizontal to oblique crypts varying in length and shape usually merged at various levels. Some merging crypts were rudimentary. Merging crypts in IG were called tributaries (Figures 1-3).

Left tributary crypts: Merging crypts located on the left side of the central luminal axis.

Right tributary crypts: Merging crypts located on the right side of the central luminal axis.

Bottom tributary crypt: A tributary crypt usually found at the bottom of the central luminal axis of an IG; this tributary crypt was often lined with distinctive goblet cells. The thickness of the intervening lamina propria between the tributary crypts also varied.

Counting of tributary crypts in IGs. The number of the tributary crypts found on the left side, on the right side, and at the bottom of the central luminal axis of the IG, was separately recorded.

Statistical analysis. The two-tailed non-parametric Mann–Whitney U-test was applied to compare difference between IGs and normal colon mucosa. Statistical significance was defined as \( p < 0.05 \).

Results

The shape and length of tributary crypts on both sides of the central lumen in normal segments and in UC. The shape and the length varied considerably in the 60 IGs. No apparent difference in length or shape between IGs in normal colonic mucosa and UC was found, despite the moderate to severe inflammation present in the lamina propria in UC. Examples of the shape and length asymmetry of tributary crypts found in the same side and at both sides of the central lumen in IG are shown in Figures 1 to 3.

Quantification of all tributary crypts merging in IGs in normal colonic segments and in UC. The results in Table I indicate that 152 (median=5) tributary crypts were found merging in the 30 IGs in normal colon, and 170 (median=5) in the 30 IG in UC. The difference was non-significant \( (p=0.08012) \).

Quantification of tributary crypts merging on the right side of the central luminal axis of IGs in normal colon segments and in UC. Table I also shows that the median number of tributary crypts merging into the right-side of the central luminal axis in normal colon segments was 2, and in UC was also 2. The difference between the two groups was not significant \( (p=0.13622) \).

Quantification of tributary crypts merging on the left side of the central luminal axis of IGs in normal colon segments and in ulcerative colitis. Table I shows that the median number of tributary crypts merging into the left side of the central luminal axis in normal colon segments was 2 and in UC was also 2. The difference between the two groups was not significant \( (p=0.4009) \).

Quantification of tributary crypts merging at the bottom of the central luminal axis of IGs in normal colon segments and in UC. The median number of tributary crypts merging at the bottom of the central luminal axis in normal colon segments was 1 and in UC was also 1 (Table I). The difference between the two groups was not significant \( (p=0.99202) \).

Discussion

The present survey provides, for the first time, qualitative and quantitative histological data on merging tributary crypts in IG. The main findings were:

i) The histological architecture of IGs consisted of a collection of tributary crypts merging at both sides of a sagittal luminal virtual axis, and a tributary crypt merging at the bottom of the central lumen. Hence, the characteristic ‘test tube’ arrangement found in anthemic folds (7) was not recreated in the colonic mucosa of IG.

ii) The length and shape of the tributary crypts varied both within the same sagittal plane, between sides and at the bottom of the central lumen (cf Figures 1-3).

iii) The difference in length and shape in tributary crypts contrasted with the similar length and shape in crypts found in the anthemic folds (7) (Figure 1, left).

iv) The amount of lamina propria found between the tributary crypts varied considerably, whereas the amount between the crypts arranged as test tubes in the anthemic folds was usually similar (7).
v) Some of the tributary crypts were ‘rudimentary’. It may be argued that these rudimentary tributary crypts were not at the same cutting plane as the other tributary crypts. On the other hand, no rudimentary crypts were found in the anheticic folds.

vi) Quantitative analysis revealed no particular differences in the number of tributary IG crypts found in the colonic mucosa in normal colonic segments or in UC. It would thus appear that chronic inflammation does not influence the
frequency of tributary crypts in IGs. These results are at variance with those found in anhemic folds in UC, where crypt distortions are often found (9).

The molecular signaling that orchestrates crypt replication in anhemic folds have been amply investigated; WNT is known to control cell proliferation, differentiation and apoptosis along the crypt (11); adenomatous polyposis coli (APC) forms a gradient of cellular adhesion along the crypt axis, prompting more proliferative cells to reach the top of the crypts (12); phosphatase and tensin homolog deleted on chromosome 10, a protein encoded by the phosphatase and tensin homolog (PTEN) gene, regulates crypt formation (13); and sodium-hydrogen exchanger regulatory factor 1 (NHERF1) protein controls colonic gland morphogenesis (14). On the other hand, surprisingly, no studies have been carried out to explore whether similar or different molecular signaling is required to coordinate the replication of tributary crypts of IG.

The pertinent question is do similar signaling molecules that orchestrate the required stimuli for anhemic folds also act in IG tributary crypts, or are tributary crypts in IGs programmed by a different molecular agenda? This question deserves to be answered before the true biological significance of IGs, both in UC and in the normal mucosa, can be fully understood.

The chief physiological function of IGs is the expansion of the colon when it is submitted to substantially increased intraluminal pressure (6). Accordingly, it is not totally inconceivable that following such a demand, the merging
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