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Histogenesis of human colorectal adenomas and hyperplastic polyps: the role of cell proliferation and crypt fission

W-M Wong, N Mandir, R A Goodlad, B C Y Wong, S B Garcia, S-K Lam, N A Wright

Background: The histogenesis of human colorectal hyperplastic polyps and colorectal adenomas is poorly understood even now. Human colorectal adenomas, hyperplastic polyps, and normal colorectal mucosae (patients with familial adenomatous polyposis and hereditary non-polyposis colorectal carcinoma were excluded) were obtained during colonoscopy and microdissected into individual crypts. Morphology, cell proliferation characteristics, and fission indices of crypts isolated from these lesions were then studied.

Results: Crypts isolated from colorectal adenomas and colorectal hyperplastic polyps were significantly larger (p<0.001) than crypts from normal colorectal mucosa. Crypt fission was an uncommon event in normal colonic mucosae but common in crypts isolated from adenomas and hyperplastic polyps (p<0.001). Analysis of the distribution of mitoses suggested an upward expansion of the proliferation compartment in adenomas to the surface of the crypt with no reversal of proliferating cell distribution, as has previously been described.

Conclusions: Sporadic human colorectal adenomas and hyperplastic polyps grow by the process of crypt fission. Expansion of the proliferative compartment was demonstrated in crypts from adenomas, consistent with deregulation of cell cycle control.

There is growing evidence to support the idea that accumulated genetic changes in important genes, occurring in vulnerable cells, ultimately lead to the development of neoplasia. This process is well illustrated by colorectal cancer in which there is progressive accumulation of mutations during the gradual transition from normal mucosa to carcinoma. One of the earliest events in colonic carcinogenesis occurs at the adenomatous polyposis coli (APC) gene. Hyperproliferation is believed to develop in preneoplastic epithelium, implying that an APC mutation is involved in the process. None the less, a recent study of preneoplastic intestine of human familial adenomatous polyposis (FAP) and multiple intestinal neoplasia (MIN) mice which harbour the APC mutations has shown that the major abnormality is elevated rates of crypt fission, a process that begins by basal bifurcation and is followed by longitudinal division of the crypt.

In the same study, direct observations of three dimensional reconstruction of FAP microadenomas showed that both normal and atypical crypt fission events were increased in FAP and MIN adenomas and indicated that adenomas expanded mainly by the process of crypt fission. Crypt fission is very prominent in neonatal animals, leading to a progressive increase in crypt number in both the small and large intestine, and it is involved in the emergence of wholly mutated crypts from partially mutated crypts in mice after injection of carcinogens. The Dib-1 locus in mice generates an intestinal binding site for the lectin Dolichos biflorus agglutinin, and study of the induction of mutated Dib-1 locus in mice by N-nitroso-N-ethylurea in SWR mice showed that asymmetrical partition or segregation of the mutated progenitors is central to the process of clonal purification. Similarly, using FACS analysis on single cell suspensions from colonic crypts isolated from patients with longstanding ulcerative colitis, considerable areas of colonic mucosa, exceeding 9 cm in length, were colonised by the same aneuploid stem cell of the same phenotype, suggesting that mutated clones in the human colon spread by the process of crypt fission.

Crypt fission events are likely to be under the control of intestinal stem cells: the three dimensional reconstruction by Wasan et al showed that adenomatous crypts can apparently arise from normal crypts in pre-existing adenomas. Such a process could play a role in segregating mutated stem cell clones from normal crypt epithelium, and also concurs with data showing that about 75% of FAP microadenomas are polyclonal. Similarly, Bjerknes et al studied the replication rate of mutant crypts (aberrant crypt foci) in FAP patients and controls (non-FAP colons) and found that both FAP crypts and mutant crypts in non-FAP patients replicate at least 1.1 times and seven times faster than normal crypts. Thus the main mode of growth of colonic adenomas in FAP patients and MIN mice is by the process of crypt fission. Hereditary colonic cancers account for about 15% of all human colorectal cancers and the majority of colonic carcinomas are sporadic in nature. It is commonly believed that the colorectal adenoma is the precursor lesion of most colorectal cancer. However, the histogenesis of human sporadic colorectal adenomas remains poorly understood.

Similarly, the histogenesis of human colorectal hyperplastic polyps was under described until the study by Araki et al who examined hyperplastic polyps by scanning electron microscopy and found that branching was observed in 20% of polyps. But the accuracy of quantitative assessment in their study was limited by the fragility of the isolated crypts and the high incidence of fracture of branched crypts during the procedure. Thus we used a microdissection based technique to examine the crypt fission process in both sporadic colorectal adenomas and hyperplastic polyps.
adenomas and hyperplastic polyps. This method is robust, reliable, and easily reproducible.

In normal colonic mucosa, proliferating cells are predominantly located in the lower two thirds of the crypts. In vitro labelling studies showed that the pattern of labelling in hyperplastic polyps is essentially similar to that of the normal mucosa. However, the pattern of labelling is different in adenomas, in which DNA synthesising cells are supposedly located mainly in the upper portion of the gland suggesting an upward migration of the proliferation zone. The classical ideas on the histogenesis of colonic adenomas are due to Dukes who proposed that tubular adenomas arise through proliferation and lateral expansion in the basal proliferative zone of the crypts. This can give similar techniques and in setting thresholds for scoring a cell as first is the difficulty in standardising immunohistochemical problems associated with the use of antibody techniques. The orientation of cells but the results generated are highly related antigens has the advantage of preserving the spatial zone of the crypts while villous adenomas arise from surface proliferation. However, others had shown that tubular adenomas arise in the superficial portion of the rectal crypts. Similarly, Lipkin showed that cell proliferation was prominent at the top of the rectal crypts, not only in the small adenomas of polyposis coli but also in the intervening morphologically normal rectal crypts, supporting an origin of adenomas from the upper crypt surface. However, some studies refute these claims. Based on the distribution of proliferating cells by proliferating cell nuclear antigen and Ki67 immunohistochemistry, and apoptotic cells denoted by terminal deoxynucleotidyl nick end labelling, Moss et al controversially proposed that the main direction of cell migration in adenomas is not towards the lumen but instead cells migrate towards the polyp base.

The use of immunohistochemical staining of cell cycle related antigens has the advantage of preserving the spatial orientation of cells but the results generated are highly dependent on prior treatment of the tissue and there are two problems associated with the use of antibody techniques. The first is the difficulty in standardising immunohistochemical techniques and in setting thresholds for scoring a cell as labelled or unlabelled. The second is related to the geometrical problem of scoring the sections, as the results may be confounded by concomitant changes in crypt size. An alternate method is to count mitotic figures or arrested metaphases in microdissected crypts. This can give similar results but is far quicker and is also more robust. The great advantage of such techniques is that results are expressed on a per crypt basis so that one does not need to count interphase cells. Furthermore, the geometric disadvantages associated with small cell crypt sections are avoided. Microdissection of crypts enables all the mitotic figures in the entire crypt to be scored, and all the crypts in a sample can be quantified. This is often more preferable to the several in vitro methods available for the assessment of biopsies which are labour intensive, require the immediate incubation of the tissue, and may not reflect the in vivo state. The microdissection method has been validated by study of human intestinal biopsy samples as well as in animal studies. Biopsies need little special treatment, apart from the use of Carnoy's solution for fixation. The tissue can then be stored in 70% alcohol for years. Crypt size and area can also be determined by the use of a drawing tube, as can the position of mitoses in the crypt. Combining all of the above factors, scoring mitoses in microdissection crypts appears to be the method of choice for the study of gastrointestinal proliferation.

Thus the aims of this study were: (1) to examine the morphology and proliferation characteristics of normal human colorectal mucosa, colorectal hyperplastic polyps, and sporadic colorectal adenomas by the microdissection method; (2) to examine the zonal distribution of mitoses in the three conditions; and (3) to determine whether crypt fission is the predominant proliferation abnormality in hyperplastic polyps and sporadic colorectal adenomas.

**METHODS**

**Specimens**

Fresh colonic polyps (adenomas and hyperplastic polyps) and normal colonic mucosa were obtained by endoscopic means during colonoscopy, after informed written consent. The colonoscope used was a Pentax EC-3801 or ES 3801 (Pentax, Hong Kong Ltd) with a 3.8 mm biopsy channel. Bowel preparation for colonoscopy consisted of a 45 ml phosphosoda solution followed by 2 litres of clear fluid. The majority of colono-scopies were performed by a single experienced endoscopist (BCYW). Patients with FAP, hereditary non-polyposis colorectal cancer, or any other form of hereditary colorectal cancer were excluded. The biopsy tissues were immediately fixed in Carnoy's fixative (ethanol:acetic acid:chloroform 6:1:3) for three hours and then stored in 70% alcohol before analysis. Part of the colonic polyps were fixed in formalin, sent for routine histological examination, and classified into hyperplastic polyps or adenomas. Polyp tissues or a definite pathological diagnosis were excluded. Eleven cases of adenomas (six cases of tubular adenomas (one with mild dysplasia and five with moderate dysplasia), four cases of tubulovillous adenomas (two with mild dysplasia, one with moderate dysplasia, and one with moderate to focal severe dysplasia), and one case of villous adenoma with moderate dysplasia) and five cases of hyperplastic polyps were studied. In 14 cases, non-involved colonic mucosa 10–15 cm away from the polyp lesions were also studied. Five cases of normal colonic mucosa from patients with normal colonoscopic findings were used as normal controls.

**Microdissected techniques for isolating crypts**

Individual crypts were microdissected using the method described by Goodlad and colleagues. The tissue was taken from its storage in 70% ethanol, hydrated (in 50% ethanol for 10 minutes followed by 25% ethanol for 10 minutes), and hydrolysed in 1 M HCl at 60°C for 10 minutes. It was then stained by Feulgen reaction for at least 45 minutes. The tissue was transferred onto a slide with 45% acetic acid and gently teased apart under a dissecting microscope at ×25 magnification. A coverslip was then placed over the wet tissue and tapped until the crypts began to separate. Microdissected crypts were examined under the compound microscope, and the outline of the crypts was traced using a Leitz drawing tube (Leica UK, Milton Keynes, UK). The number of mitoses in each gland or crypt was counted. The criteria used to define mitoses were strict and only distinct late prophase, metaphases, anaphases, and telophases were scored. Crypt size (area) was obtained by scanning the tracings with a flatbed scanner connected to an Apple Macintosh computer and the area was measured using the program Image (NIH public domain).

**Measurement of crypt fission index, cell proliferation, and distribution of mitoses**

Microdissected Feulgen stained crypts were counted under ×40 magnification. Mitoses were counted in at least 20 microdissected crypts per biopsy using a compound microscope at 160 magnification. By focusing up and down through the width of the crypt, a total mitotic count for each crypt was obtained. Crypts were then divided into five zones (zones 1–5, from base to top) by projecting a grid into the microscope field or view, and the distribution of mitoses per zone was recorded. Crypts with complex branching were orientated correctly before calculation of zonal mitotic count. The mean crypt area of all crypts isolated was plotted against the crypt fission index to determine the relationship between the two variables.

**Statistical analysis**

All results are presented as group means (SEM). The statistics used included the two sided t test using the SPSS 7.5 (Inc.,
The association between crypt fission index and crypt area was measured by calculating the Pearson correlation coefficient. A p value $< 0.05$ was considered to be statistically significant.

**RESULTS**

**Morphology of colonic crypts from uninvolved mucosa and normal controls**

Normal colonic crypts were cylindrical in shape and mitotic figures were mainly concentrated at the base of the crypt—that is, zone 1 (fig 1A, B). Crypt fission was rare and began with basal bifurcation at the base of the gland (fig 1B). No asymmetrical or atypical types of branching were observed.

**Morphology of adenomas and hyperplastic polyps**

The mean area of colonic crypts isolated from adenomas was significantly larger compared with uninvolved mucosae and normal controls ($p<0.001$) (fig 2B), and significantly larger than hyperplastic polyps ($p=0.001$). The shapes of these crypts were irregular and asymmetrical. Multiple branching and budding were observed in a single crypt, suggesting an accelerated rate of crypt fission (fig 1E–G). Fission was mostly asymmetrical as opposed to fission in normal colonic mucosae (fig 1B). Superficial budding was commonly seen in contrast with normal controls in which all budding was basal. Crypts isolated from hyperplastic polyps were significantly larger than crypts isolated from non-involved mucosae ($p<0.001$) (fig 2B). Mean crypt area of the hyperplastic polyps was also larger than that of normal controls but this difference was not statistically significant. Crypts from hyperplastic polyps appeared less irregular in shape but fission events were still very common. Crypt fission was more symmetrical in these crypts compared with adenomatous crypts and usually started from the crypt base (basal budding). Multiple fission events...
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Figure 2 Mitoses per crypt (A), mean crypt area (B), mitoses per area (C), and crypt fission index (CFI, proportion of crypts in fission) (D) for adenomas, hyperplastic polyps, non-involved mucosae, and normal controls.

Figure 3 Relationship between crypt area and crypt fission index ($r=0.83$, $p<0.001$).

Figure 4 Percentage mitoses per crypt zone for adenomas, hyperplastic polyps, non-involved mucosae, and normal controls. **$p<0.001$** compared with controls and non-involved mucosae.

were observed (fig 1C, D) in some of the crypts isolated. Interestingly, we also observed the presence of some very elongated crypts adjacent to adenomas (fig 1H).

**Proliferation characteristics of adenomas and hyperplastic polyps**

The mean crypts counted for mitotic scores were 26.5 (1.6).Crypt mitotic scores were significantly greater in adenomas ($p<0.001$) and hyperplastic polyps ($p=0.03$) compared with non-involved mucosae (fig 2A). Mitoses per crypt for adenomas were also significantly greater than for normal controls ($p=0.02$) (fig 2A). Mean mitoses per crypt for hyperplastic polyps was greater than that for normal controls but this was not statistically significant. When the effect of size was taken into account—that is, mitoses per area—there were no differences between adenomas, hyperplastic polyps, and normal controls (fig 2C).

Mean crypts counted for crypt fission index per control and per adenoma were 292 (SEM 39; range 100–828) and 91 (SEM 17; range 21–202), respectively. The crypt fission index (proportion of crypts in fission) for adenomas was significantly greater than that for hyperplastic polyps ($p=0.049$), non-involved mucosae ($p<0.001$), and normal controls ($p=0.003$) (fig 2D). The crypt fission index for hyperplastic polyps was significantly greater than that for non-involved mucosae ($p<0.001$) and normal controls ($p=0.01$) (fig 2D). The correlation between crypt fission index and crypt area was 0.83 ($p<0.001$) (fig 3). Crypt mitotic scores were higher in adenomas with moderate dysplasia compared with adenomas with mild dysplasia (30.6 ± 11.1; $p=0.03$) but mean crypt area, mitoses per area, and crypt fission index were similar.

**Zonal distribution of mitoses (fig 4)**

The zonal distribution of mitoses in adenomas, hyperplastic polyps, non-involved mucosae, and normal controls is shown in fig 4. In normal colonic mucosae, the mitotic figures were mainly located in zones 1 and 2—that is, the base of the crypts. In hyperplastic polyps, mitotic figures were mainly concentrated in zones 1–3, but in adenomas mitoses were almost evenly distributed between the five zones. The mean percentage zonal mitoses of zones 1 and 2—that is, the base of the crypts—were significantly smaller in adenomas (zone 1 = 16%; zone 2 = 22%) compared with non-involved mucosae (zone 1 = 34%, zone 2 = 43%; $p<0.001$ for both zones) and normal controls (zone 1 = 29%, zone 2 = 47%; $p=0.001$ for both zones). On the other hand, mean percentage zonal mitoses of zones 4 and 5—that is, the top of the crypts—were significantly greater in adenomas (zone 4 = 23%, zone 5 = 17%) compared with non-involved mucosae (zone 4 = 4%, zone 5 = 1%; $p<0.001$ for both zones) and normal controls (zone 4 = 2%,
mitoses were almost evenly distributed between the five colleagues' findings support the mouse studies of Totafurno and larger in both hyperplastic polyps and adenomas (fig 2D). Our crypts (fig 2B). The crypt fission index was also significantly larger were significantly larger than those from normal colonic adenomas. Here we have reported a microdissection study in which colonic hyperplastic polyps and adenomas grew by the process of crypt fission and was the major proliferation abnormality.

Colonic crypts isolated from both hyperplastic and adenomas were significantly larger than those from normal colonic crypts (fig 2B). The crypt fission index was also significantly larger in both hyperplastic polyps and adenomas (fig 2D). Our findings support the mouse studies of Totafurno and colleagues in which dividing crypts were generally larger (fig 3) and crypt size was therefore believed to be an important factor in initiating crypt fission. It has been proposed by Löefl et al that crypts will divide if the number of stem cells exceeds a certain threshold, called the S th. Thus it is reasonable to assume that a larger crypt will contain a greater number of stem cells and hence a greater rate of crypt fission compared with normal sized crypts. Furthermore, we also found that the proliferation characteristics of non-involved mucosae adjacent to adenomas were similar to those of normal controls. Thus hyperproliferation appears to be localised on colonic adenomas rather than a generalised and diffuse hyperproliferation of the whole colon, as previously suggested. In conclusion, sporadic human colonic adenomas and hyperplastic polyps grow by the process of crypt fission. Expansion of the proliferative compartment was demonstrated in crypts from adenomas, consistent with deregulation of cell cycle control.

Note added in proof
A recent article by Shih et al (Proc Natl Acad Sci USA 2001;98:2640–5) has suggested that adenomas grow by a "top-down morphogenesis," taking origin from mutant stem cells on the surface of the colonic mucosa, and the adenomatous epithelium grows downwards to occupy adjacent crypts. This concept diverges markedly from the present study, which indicates that adenomas grow, in the main, by crypt fission. The debate also extends into how clonal patches of dysplasia branch in the colon in ulcerative colitis—"top down" by lateral migration or "bottom up" by crypt fission, or both? Which mode of morphogenesis prevails will have considerable implications for stem cell biology in the gut.

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