Colon Preneoplastic Lesions in Animal Models

Masumi Suzui1*, Takamitsu Morioka2, and Naoki Yoshimi3

1 Department of Molecular Toxicology, Graduate School of Medical Sciences and Medical School, Nagoya City University, 1 Kawasumi, Mizuho-ku, Mizuho-cho, Nagoya 467-8601, Japan
2 Radiation Effect Accumulation and Prevention Project, Fukushima Project Headquarters and Radiobiology for Children’s Health Program, Research Center for Radiation Protection, National Institute of Radiological Sciences, 4-9-1 Anagawa, Inage-ku, Chiba 263-8555, Japan
3 Department of Pathology and Oncology, Graduate School of Medicine and Faculty of Medicine, University of the Ryukyus Faculty of Medicine, 207 Uehara, Nishihara-cho, Okinawa 903-0215, Japan

Abstract: The animal model is a powerful and fundamental tool in the field of biochemical research including toxicology, carcinogenesis, cancer therapeutics and prevention. In the carcinogenesis animal model system, numerous examples of preneoplastic lesions have been isolated and investigated from various perspectives. This may indicate that several options of endpoints to evaluate carcinogenesis effect or therapeutic outcome are presently available; however, classification of preneoplastic lesions has become complicated. For instance, these lesions include aberrant crypt foci (ACF), dysplastic ACF, flat ACF, β-catenin accumulated crypts, and mucin-depleted foci. These lesions have been induced by commonly used chemical carcinogens such as azoxymethane (AOM), 1,2-dimethylhydrazine (DMH), methylnitrosourea (MUN), or 2-amino-1-methyl-6-phenylimidazo[4,5-b]pyridine (PhIP). Investigators can choose any procedures or methods to examine colonic preneoplastic lesions according to their interests and the objectives of their experiments. Based on topographical, histopathological, and biological features of colon cancer preneoplastic lesions in the animal model, we summarize and discuss the character and implications of these lesions. (DOI: 10.1293/tox.2013-0028; J Toxicol Pathol 2013; 26: 335–341)

Key words: preneoplastic lesion, colon carcinogenesis, animal model, topographic view

Aberrant Crypt Foci (ACF)

Bird1 first reported in 1987 that when C57BL/6J mice were treated with azoxymethane (AOM), aberrant dysplastic crypts appeared in the colonic mucosa. After fixation with 10% buffered formalin and staining with methylene blue, these crypts were easily visualized in the topographic view of the colonic mucosa using a x4 objective (Fig. 1A). These lesions were referred to as aberrant crypts (AC) or aberrant crypt foci (ACF) in the colon of both animals and humans2–4. ACF were cryptic lesions distinguished by their increased size, thicker epithelial lining, and increased pericryptic zone1. ACF have only been seen in the colon of carcinogen-treated mice and rats. They have not been seen in the colon treated with a noncarcinogen or in untreated animals2,3. After carcinogen treatment, they appeared as early as within 2 weeks and persisted until the experimental termination of animals (16 weeks); histological changes from mild atypia to dysplasia5 were also revealed. Two heterocyclic amines, 2-amino-3-methylimidazo[4,5-f]quinoline (IQ) and 2-amino-1-methyl-6-phenylimidazo[4,5-b]pyridine (PhIP), were shown to be able to induce ACF in the colon, respectively, after 4 and 10 weeks of exposure2. The number of ACF increased significantly over time, and small-sized ACF were predominant at all time points5. In histological slides, the large ACF exhibits dysplasia and thus can be termed a microadenoma2.

ACF are also induced in the colonic mucosa of rats or mice treated with carcinogens such as AOM, methylazoxymethanol (MAM) acetate, 1,2-dimethylhydrazine (DMH), methylnitrosourea (MNU), PhIP, IQ, 2-amino-3,8-dimethylimidazo[4,5-f]quinoline (MeIQ) and 2-amino-6-methylidipyrido[1,2-a:3',2'-d]imidazole (Glu-P-1)6–11. In our previous experiments7,12, F344 rats were subcutaneously (sc) injected with AOM (20 mg/kg body weight) twice. Five weeks after the beginning of the experiment, 93–139 ACF per colon occurred. When F344 rats were treated with AOM (15 mg/kg body weight, sc injection) 3 times, 240 ACF/colon occurred at 11 weeks after the beginning of the experiment13. When F344 rats were treated with DMH (40 mg/kg body weight, sc injection) 3 times, 240 ACF/colon occurred at 11 weeks after the beginning of the experiment14,15. When F344 rats were treated with DMH (40 mg/kg body weight, sc injection) twice, 175–200 ACF/colon were induced at 5 or 8 weeks after the beginning of the experiment13. These ACF usually contained 1–3 or more crypts per focus. The diameter of an aberrant crypt measured at least 3 to 4 times larger than that of a normal crypt...
in mice and up to 1.5 times larger than a normal crypt in humans. Pretlow et al. reported that ACF were at least 3 times larger in diameter than normal crypts, and most ACF had lumina that were oval or slit shaped rather than circular. ACF range in size and have from 1 to 412 aberrant crypts per focus. The size in the topographic view and the histologically dysplastic character of ACF are critical factors when we distinguish ACF as preneoplastic lesions. We consider that large ACF consisting of more than 10–20 crypts and manifesting dysplasia could be termed a microadenoma. In mouse models, for instance, B57BL/6J and CF1 mice were given a single intraperitoneal (ip) injection of AOM (5 mg/kg body weight), and 4 weeks later, mice developed 2.6 and 3 ACF per colon, respectively. BALB/c mice were ip injected with AOM (10 mg/kg body weight) twice, and 14 ACF were induced 4 weeks after the injection. In C57BL/6J-Min+/(Min) and C57BL/6J-+/+(wild type) mice, PHIP was ip injected 4 times. Ten weeks after the injection, male mice developed 3 and 0 ACF, respectively, and female mice developed 1.9 and 0.2 ACF in their colons. These findings indicate that duration of the experimental period, strain of animals, method of administration of carcinogens, and nature of carcinogen used as an initiator, may affect the number of ACF in the colonic mucosa.

In terms of the distribution of ACF, McLellan et al. demonstrated that AOM-treated CF1 mice developed ACF, 67% of which were in the rectal segment, 29% of which were in the middle segment and 4% of which were in the cecal segment. ACF were seen mainly in the rectal and middle segments when the animals were treated with DMH, NMU, MeIQ, or Glu-P1. Most ACF were found in the middle and distal colon in F344 rats treated with AOM. In contrast, Hata et al. demonstrated that ACF were frequently found in the proximal colon (cecal segment of the colon) when AKR/J and SWR/J mice were treated with AOM. The carcinogen IQ also induced ACF primarily in the middle and cecal segments of the colon. Colon tumors induced by AOM were primarily found in distal colon rather than in proximal colon in the rat and mouse models, indicating that the correlation between ACF formation and carcinogenesis is not necessarily straightforward. This is presumably because of the heterogeneous nature of ACF. Also, experimental protocol and species used may affect the difference in distribution of ACF.

The shape of the lumen of the ACF is related to the histology of the ACF. Histological criteria of rat/mouse ACF have been described by several investigators. Accordingly, ACF may be classified into the following 3 categories. In brief, these are (1) non-dysplastic foci, which exhibit hypercellularity of uniform or normal looking goblet cells with basal-oriented nuclei and apical localization of mucus; (2) mild to moderate dysplastic foci, which exhibit hypercellularity of cells with elongated nuclei and focal nuclear stratification; and (3) moderate to severe dysplastic foci, which exhibit hypercellularity of elongated cells with abundant basophilic cytoplasm. These foci display enlarged and vilicated nuclei, sometimes with prominent nucleoli.

### Dysplastic ACF

The dysplastic nature of ACF was described by McLellan and Bird. In a hematoxylin-eosin (HE) stained transverse section, ACF exhibited a focal appearance and mild cellular atypia, and dysplasia was observed in the large focus. Bird and Pretlow mentioned that use of the term dysplastic crypt foci to describe abnormal crypts is valid only if the investigators examined histologically all of the methylene blue-identified lesions and found dysplasia in all of them. Ochiai et al. described two distinct types of ACF in the PHP-induced rat model. One was dysplastic ACF, and the other was nondysplastic ACF. In their reports, dysplastic ACF are histologically characterized by distortion of the crypt structure, a decrease in goblet cell number, existence of nuclear stratification, and enlarged nuclei. Nondysplastic ACF indicated the hyperplastic change in crypts. One-fourth of PHP-induced ACF were dysplastic ACF, and the remaining ACF were nondysplastic ACF. Two-week dietary administration of 400 ppm PHP was repeated three times with a 4-week interval. The average number of dysplastic ACF was up to 0.8 per colon, and they were larger in size than nondysplastic ACF after 32 weeks of experimentation. In the dysplastic ACF, cytoplasmic β-catenin protein accumulation and β-catenin gene mutation were found. The mutations were A→G (Asp→Gly), G→T (Gly→Val), and C→T (His→Tyr). By a staining method that uses 70% methanol followed by 0.2% methylene blue staining, dysplastic ACF can be topographically contrasted with nondysplastic ACF on the colonic mucosa and identified without performing histological examination. The average number of dysplastic ACF/colon was 2.0–3.2 in F344 rats treated with PHP (400 ppm in diet), MeIQ (300 ppm in diet), and IQ (300 ppm in diet). Two-week dietary administration of PHP, MeIQ, or IQ was repeated three times with a 4-week interval. Other investigators have also described dysplastic ACF. Thorup found that a correlation between degree of dysplasia and crypt multiplicity, indicating that chemically induced ACF can increase in crypt multiplicity over time and progress into a tumor and that hyperplastic human ACF can also develop into adenomatous ACF, as reported elsewhere. However, this view disagrees with that of other studies demonstrating that the degree of dysplasia is not necessarily related to the crypt multiplicity.

### Flat ACF

Paulsen et al. examined unsectioned methylene blue-stained colon tissues obtained from male F344 rats treated with AOM (sc injection ×2 times, 15 mg/kg body weight), and found two types of early lesions. One was classic elevated ACF, and the other was flat ACF. Classical ACF were seen as enlarged crypts that were elevated from the surrounding epithelium and had elongated luminal openings. However, Paulsen et al. described flat ACF as structures that were not elevated. The bright blue appearance and compressed pit pattern of flat ACF were used as criteria for identification.
Flat ACF were characterized by enlarged or small crypts that were not elevated from the epithelium and had round or elongated luminal openings. The investigators also described histological findings of flat ACF with severe dysplasia. In immunohistochemical analysis, classic elevated ACF did not show (0 of 99) cytoplasmic/nuclear expression of the β-catenin protein. In contrast, all flat ACF (8 of 8) displayed cytoplasmic/nuclear expression of the β-catenin protein. The number of classic elevated ACF decreased along with time. Their crypt multiplicity increased during the time period. The number of flat ACF decreased along time, and that of tumors increased correspondingly. The numbers of flat ACF plus tumors were virtually constant. In view of these findings, Paulsen et al. concluded that flat ACF display a continuous development from early stages into a tumor.

β-Catenin Accumulated Crypts (BCAC)

In a previous study, we found that focal lesions that display accumulation of the β-catenin protein predispose to carcinogenic-induced colon carcinogenesis. We named these lesions β-catenin-accumulated crypts (BCAC) (Fig. 1B). F344 rats were treated with AOM (sc injection ×3 times, 15 mg/kg body weight), and a complete autopsy was performed at 10 weeks after the first AOM treatment. In the topographical view in which colon tissues were stained with methylene blue, we found distinct populations of altered crypts named histologically altered crypts with macroscopically normal-like appearance (HACN) among the tissue samples. In HACN, which are equivalent to BCAC, the β-catenin gene was frequently mutated in 10 of 15 samples (67%), and the cytoplasmic β-catenin protein was accumulated in 13 of 15 samples (86%). Among these lesions, there were 28A→T (Gln→His), 29C→G (Ser→Cys), 30T→C (Tyr→His), 32G→A (Asp→Asn), 34G→A (Gly→Glu), 34G→T (Gly→stop), and 41A→T (Thr→Ile) mutations. Because the lesion in which the β-catenin protein accumulated was considered to be valid in AOM-treated rat colonic mucosa, a time course study was done to examine the status of the protein accumulation, the number of crypts/lesion, and the diameter of the crypts. Both the number of crypts/lesion and the diameter of the β-catenin accumulated crypts that were identified with immunohistochemical analysis significantly increased with the time course of BCAC. The number of BCAC induced by AOM in AKR/J and SWR/J mice varied by 3–12 per cm², and multiplicity was about 3–4 in both strains. Histological abnormality of the crypts and cell proliferation also significantly increased when compared with those of ACF, indicating that BCAC are neoplastic lesions in AOM-induced colon carcinogenesis.

Mucin-depleted Foci (MDF)

Caderny et al. identified specific lesions in the colon of rats treated with AOM. When unsectioned colon tissues were stained with high-iron diamine-Alcian blue (HID-AB), foci of crypts with scarce or absent mucins were seen, and such lesions were first defined as mucin-depleted foci (MDF) (Fig. 1C). In that study, male F344 rats received sc injection of AOM (15 mg/kg body weight) twice. The rats developed approximately 4 and 8 MDF/colon at 7 and 15 weeks, respectively, after the start of the experiment, while 271–289 ACF/colon developed during the same period. Mutations in β-catenin, Apc, and K-ras genes and cytoplasmic β-catenin expression were found in MDF induced by DMH. Among these, β-catenin gene mutations included 32G→A (Asp→Asn), 33C→T (Ser→Phe), 33C→T (Ser→Phe), and 34C→T (Thr→Ile). In DMH studies, MDF exhibit dysplastic features, and the induction rate of MDF is dose dependent. Also, MDF increase in size with time. To examine the multiplicity and distribution of ACF, MDF, and tumors, six-week-old F344 rats were treated with DMH (40 mg/kg body weight sc injection twice a week) followed by 1% dextran sodium sulfate in drinking water. At ten and fourteen weeks after the start of the experiment, animals were euthanized. ACF were mainly found in the middle portion of the colon (Fig. 2A). MDF and tumors occurred more in the distal portion than in the proximal portion (Fig. 2B and C). These results were in accordance with those in the report of Femia and Caderny. They found that DMH-induced MDF and tumors were mainly found in the distal portion of the colon, while “classical” ACF were found more predominantly in the middle portion of the colon. Also, Femia et al. mentioned that with regard to the ability of ACF/MDF as a biomarker predicting the carcinogenesis status, the heterogeneous nature of each lesion may be related.

Only a limited number of findings on MDF are currently available; based on those that are available, Femia and Caderny conclude that MDF are premalignant lesions for colon carcinogenesis and a promising biomarker for study of the effect of chemopreventive agents in colon carcinogenesis. MDF may provide a reliable option as biomarkers for colon carcinogenesis, and it is thought that production or deletion of mucin or both plays some roles in the development of colon tumors. To reiterate, MDF may have both morphological and biochemical aspects as a biomarker. To identify MDF, we demonstrated a simple staining method using 1% Alcian blue (pH 2.5) solution instead of the original HID-AB staining method. In this study, male F344 received sc injections of DMH (40 mg/kg body weight) twice, and the rats developed 19 MDF/colon and 150 ACF/colon at 8 weeks after the start of the experiment. By comparing exact locations of MDF and BCAC on the face-up mucosal samples and by conducting Alcian blue/HE/immunohistochemical staining, we found that MDF are practically identical to BCAC and useful as an early biomarker in rat colon carcinogenesis. In human specimens obtained from patients with colorectal carcinoma (CRC) and familial adenomatous polyposis (FAP), MDF were also identified. The mean numbers of crypts/MDF were 60 and 33 in specimens of patients with CRC and FAP, respectively. In a CRC case, the histological diagnosis of MDF was microadenoma with moderate grade dysplasia, while in cases of FAP, the diagnosis was microadenoma with low-grade dysplasia.
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Our study, our group examined human CRC cases and found MDF on the colonic mucosa. The lesion was histologically classified into two categories: flat MDF and protruded MDF. The former lesion did not show nuclear stratification or loss of polarity, but showed Paneth cell metaplasia and decrease/loss of goblet cells, indicative of low-grade dysplasia. Protruded MDF displayed the features of both ACF and MDF, also corresponding to low-grade dysplasia. A topographic view of human MDF is shown in Fig. 3.

Conclusions

This review summarizes topographical, histopathological, and biological features of preneoplastic lesions that have been described in colon carcinogenesis models of the rodent (Table 1). The early lesion has been identified and documented as a preneoplastic lesion in the carcinogenesis process. However, the fact that even the verified lesions appear to contain neoplastic lesions such as a microadenoma indicates the need for further investigation. This may be due to complicated categories or classifications of preneoplastic lesions. Considering the 3R principles (which commonly consist of replacement of methods with no animal use, reduction of the number of test animals, and refinement of methods that minimize the suffering of test animals), a short-term experiment in which an early preneoplastic lesion occurs and can be used as a biomarker should be used to examine toxicity and/or carcinogenicity of test compounds in a specific organ site. In this context, investigators can choose any procedures or methods to examine colonic preneoplastic lesions according to their interests and the objectives of their experiments.
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Table 1. Summary of Characters of Prenoeplastic Lesions in the Animal Model

| Practical name of the lesion | Ref. no. of the current article | Staining method | Features in topographic and/or histological views |
|-----------------------------|---------------------------------|-----------------|--------------------------------------------------|
| ACF                         | 1, 2, 11, 26, 32                | Methene blue staining | In topographic view, increased cryptal size, thicker epithelial lining, and increased pericryptical zone. |
| Dysplastic ACF              | 26, 32, 34–36                   | HE staining      | Histologically characterized by distortion of the crypt structure, a decrease in goblet cell number, nuclear stratification, and enlarged nuclei. |
| Flat ACF                    | 42                             | Methene blue staining | Characterized by bright blue staining, enlarged or small crypts not elevated from the epithelium and round or elongated luminal openings. Because the flat ACF were not observed as elevated structures, their bright blue appearance and compressed pit pattern were used for identification. |
| BCAC                        | 24, 43, 44                      | Immunohistochemical staining | Accumulation of cytoplasmic β-catenin protein. Crypts of BCAC do not display prominent epithelial cells in a topographic view. |
| MDF                         | 11, 45–50                      | Alcian blue (HID-AB) staining 1% AB, pH2.5 | When colon tissues were stained with HID-AB, foci of crypts with scarce or absent mucin were defined as MDF. MDF can be stained with 1% AB solution. |

ACF, aberrant crypt foci; BCAC, β-catenin accumulated crypts; MDF, mucin-depleted foci.
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