MUCOUS SECRETION IN RAT COLONIC MUCOSA DURING CARCINOGENESIS INDUCED BY DIMETHYLDRAZINE. A MORPHOLOGICAL AND HISTOCHEMICAL STUDY

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Summary.—Our previous studies, in specimens of large intestine resected for carcinoma, have shown abnormal patterns of mucous secretion in areas of apparently "normal" mucosa, where goblet cells produce mainly sialomucins as compared with the true normal colonic mucosa in which sulphomucins predominate. In the present work, large bowel cancer was induced in rats by the administration of 1,2-dimethylhydrazine-2HCl (DMH). We attempted to study the sequential histological and secretory abnormalities which developed in the colonic epithelium during carcinogenesis, and to correlate these changes with those described above in the human. The microscopical and histological lesions observed in the colonic mucosa of DMH treated rats confirmed the findings of other authors and resembled the human colorectal cancer. The earliest changes detected were small foci of hyperplasia accompanied from the 6th week onwards by several foci of dysplasia. Carcinoma in situ appeared at the 15th week and finally invasive carcinoma developed from the 19th week onwards. Changes in the type of mucous secretion, with predominance of sialomucins, were observed in the majority of the areas showing mild to moderate dysplasia whilst the surrounding normal epithelium produced sulphated material. Mucous depletion was a common feature in areas of severe dysplasia and carcinoma.

These findings correlated well with the similar variations in the mucin composition observed in human colonic mucosa in carcinoma and further supported our previous hypothesis that mucin changes characterized by an increase in sialomucins might reflect early malignant transformation. If this hypothesis proved to be correct, the use of a simple method for the identification of mucins in large bowel biopsies would be of great help in detecting early malignancy.

In man, the non-neoplastic mucosa adjacent to carcinoma of the large intestine ("transitional" mucosa as we provisionally termed it) is quite often histochemically and ultrastructurally abnormal (Filipe, 1969, 1971a; Dawson and Filipe, 1975). It may be normal in haematoxylin–eosin stained sections or may show longer crypts with branching and larger goblet cells distended with mucus (Filipe and Branfoot, 1974). The histochemical changes in the composition of the goblet cell mucin consist of an increase in sialomucins as compared with the normal mucosa in which sulphomucins predominate. The higher level of sialic acids is accompanied by a rise in neuraminidase-sensitive sialic acids (Filipe and Cooke, 1974), a feature described in the human foetal gut mucin (Lev and Orlic, 1974).

From these previous findings, it is possible that the mucin changes described are either secondary to the tumour growth or are a primary cellular response to unknown stimuli (e.g. carcinogens) and thus indicative of early malignant transformation.

In an attempt to clarify these points, we studied in detail the mucosa of the
entire length of large intestinal specimens resected for carcinoma (Filipe and Branco-foot, 1974). We observed that mucin changes with a predominance of sialomucins were present not only in the apparently normal mucosa adjacent to the carcinoma but also in patches of normal mucosa at various distances from the tumour. These observations suggest that the modifications in the secretory product may represent an early feature of malignancy.

To elucidate this problem further, it is of the utmost importance to use an animal model for colorectal carcinoma. The present work reports our findings in rats in which neoplasms of the large intestine were selectively induced by 1,2-dimethylhydrazine-2HCl (Druckrey et al., 1967; Druckrey, 1970).

Our purpose was to investigate the morphological and mucin changes occurring in the rat colonic epithelium, with particular emphasis on the early stages of carcinogenesis, and also to determine whether changes in the mucin composition in the rat are similar to those described in the human colonic mucosa in cases with carcinoma.

MATERIALS AND METHODS

Animal experiments.—75 white Wistar female rats (A. Tuck & Son Ltd, The Nursery, Rayleigh, Essex, England) weighing approximately 140 g at the beginning of the experimental period were used in this study. The animals were fed with standard diet 41B and water ad libitum. The “test” rats (50) were injected subcutaneously with a weekly dose of 20 mg/kg body weight, expressed as the base of 1,2-dimethylhydrazine-2HCl (DMH). The DMH solution was prepared according to Wiebecke et al. (1969) modified by P. Magee (personal communication) as follows: the 1,2-dimethylhydrazine-2HCl (Ralph Emanuel Ltd, Wembley, Middlesex, England) was freshly dissolved in 0.9% NaCl containing 1.5% EDTA to give a 0.35% solution with respect to the base and the pH was adjusted to 6.5 with 1% NaOH. The control animals (25) were given a weekly subcutaneous injection of 0.9% NaCl containing 1.5% EDTA. The experiment lasted for 29 weeks.

Preparation of tissues.—Groups of 3 rats (2 tests and 1 control) were killed each week from the start of the experiment. The whole of the large intestine was removed, opened, pinned down onto cork and fixed in 10% neutral formol-calcium. After fixation, the entire specimen was divided into lengths and coiled up into “swiss-rolls” (Fig. 1). Blocks were also taken when the presence of large neoplasms made the use of “swiss-rolls” impractical.

All tissue was processed routinely and embedded in paraffin. Not less than 4 serial sections were cut at 5 μm thickness. Blocks containing tumours were sectioned at 3 or more different levels and serial sections were taken, as above, from each level. The sections were stained with haematoxylin and eosin (H. and E.) and by the following histochemical techniques for the identification of mucins: periodic acid-Schiff (PAS) (Pearse, 1968), high-iron-diamine (HID) and high-iron-diamine-Alcian blue (HID-AB) for the visualization of both sialo- and sulphated mucins (Spicer, 1965; Gad and Sylvén, 1969; Sorvari, 1972).

Mapping of morphological changes and mucin distribution.—The images of each of the H. and E. and HID-AB stained serial sections were projected onto paper and tracings were made of the mucosal outlines (Fig. 2). Areas showing alterations in the morphology and in the mucin composition were recorded on the graphs so that we could obtain an accurate chart correlating the histological features and the patterns of mucous secretion (Fig. 1, 2).

RESULTS

During all stages of the experimental period in which the rat colonic mucosa is subjected to the effect of the carcinogen, a wide spectrum of histological and histochemical (mucous secretion) modifications occur. The lesions are more severe and extensive in the distal colon and rectum than in the proximal colon and caecum.

Macroscopy

Caecum, proximal and distal colon can easily be distinguished by the different arrangement of the mucosal folds. In the caecum the folds are prominent and
Fig. 1.—DMH treated rat. Distal colon coiled up in a "swiss roll" (arrows indicate foci of dysplasia and carcinoma in situ). H. and E.

Fig. 2.—Tracing of mucosal outlines and mucin distribution of HID-AB stained serial section to Fig. 1. Secreting glands are indicated as circles (○) for sulphomucins and (●) for sialomucins. The latter are present in the areas showing dysplasia in the H. and E. section. Areas of carcinoma in situ (arrows) show mucus depletion.
### Table I.—Rat left colon: Distribution of Macroscopical, Histological and Mucin Changes during Carcinogenesis

#### Number of Foci

| Duration of Experiment (weeks) | Hyperplasia | Dysplasia | Carcinoma in situ | Carcinoma invasive | Macroscopical lesions |
|-------------------------------|-------------|-----------|-------------------|-------------------|----------------------|
| 0-1                           | ●           | ●         |                   |                   |                      |
| 1-2                           | ●           | ●         |                   |                   |                      |
| 1-3                           | ●           | ●         |                   |                   |                      |

Each column represents one DMH treated rat. Each symbol = one focal lesion, secreting either sulphomucins (●), or sialomucins (○) or both (●). △ = Mucous secretion absent or scanty (△ sialo- or sulphomucins △). □ = macroscopical lesions.

* = Invasive carcinoma also present in the right colon.
** = Generalized dysplasia.
*** = Superficial dysplasia.
at varying angles whilst in the proximal colon they display a "herring-bone" pattern. In the distal segment the mucosa is often smooth with the folds running longitudinally.

In the controls no macroscopical changes were observed in the mucosa, during the 29 weeks duration of the experiment.

In the tests no recognizable lesions are detected up to 12 weeks (Table I). At the 13th week one of the tests showed 2 small excrescences in the transitory distal-proximal colon (at 15 and 15.5 cm from the anus respectively). With light microscopy they represent areas of dysplasia. Other small mucosal protrusions are noted in the distal colon at the 15th week, one of them being histologically an area of carcinoma in situ.

The number and size of the macroscopical lesions increase with longer exposure to DMH (Table I) and from the 19th week onwards all rats develop neoplasms either as plaques or as sessile polyps measuring between 0.3 and 0.8 mm in diameter.

Of the 54 macroscopically recognizable lesions found between the 19th and 28th weeks of the experiment, 29 are invasive carcinoma, out of which 79% are located in the distal colon.

**Histology (Table I)**

**Controls.**—The distal colon reveals similar histological features to the human colorectal mucosa, with straight crypts lined by mucus secreting goblet cells (Fig. 3). The goblet cells predominate in the ⅔ upper crypt and are interspersed

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**Fig. 3.**—Control rat. Distal colon. HID-AB staining to show the predominance of dark coloured sulphomucins. × 600.
by absorptive cells. In the proximal colon, the crypts are branched and the goblet cells and absorptive cells are present in the $\frac{2}{3}$ or $\frac{1}{3}$ upper crypt, whilst the lower compartment is occupied by a further type of epithelial cell. This third type of cell is globular, with a vast pale cytoplasm in H. and E., and producing mucus which differs histochemically from the secretory product of the goblet cells (Fig. 4) (see Histochemistry section and Table II).

In this group of animals, no histological abnormalities other than the occasional mild oedema in the lamina propria are demonstrated.

Tests.—The morphological lesions observed, involving 1, 2 or more glandular tubules, may be grouped into 4 categories according to the degree of histological dedifferentiation (Muto and Morson, 1975) and defined as follows:

(1) Epithelial hyperplasia: Where the crypts are longer, the lumen may be dilated and the goblet cells are more numerous, taller, and distended with mucus. The nuclei are aligned regularly on the basement membrane.

(2) Dysplasia is graded 0–2. In grade 0 (Fig. 5) the crypts are not usually longer but may be dilated, the goblet cells are globular and between them the

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**Fig. 4.—Control rat. Proximal colon.** HID-AB Technique to show the predominance of sialomucins (grey) in the $\frac{2}{3}$ lower crypt; a few goblet cells containing sulphated material (black) may be seen in the upper crypt. The HID-AB method stains sialomucins blue (grey), whilst sulphomucins are stained brown-black (black). × 600.
TABLE II.—Histochemical Characteristics of the Mucins in the Various Segments of the Rat Large Intestine

|                  | Distal colon | Proximal colon | Caecum     |
|------------------|--------------|----------------|------------|
|                  | PAS     | HID-AB         | PAS     | HID-AB         | PAS     | HID-AB         |
| 1/2 upper crypt  | +++    | Brown          | +/+++  | Blue/brown     | +++    | Brown          |
| Middle crypt     | +++    | Brown          | +/+++  | Blue           | +++    | Brown          |
| 1/2 lower crypt  | +++    | Brown          | +/+++  | Blue           | +      | Brown/blue     |

absorptive cells have a more eosinophilic ground-glass cytoplasm. These features are more commonly noted in the upper half of the crypt. In grade 1 (Fig. 6 ABC) there is a distortion of the crypts' contours, hyperchromatic nuclei, moderate pseudostratification but with the preservation of nuclear polarity, and a slight reduction in the number of goblet cells. Mitotic figures are not present.

As it is not rare to find two grades of atypism in the same focus of dysplasia, or even in the same crypt, we will group them together under grade 0–1 (Fig. 5.—DMH treated rat. Grade 0 dysplasia in a single gland. The HID-AB method shows that the atypical gland (arrow) produces sialomucins (grey) compared with the surrounding glands secreting sulphomucins (black). ×600.
7A, B). Grade 1 (Table I) groups foci consisting of crypts of grade 1 only; grade 2 groups one or more crypts showing marked hyperchromatism, loss of nuclear polarity and mitotic figures. Also, tubular irregularity is common, goblet cells are rare and thus mucus production is very much reduced. Quite often, grade 1 and 2 can be demonstrated in one focus of dysplasia and it is easier to put them together in grade 1–2 (Fig. 8A, B).

(3) Carcinoma in situ (Fig. 9A, B) corresponds to the severe dysplasia grade 3 of Muto and Morson (1975). The crypts are tortuous and the nuclei larger, hyperchromatic and located at differing heights in the cell. Mitotic figures are present. There is no mucus production and goblet cells are either reduced to a tiny "goblet" close to the crypt lumen or more often are absent. Carcinoma in situ may be demonstrated in as small a group as 3–4 tubules either in a flat mucosa or in a polypoid lesion.

(4) Invasive carcinoma (Fig. 10): There is severe dysplasia with definite evidence of submucosal invasion. The tumours are then classified into grades I–IV of histological dedifferentiation according to Dukes (1940).

Both carcinoma in situ and invasive carcinoma are often surrounded by areas of hyperplasia and tubules showing various degrees of dysplasia (Fig. 9A, B).

During the first 3 weeks of the experiment (Table I), no histological changes are detected. In the following 3 weeks
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FIG. 7A

Fig. 7.—DMH treated rat. Grade 0–1 dysplasia. (A) H. and E. ×600. (B) HID-AB revealing the presence of sialomucins (blue) in the area of dysplasia, surrounded by normal glands producing sulphomucins (brown). ×150.
foci of hyperplasia are the only findings, with an occasional crypt showing mild or rarely severe dysplasia. From the 7th to the 13th week the number and severity of the histological lesions increase. During this period hyperplasia and all grades of dysplasia are observed. However, foci with severe atypia (grade 2) are still uncommon. From the 13th week onwards (Table I) the number of focal areas showing severe dysplasia (grade 2) increases markedly and histological features of carcinoma in situ appear. Invasive carcinoma is first demonstrated in rats killed after receiving 19 injections of DMH and in all rats with longer periods of treatment, except in one case where 5 areas of carcinoma in situ appeared in the mucosa but no invasive carcinoma was detected.

All the carcinomata found in the distal colon are well differentiated adeno-
Fig. 9.—DMH treated rat. Carcinoma in situ (T) surrounded by an area of hyperplasia and dysplasia. (A) HID-AB-sialomucins (grey) are the mainly secretory product in the areas of dysplasia accompanied by a decrease in sulphated material as shown in section (B) stained by HID. Areas of hyperplasia contain predominantly sulphomucins (black). Note absence of mucous secretion in the area of carcinoma in situ. × 600.
carcinomata (grade I), a few of them showing "endophytic" growth (Fig. 10). Three of 5 invasive carcinomata in the proximal colon produced abundant mucus and presented features of "signet ring" carcinoma whilst the other 2 were grade 1.

**Mucous secretion (histochemistry)**

*Controls.*—The composition and distribution of the epithelial mucins vary in the different segments of the colon and caecum (Table II). In the distal colon, the histochemical characteristics of the goblet cell mucins and their distribution in the crypt epithelium are similar to the human colonic mucosa. The goblet cells located along the whole crypt or in the $\frac{3}{4}$ lower part contain mainly sulphomucins (Fig. 3). Sialomucins may be present in the goblet cells of the upper crypt and surface epithelium. They also show a strong to moderate PAS reactivity. This pattern changes in the proximal colon, where sialomucins are the predominant content of the mucus secreting cells all along the crypt epithelium. Occasional goblet cells in the upper crypt and surface epithelium may show sulphated material (Fig. 4). The goblet cells of the upper crypt and surface epithelium have a strong to moderate PAS reaction whereas the mucus secreting cells in the lower crypt react weakly with PAS.

The transition from distal to proximal colon is gradual, sialomucins being present in the lower half of the crypt whereas in the upper half sulphomucins predominate.

In the caecum, goblet cells are distributed in reduced numbers from the middle crypt upwards and their contents have the staining properties of the sulphomucins, whilst in the bottom of the crypt both types of acid mucosubstances may be present.

Owing to the morphological and histochemical similarities between the rat distal colon and the human colorectal mucosa, we will refer only to the changes observed in the distal colon which may
be accepted as a suitable experimental model for the present work.

**Tests.**—The rats treated with DMH develop changes in the mucin composition. These consist of a predominance of sialomucins in the goblet cells, accompanied by a decrease or absence of sulphomucins, similar to those changes described in the human colonic mucosa in specimens with carcinoma. These modifications in the secretory product correlate well with the changes in the morphology.

Based on the relationship between mucin changes and focal areas of dysplasia, we may consider 3 or possibly 4 stages in the process of chemically induced carcinogenesis in the rat, as follows (Table I):

A. The mucosa in the first 6 weeks after the initiation of the treatment is either histologically normal or may show areas of hyperplasia. No alteration is observed in the mucin composition even in the areas of hyperplasia where goblet cells contain sulphomucins as in the controls.

B. From the 7th to 18th week, the number and severity of the lesions gradually increase and this increase is accompanied in general by marked qualitative mucin changes. Again, sulphomucins persist in the hyperplastic glands whereas goblet cells, in the majority of the areas with dysplasia (grade 0–1 and 1), produce mainly sialomucins (Fig. 5–8). Secretion is markedly reduced in grade 2 dysplasia and in the presence of a carcinoma in situ (Fig. 9A, B). In both, either sialo- or sulpho- or both types of acid mucins can be detected in the secretory product. During this period of time no carcinoma had yet developed in any of the rats.

C. The period between the 19th and 27th week is characterized histologically by the presence of invasive carcinoma and a greater number of areas with carcinoma in situ and severe dysplasia (grade 2). Areas of mild dysplasia are less frequent.

Mucin changes are not commonly seen and, with a few exceptions, foci of dysplasia show either sulphomucins or a mixture of both types of acid mucosubstances. In the distal colon all but 2 carcinomata do not show secretion, or else it is scanty. In one mucus secreting carcinoma there is abundant mucus rich in sialomucins; in the other, only moderate amounts of both sulpho- and sialomucins are present. In the proximal colon 3 "signet ring" carcinomata secreted a mixture of acid mucins (Fig. 11).

D. From the 28th week onwards the mucosa shows the same histological aspects as above but changes in mucin composition are more frequently demonstrated in a greater number of areas of dysplasia (grade 0–1 and 1).

**DISCUSSION**

The macroscopical and histological lesions we observed in the colonic mucosa of rats treated with DMH are similar to those described by several other authors (Druckrey *et al.*, 1967; Schauer, Völlnagel and Wildanger, 1969; Druckrey, 1970; Deschner, 1974; Ward, 1974) and resemble the human colorectal cancer, thus providing a suitable animal model for the study of various aspects of carcinogenesis.

The sequential changes occurring in the rat colonic mucosa were followed weekly for a period of 29 weeks and our observations correlate well with the features reported in a similar stepwise investigation carried out in mice by Thurnher *et al.* (1973) and Deschner (1974). The earlier changes detected at the 4th week are focal areas of hyperplasia in which sulphomucins still predominate as in the normal control musoca. As the experiment progresses the number of lesions with severe dysplasia increases, with carcinoma in situ developing at the 15th week and invasive carcinoma appearing at the 19th week. The areas of mild or moderate dysplasia show marked changes in the composition of the goblet cell mucin, characterized by the pre-
dominance of sialomucins and accompanied in general by a decrease or absence of sulphated material. Studies with $^{35}$S, in fact, show decreased isotope uptake in the non-neoplastic mucosa of rats treated with DMH (Springer, Springer and Oehlert, 1970; Filipe, in preparation) and also in the “normal” mucosa adjacent to colorectal carcinoma in man (Filipe, 1971b). It is of interest to note that similar qualitative changes of the epithelial glycoproteins have been found in the human large intestine not only in the “normal” mucosa around carcinoma but also in patches of “normal” mucosa far from the tumour (Filipe, 1969, 1971a; Filipe and Branfoot, 1974).

It is not yet possible to answer whether these modifications in the secretory product, in the human, represent an early feature of malignancy. The numerous genetical, physiological and environmental factors which may play a role in human carcinogenesis, as well as other unknown stimuli to which the colonic mucosa is exposed continuously, make it difficult to assess the “specificity” of the mucin changes in the human colonic mucosa. However, our present data from a rat model strongly suggest a relationship between the variations in the mucin composition and the process leading to malignancy, rather than these being the result of other stimuli (Smith and Butler, 1974) or of the toxic effects of DMH (Haase et al., 1973). The mechanism of action of DMH is not yet fully understood but it may interact with nucleic acids and/or protein (Miller and Miller, 1966; Shank and Magee, 1967; Farber, 1968) and thus alter the normal process of cell differentiation and maturation. Indeed, Lipkin and co-workers have described changes in the nucleic acid metabolism and proliferative capacity of the colonic epithelium in DMH treated mice (Thurnherr et al., 1973; Deschner, 1974) which are similar to those found in
the human colonic mucosa adjacent to carcinoma and neoplastic polyps (Immondi, Balis and Lipkin, 1969; Deschner and Lipkin, 1970; Troncale, Hertz and Lipkin, 1971) and in familial polyposis (Lipkin, 1974). These changes may reflect a loss of suppressor genes and a regression of the cell to a more embryonic state, a hypothesis supported in the demonstration of embryonic specific antigens in malignant tumours (Gold and Freedman, 1965; von Kleist and Burtin, 1969; Stonehill and Bendich, 1970; von Kleist, 1971; Bordes, Michiels and Martin, 1973).

Whether the raised levels of sialomucins in the colonic epithelium are considered as an expression of cell immaturity or as a result of a direct effect of the carcinogen on the mechanism(s) of the glycoprotein synthesis, and what effect it has on carcinogenesis, we do not know. However, there is cumulative biochemical evidence that the carbohydrate metabolism is profoundly altered in malignant transformed cells, and sialic acids may play an important role. The presence of sialic acids in the glycoproteins of the cell membrane seem to confer certain properties to the membrane expressed in cell adhesion (Deman, Bruyneel and Mareel, 1974), intercellular contact (Emmelot, 1973), masking antigens (Currie and Bagshawe, 1968; Rios and Simmons, 1973) and changes in its content may be related to the difference in behaviour of normal and malignant cells. Increase in sialic acids and sialyltransferases have been found in malignant cells (Warren, Fuhrer and Buck, 1972; Emmelot, 1973). Changes in glycosyltransferases, as they have been described in malignant cells, may also alter cell contact and the mechanism controlling cell growth and differentiation (Emmelot, 1973). Higher levels of glycosidases have also been reported in the “normal” colonic mucosa and tumours in animals treated with DMH, as compared with controls (Mian and Cowen, 1974; Mian, Ccwen and Nutman, 1974).

It has been suggested (Currie and Bagshawe, 1968; Rios and Simmons, 1973) that tumour cells are coated with neuraminidase sensitive sialic acids which may not only hide their antigens from the host but also shield them from his immunocompetent cells. If this hypothesis is correct, then they may play an important role in the immunological reaction to tumour specific antigens in the control of tumour growth (Baldwin, 1970). In fact, we have found greater amounts of neuraminidase sensitive neuraminic acids in the mucosa around carcinoma in man (Filipe and Cooke, 1974) and they are more abundant in our most invasive tumours (Filipe and Branfoot, 1974). In DMH treated rats we have also observed that the amount of sialomucins present in the focal areas of dysplasia varies during the progress of the experiment, possibly reflecting the immunological status of the host and thus affecting the tumour growth potential.

The changes in the type of glycoprotein secreted by the colonic goblet cells in DMH treated rats further support our previous hypothesis that the increase in sialomucins in the apparently normal mucosa in specimens with carcinoma may represent an early feature of carcinogenesis. At present there is no method which enables us to detect early malignant changes. We feel, therefore, that further investigation of glycoproteins may prove valuable in the search for such a method.

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