Histochemistry of the surface mucous gel layer of the human colon

K Matsuo, H Ota, T Akamatsu, A Sugiyama, T Katsuyama

Abstract

Background and aims—Histochemical analysis of the surface mucous gel layer of the human colon is difficult, as it dissolves in fixatives. This study was undertaken to explore the surface mucous gel layer on the normal mucosa and neoplastic tissues of the large intestine. In addition, the distribution of different mucins secreted from goblet cells was studied with a series of histochemical stains for mucins.

Methods—Twenty four surgically resected specimens were fixed in Carnoy’s solution and embedded in paraffin. In four cases, the surface mucous gel layer was also studied in frozen sections. Serial sections were stained by a battery of histochemical techniques characterising mucins.

Results and conclusion—The surface mucous gel layer consisted of the inner and outer layers. The first covered the luminal surface of the mucosa, consisted of mucins, and showed a vertical striped pattern. The second overlaid the first, showed a lateral striped pattern, and was contaminated with bacteria and other substances. Their thickness in paraffin sections varied considerably among the sites in the large intestine, but was the thickest in the rectum and measured 12.7 (SEM 6.0) μm and 88.8 (SEM 80.1) μm respectively. Mucins forming the inner layer were obviously derived from goblet cells underlying it.

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Keywords: surface mucous gel layer, mucin, histochemistry, colon cancer, colon adenoma.

The surface mucous gel layer of the large intestine probably serves as a lubricant to protect the mucosa. Its study in ordinary microscopical preparations is hampered by its loss during fixation. Stabilisation of the surface mucous gel layer was attained by Ota and Katsuyama by fixing surgically removed stomach in cooled Carnoy’s solution, clearing in xylene, embedding in paraffin wax, and sectioning. The surface mucous gel layer was well preserved. Some of us applied this technique to study the surface mucous gel layer of the large intestine and showed that it consisted of two layers, an inner obliquely striped layer and an outer multilaminated layer. The present study was undertaken to elucidate the distribution of the surface mucous gel layer on normal mucosa and neoplastic tissues of the large intestine. In addition, the distribution of different mucins secreted from goblet cells was studied by using a series of histochemical stains for mucins.

Materials and methods

Twenty four samples of surgically removed human colon were used in this study. These materials were obtained from 22 cases of colon cancer, one case of adenoma, and one case of familial adenomatosis coli. Locations of these specimens were caecum (one), ascending colon (three), transverse colon (four), sigmoid colon (eight), and rectum (eight). Immediately after resection all materials were opened along the contralateral portion to the lesion. Without rinsing, the specimens were laid flat with the mucosal surface up and pinned on cardboard. In 13 cases, 3% alcian blue solution in distilled water was sprayed onto the mucosal surface before immersing in fixative to investigate whether the mucous gel layer was an artifact during fixation. All specimens were immersed in Carnoy’s solution (ethanol 6:acetic acid 3:chloroform 1, v/v/v) for two hours at 4°C. They were then placed in 100% alcohol. Materials were sliced longitudinally at regular intervals of 5 mm width, cleared in xylene, and embedded in paraffin. After histopathological examination, several blocks, which were obtained from the cut end of the materials or included non-neoplastic mucosa as well as neoplastic tissues, were selected for histochemical staining. Serial paraffin sections of 3 μm thickness were prepared and stained to analyse mucins in the surface mucous gel layer. The Table gives the histochemical stains used and their histochemical relevance. Alkaline hydrolysis (1% potassium hydroxide for 15 minutes at room temperature) was performed to remove O-acetylated groups of 8-O-acetylated N-acetylnearuminic acid (8-O-AcNeuAc).

In addition, the surface mucous gel layer was explored in frozen sections in four samples, including two of ascending colon, one of transverse colon, and one of rectum. These materials were removed because of colon cancer, and tissue sections were obtained from macroscopically normal portions. Immediately after removal, these sections were laid in plastic cases (Cryomold, Miles Laboratories, Naperville, IL, USA) and snap frozen in OCT embedding medium (Miles Laboratories, Naperville, IL, USA) by submersion in liquid nitrogen. Sections of 5 μm thickness were cut on a cryostat and mounted on poly-L-lysine coated slides (Muto Pure Chemicals, Tokyo, Japan), dried at room temperature, coated with...
Histological staining: relevance and sources of reagents

<table>
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<tr>
<th>Method</th>
<th>Histochemical relevance</th>
<th>Sources of reagents</th>
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<tr>
<td>Alcian blue pH 2-5-PAS (AB/PAS)</td>
<td>To identify acid and neutral mucins.</td>
<td>Alcian blue (Chroma, Koengen, Germany); basic fuchsin (Wako Pure Chemical, Osaka, Japan)</td>
</tr>
<tr>
<td>High iron diamine/alcian blue pH 2-5 (HID/AB)</td>
<td>To differentiate sulphated mucins from non-sulphated sialomucins</td>
<td>N,N-dimethyl-p-phenylenediamine dihydrochloride (Kanto Chemicals, Tokyo, Japan); N,N-dimethyl-p-phenylenediamine monohydrochloride (Sigma, St Louis, MO, USA)</td>
</tr>
<tr>
<td>Periodic acid/sodium borohydride/potassium hydroxide-PAS (PA/SB/KOH/PAS)</td>
<td>To identify sialic acid with an O-acetylated side chain (8-O-acetyl-N-acetylamaminic acid or 8-O-AcNeuAc). Such a sialic acid is most abundant in goblet cells of the large intestine</td>
<td>Periodic acid (Wako Pure Chemical, Osaka, Japan); sodium borohydride (Wako Pure Chemical, Osaka, Japan); potassium hydroxide (Wako Pure Chemical, Osaka, Japan)</td>
</tr>
<tr>
<td>Periodic acid-cold thionin Schiff-potassium hydroxide-PAS (PA/TS/KOH/PAS)</td>
<td>To differentiate sialic acid with an adjacent hydroxyl radical (blue) from those without adjacent hydroxyl radicals (8-O-AcNeuAc; magenta)</td>
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0.2% celluloidin in 1:1 v/v ethanol-diethyl ether, and stained by haematoxylin and eosin (H and E), alcian blue/periodic acid Schiff (AB/PAS), high iron diamine/alcian blue (HID/AB), periodic acid/sodium borohydride/potassium hydroxide/potassium hydroxide/potassium hydroxide/potential acid Schiff (PA/SB/KOH/PAS), or periodic acid/cold thionine Schiff/potassium hydroxide/potential acid Schiff (PA/TS/KOH/PAS).

The thickness of the surface mucous gel layer was obtained by measuring at 10 randomly selected points in each section with an ocular scale. The thickness was defined as the distance from the outermost layer of the surface mucous gel to the luminal surface of the surface lining cells. The data are expressed as means (SEM).

**Results**

**HISTOLOGY OF THE SURFACE MUCOUS GEL LAYER**

In H and E preparations of ileum, no surface mucous gel layer was seen and only amorphous materials were scattered among villi. In preparations in which alcian blue solution was sprayed on before fixation, the luminal surface of the surface mucous gel layer was continuously covered by this pigment.

The surface mucous gel layer seemed to be a continuous eosinophilic layer coating the mucosal surface (Fig 1 also Fig 5A), although its thickness differed considerably among different sites in the large intestine. This layer was differentiated into an inner layer and an outer layer (Fig 2 and inset). The inner layer was consistently attached to the apical surface of the covering epithelia facing the intestinal lumen and was continuous with the intracrypt mucus. An obliquely striped pattern was often evident (Fig 2 and inset), except in the caecum, where this layer appeared as a homogeneous thin band. The outer layer overlaid the inner...
one, was less eosinophilic than the inner layer, and mostly showed a lateral striped pattern (Fig 2 and inset). The surface gel layer was thickest in the rectum and thinnest in the caecum and measured 31·1 (7·2) μm (range 26·0–36·3 μm) in the caecum, 34·4 (8·9) μm (range 26·8–44·1 μm) in the ascending colon, 50·5 (14·0) μm (range 38·3–45 μm) in the transverse colon, 62·0 (31·9) μm (range 17·3–115 μm) in the sigmoid colon, and 88·8 (80·1) μm (range 46·8–284·5 μm) in the rectum. Except in the ascending colon, the inner layer was also thicker in the distal colon and measured approximately 5·6 (0·2) μm in the caecum, 4·7 (1·4) μm in the ascending colon, 7·0 (3·7) μm in the transverse colon, 7·6 (3·4) μm in the sigmoid colon, and 12·7 (6·0) μm in the rectum. Cellular debris, food residues, and bacilli were often seen sandwiched between laminated arrays of the outer layer. The surface mucous gel layer was thicker on the transitional mucosa but had almost disappeared on the adenoma and carcinoma tissues (Fig 1), as described below.

**HISTOCHEMISTRY**

*Caecum*

In AB/PAS preparations, the goblet cells and intracryptal mucus stained blue/purple. The inner layer stained intensely. The outer layer, on the other hand, stained magenta with horizontal stripes of blue/purple. Staining with HID/AB showed two layered patterns in each crypt; the goblet cells lining the upper crypt and luminal surface stained predominantly for sulphated mucins, whereas those lining the lower crypt stained for non-sulphated sialomucin. The inner layer of surface mucous gel layer stained black, whereas the outer layer was a greyish blue colour with black horizontal stripes. In preparations stained with PA/SB/KOH/PAS goblet cells lining the crypts showed almost equal reactivities, as did the intracryptal mucus. The inner layer stained more intensely than the outer layer.

**Ascending colon**

With AB/PAS staining, goblet cells and intracryptal mucus predominantly stained blue/purple throughout the crypts (Fig 3A). The surface mucous gel layer stained blue/purple, showing a laminated structure (Fig 3A inset). With HID/AB staining, sulphated mucins predominated in the goblet cells and also were the major component of the overlying surface mucous gel layer (Fig 3B and inset). Multilaminated structure of the outer layer was evident by mucin staining. In PA/SB/KOH/PAS preparations, the goblet cells disclosed definite but weak reactivity at all levels of the mucosa (Fig 3C). The surface mucous gel layer also stained faintly (Fig 3C inset). Under the inner layer, the surface coat of the lining cells stained clearly with both AB/PAS and HID/AB. This finding was consistently found throughout the colon (Fig 4A and B).

**Transverse colon**

This region of the large intestine showed transitional patterns from the ascending to the sigmoid colon. In AB/PAS preparations, the colour of intramucosal goblet cells seemed to show more alcianophilia than that in the ascending colon, especially in the upper and middle portions of the crypts. The inner layer
of the surface mucous gel layer stained blue/purple and the outer layer showed a multilaminated structure. In HID/AB preparations, the number of goblet cells containing non-sulphated sialomucins slightly increased in the upper portion of the crypts, and non-sulphated sialomucins increased in the inner layer, although those containing sulphated mucins remained predominant. The outer layer also showed a laminated structure in PA/SB/KOH/PAS preparations, goblet cells lining crypts stained less than those in the ascending colon. The surface mucous gel layer also stained faintly.

**Sigmoid colon**

In the preparations stained with AB/PAS, goblet cells stained purple/red. The inner layer stained intensely red, whereas the outer layer stained pale magenta with a lateral striped pattern. In preparations stained with HID/AB, the number of goblet cells containing sulphated mucins decreased compared with the transverse colon, especially in the upper layer of the mucosa. The inner layer and outer layer stained only faintly, and the outer layer showed multilaminated structures of fine grey stripes on an almost colourless background. In preparations stained with PA/SB/KOH/PAS, both goblet cell mucins and the surface mucous gel layer were stained with moderate intensity. For the surface mucous gel layer, the outer layer almost always exhibited the multilaminated structure consisting of alternating PAS positive and PAS negative layers.

**Rectum**

With AB/PAS staining, goblet cells lining crypts and intracryptal mucins stained red/purple (Fig 5B). The inner layer stained similarly but occasionally showed intense alcianophilia. The outer layer, on the other hand, showed alternating laminated structures consisting of blue/purple and blue layers (Fig 5B inset). With HID/AB staining, goblet cells lining the upper crypts predominately stained for non-sulphated sialomucins, whereas those lining the lower crypts stained for sulphated mucins (Fig 5C). The inner layer contained mostly non-sulphated sialomucins, and the outer layer consisted of non-sulphated sialomucins and sulphated mucins abutting each other (Fig 5C inset). With PA/SB/KOH/PAS staining, goblet cells and the surface mucous gel layer stained as light as those in the transverse colon (Fig 5D and inset).

**Colon adenoma**

Adenomas were recognised in 16 lesions from seven cases. There were two cases of tubulovillous adenoma with moderate dysplasia, eight cases of tubular adenoma with mild dysplasia, and six with moderate dysplasia. In most cases examined, the surface mucous gel layer was not identified on the adenoma tissues, regardless of the abundance of neoplastic goblet cells (Fig 6A and B). A very thin surface mucous gel layer was seen in three cases. In these cases, only the outer layer persisted.
Colon cancer

Hardly any surface mucous gel layer overlaid cancerous tissues in most lesions, although mucins derived from neoplastic goblet cells occasionally adhered to the surface. Even in the region where numerous goblet cell type carcinoma cells lined the luminal surface, the inner layer was never found (Fig 7A and B).

So called “blue crypts” and the surface mucous gel layer

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Effect of alkaline hydrolysis on AB/PAS and HID/AB staining

Prior alkaline hydrolysis with potassium hydroxide greatly enhanced PAS reactivity of most goblet cells throughout the colon. A similar effect was confirmed for their alcianophilia. This pretreatment also enhanced HID reactivity of mucins. The surface mucous gel layer also gained higher reactivities for these stains. The effect of this pretreatment was most evident in the sigmoid colon, although the surface mucous gel layer still stained less than that of the rectum with HID/AB stain. After this pretreatment, goblet cells and the surface mucous gel layer remained unstained with PA/SB/KOH/PAS and stained homogeneously blue with PA/Ts/KOH/PAS. The “blue crypts” and their mucins secreted, on the other hand, were not influenced by this pretreatment and therefore became indistinguishable from the surrounding crypts.

In the four cases examined by frozen sections, the surface mucous gel layer appeared as a continuous layer coating the mucosal surface and consisted of an inner and an outer layer (Fig 10A, B, and C). The inner layer clearly showed a vertical striped pattern which reflected the mucus spouting from each goblet cell. The outer layer showed no particular pattern, and its luminal surface was not clear. The thickness of the inner layer was 19-6 (3-0) μm (range 17-3-23 μm) (Fig 10A, B, and C). In the inner layer, bacteria were seen along the vertical stripes (Fig 10A and B).

Discussion

The present study showed that fixation in Carnoy’s solution preserves the surface mucous gel layer in ordinary paraffin sections and allows these to be analysed by histochemical techniques.

Several attempts have been made to observe the surface mucous gel layer of the gastrointestinal tract. Kerss et al explored it by preparing unfixed thin sections of rat, frog, and human stomach and recently Pullan et al prepared sections of human colon and measured its thickness. They also used Carnoy’s fixation to support the existence of the surface mucous gel layer of the colon. As stated previously, sectioning of the surface mucous gel layer by razor blades is not easy because of its stickiness. In addition, histochemical analysis of this layer is difficult as it dissolves in solution during staining. Ota and
Katsuyama showed that Carnoy's solution was a useful fixative to preserve the mucus in paraffin blocks. The surface mucous gel layer was histochemically analysed by preparing thin sections, although there was obvious shrinkage of the mucus during fixation and preparation of paraffin wax blocks. The present study confirmed two layers in the surface mucous gel, the inner layer and the outer layer, as suggested previously. This structure was consistent in all cases examined but the question still remained whether the structure was an artificial one or not. Several findings, however, indicate the presence of this pattern. Firstly, the inner and outer layers were clearly identified in frozen sections, although the inner frozen layers were thicker than they were in paraffin sections. The luminal surface of the outer layers was not very clear in the frozen sections but was obvious in the paraffin sections. Secondly, size, distribution, and mucin content of goblet cells in the tissue preparations fixed in Carnoy's solution did not differ significantly from these features of goblet cells fixed in ordinary formalin. This finding indicates that abrupt mucin discharge did not occur during fixation in Carnoy's solution. Thirdly, the inner layer was consistently absent on adenoma or carcinoma tissues which contained numerous goblet cell type neoplastic cells. Fourthly, the surface coat, which characterises the apical surface of covering epithelium including goblet cells, preserved well under the inner layer, suggesting no disruption of the apical plasma membrane. Fithly, the inner layer was not found in ileum, which is also rich in goblet cells. Other support comes from our study on the human stomach. Gastric mucosa often showed intestinal mucosa which contained plenty of well developed goblet cells. In the stomach, however, we have never found the inner layer in the surface mucous gel layer.

So called "blue crypts" provided the clue to explore the origin of mucins in the surface mucous gel layer. The term "blue crypt" was coined by Kato et al because of the characteristic reaction to PA/T/S/KOH/PAS. Its relevance as a precancerous lesion was contradicted by recent study, however. The present study showed that goblet cells lining blue crypts lacked 8-O-AcNeuAc, a histochemical marker of the large intestine, as goblet cell mucins remained negative by PA/

Figure 8: (A–D) So called "blue crypts" in the sigmoid colon prepared from serial sections. Originally ×96. (A) Goblet cells lining blue crypt and the inner layer which cover only the extent of these crypts stain blue/purple. AB/PAS. (B) They stain more intensely than crypts of the surrounding mucosa. HID/AB. (C) They are entirely unstained by the PA/SB/KOH/PAS sequence (arrows). (D) With the PA/T/S/KOH/PAS sequence they stain intensely blue and, therefore, were called "blue crypt".

Figure 9: (A,B) Higher magnification of fig 8 (A,D). Evidence that mucous derived from "blue crypts" spreads along the boundary layer between the inner and outer layers (arrow heads) or spreads into the outer layer (arrows). Originally ×192.
groups on C-7,8, and 9 of sialic acid. In addition it might also cleave the ester linkage between acid groups and hydroxyl groups of sugar moieties, resulting in increased HID and AB reactivities of acid mucins.

Histochemical staining suggested that the mucins in the outer layer stained similarly to goblet cell mucins. They also suggested that at least sugar moieties of mucins were maintained in the outer layer, as, for example, 8-O-AcNeuAc occupies the non-reducing terminal portion of oligosaccharides, and its existence suggested the persistence of saccharide chains. Various mucins derived from goblet cells did not mix homogeneously in the surface mucous gel layer but rather stacked on each other and formed the laminated structure, as exemplified by 8-O-AcNeuAc free mucins spread along the boundary layer and in the outer layer.

The surface mucous gel layer varied in thickness in different regions of the colon. The descending order of thickness was rectum, sigmoid colon, transverse colon, ascending colon, and caecum. Pullan et al also reported that the the surface mucous gel layer was thinnest in the left colon and the thickest layer was in the rectum. The thickness in their report was about twice that of ours. The difference between the two studies reflects the method used, because, as described previously, there was appreciable shrinkage in Carnoy’s solution. It remains to be clarified what factors determine the thickness of this layer.

The surface mucous gel layer probably reduces friction between the mucosa and faeces. In this study, the inner layer only occasionally contained bacteria. Thus it may also play a part in inhibiting the direct adherence of bacteria to colonic epithelia.

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