Colonic mucin synthesis is increased by sodium butyrate

I A Finnie, A D Dwarakanath, B A Taylor, J M Rhodes

Abstract

The effects of sodium butyrate and sodium bromo-octanoate (an inhibitor of \( \beta \) oxidation) on colonic mucus glycoprotein (mucin) synthesis have been assessed using tissue from colonic resection samples. Epithelial biopsy specimens were incubated for 16 hours in RPMI 1640 with glutamine, supplemented with 10% fetal calf serum and \( N \)-acetyl-[\( ^{3} \)H]-glucosamine (\( ^{3} \)H-Glc NAc), and differing concentrations of sodium butyrate. Incorporation of \( ^{3} \)H Glc NAc into mucin by normal epithelium at least 10 cm distant from colonic cancer was increased in the presence of sodium butyrate in a dose dependent manner, with maximum effect (476%) at a concentration of 0-1 mM (number of specimens=24 from six patients, \( p<0.001 \)). The increase in response to butyrate was not seen when specimens were incubated in the presence of the \( \beta \) oxidation inhibitor sodium bromo-octanoate 0-05 M. The striking increase in mucin synthesis that results when butyrate is added to standard nutrient medium suggests that this may be an important mechanism affecting the rate of mucin synthesis in vivo and may also explain the therapeutic effect of butyrate in colitis.

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Keywords: sodium butyrate, colonic mucin synthesis.

The primary function of colonic mucus is considered to be protection of the surface mucosal cells.\(^1\) Mucus forms a gel that adheres to the epithelium, preventing invasion by colonic bacteria and protecting against damage by bacterial toxins and enzymes. Most of the protective effect of mucus is thought to relate to the presence of mucin, the predominant glycoprotein in mucus. Abnormalities of mucin have been shown in ulcerative colitis using histochetical,\(^2\) biochemical,\(^3\) and tissue culture techniques,\(^4,5\) and an attractive hypothesis for ulcerative colitis is that a primary abnormality of mucin synthesis predisposes subjects to an as yet undefined environmental factor that induces the disease.\(^6\)

Exacerbations of ulcerative colitis can be successfully treated with sodium butyrate,\(^7\) although the exact mode of its action is not clear. Butyrate is an important source of energy for the colonic epithelial cells,\(^8\) and a local deficiency of butyrate or an inborn error in its metabolism\(^9\) have been implicated in the disease. We have investigated the effects of sodium butyrate on colonic mucin synthesis; if mucin synthesis were enhanced by butyrate this might help to explain its therapeutic mode of action.

Methods

MUCOSAL BIOPSY SPECIMENS

Fresh colonic tissue was obtained from patients undergoing colectomy for cancer (n=6, median age 70, 3 men 3 women), colonic inertia (n=2, ages 26 and 39, both women) or ulcerative colitis that was unresponsive to medical treatment (n=2, ages 55 and 32, both men). Bowel preparation consisted of clear fluids for 48 hours and Picolax two sachets on the day before operation, except in the patients with ulcerative colitis in whom no formal bowel preparation was used (although as a result of their illness they also had been taking very little residue). The site of colonic cancer was rectum (n=1), sigmoid (n=3), and ascending colon (n=2); the tissue studied was from sites at least 10 cm distant from the macroscopic disease, and in each case histological examination of adjacent mucosa confirmed that there was no abnormality. In the patients with colonic inertia, tissue from the ascending and descending colon was studied separately. In the patients with ulcerative colitis, the comparatively uninvolved right colon was studied. These two patients had been treated with high dose intravenous corticosteroids for at least 10 days, and had been receiving maintenance 5-aminosalicylic acid drugs before relapse, but neither had been treated with butyrate enemas.

The resected tissue was pinned out, and multiple mucosal specimens each weighing about 20 mg were separated from underlying tissue using a scalpel. Specimens were then cultured as described below.

TISSUE CULTURE

Mucosal specimens were cultured using a technique based on that described by Macdermott \textit{et al.}\(^1,11\) Each specimen was placed on an alloy mesh (David, Wellingborough, UK), orientated so that the epithelial surface was uppermost. The wire grid was floated on 1 ml culture medium in an organ culture dish (Becton Dickinson, NJ, USA). The culture medium was 90% RPMI 1640 with glutamine (Gibco, Paisley, UK) and 10% fetal calf serum (Gibco) with gentamicin 100 \( \mu \)g/ml and nystatin 60 \( \mu \)g/ml, to which was added 1-5
was for 18 hours at 37°C in 95% O₂/5% CO₂, except in one experiment, when varying periods of culture were used to assess specimen viability. A parallel time course experiment was conducted to exclude the possibility that colonic bacteria were incorporating the radiolabel, whereby 50 μl of faecal fluid taken from the lumen immediately after colectomy, in the absence of a mucosal biopsy specimen, was added to the culture system.

**ESTIMATION OF MUCIN SYNTHESIS**

At the end of the culture period, the mucosal surface of each specimen was gently rinsed in its culture medium to remove secreted but adherent mucus, placed in 1·3 ml 0·1 M TRIS/HCl at pH 8 (TRIS/HCl) containing thimerosal 100 mg/l and sonicated using an MSE ultrasonic disintegrator (MSE Instruments, Crawley, UK). Each sample was then centrifuged for 15 minutes at 11 000 g and 1 ml of the supernatant retained and lyophilised after removal of a 100 μl aliquot for protein estimation using a bicinchoninic acid protein assay kit (Sigma, St Louis, USA). The culture medium was also retained and lyophilised separately from the specimen.

After lyophilisation, each sample was resuspended in 200 μl water and subjected to gel filtration using gravity fed mini-columns (PD 10) packed with Sepharose CL-2B (Pharmacia, Uppsala, Sweden). The samples were then eluted with deionised distilled water at a rate of 0·5 ml/min, the samples containing mucin (void volumes) retained, mixed with 5 ml scintillant (Optiphase Safe, Pharmacia), and counted in a Rackbeta β counter (LKB Wallace, Turku, Finland).

Total mucin synthesis was expressed as dpm [³H] N-acetylglucosamine incorporated into purified mucin (biopsy specimen plus culture medium) per μg biopsy protein.

**CALIBRATION OF GEL FILTRATION COLUMNS AND VERIFICATION OF MUCIN PURITY**

The gel filtration columns containing Sepharose CL-2B were calibrated using purified [³H] Glc NAc labelled mucin, which was obtained by culturing mucosal biopsy specimens as described above, preparing crude mucin by sonication of the specimens in TRIS/HCl, and subjecting this crude mucin to gel filtration using monodisperse agarose (Superose 6) (Pharmacia) using the method of Parker et al. The fractions containing purified mucin were pooled, lyophilised, reconstituted with 200 μl water, and then subjected to gel filtration using the Sepharose CL-2B columns. Deionised distilled water was added in 0·5 ml aliquots at a rate of 0·5 ml/min. Twenty 0·5 ml fractions were collected, mixed with scintillant, and β counted. The elution profile of 1 μCi [³H] Glc NAc in 200 μl TRIS/HCl was studied as a comparison.

Superose 6 purified mucin that was labelled with [³H] Glc NAc was found to elute reproducibly from the Sepharose CL-2B columns between 2 and 3·5 ml (Fig 1A). Figure 1B
Figure 2: Protease digestion of Sepharose CL-2B purified [3H Glc NAc] labelled mucin. Mucin was incubated with (A) phosphate buffered saline as a control, (B) pronase, or (C) trypsin, and then subjected to gel filtration using Sepharose CL-2B mini-columns. The protease incubated mucin fragments mainly into large and medium sized glycoproteins (see comparison with Fig 1B). The profile shown is for mucin secreted into culture medium, but similar profiles were obtained for biopsy specimen homogenates and incubation with proteoglycan degrading enzymes had no effect on subsequent elution profiles.

shows the elution profile of a mixed solution of blue dextran (MW 2×10⁶ Da) (Pharmacia) and bovine serum albumin (MW 67 000 Da) on the mini-columns. Elution of [3H] Glc NAc occurred separately, and addition of 1 µCi to labelled mucin produced no significant change in scintillation in the mucin containing fractions confirming lack of non-specific binding (Fig 1C). The yield of [3H] Glc NAc labelled mucin (mean (SD)) from the Sepharose CL-2B columns was 95 (4)% (n=6).

Fractions of Sepharose CL-2B separated mucin were also analysed using an enzyme linked mucin antibody (CAM 17·1) binding assay (Fig 1D), which confirmed the elution of mucin in the void volume.

To assess the comparative purity of the mucin samples obtained using Sepharose CL-2B gel filtration, aliquots of [3H] Glc NAc labelled mucin (about 0·2 µg protein as determined by bicinchoninic acid) purified from specimens and medium using this technique were incubated in the presence of (a) hyaluronidase (hyaluronoglucosaminidase, Boehringer, Lewes, UK) 1 mg/ml at pH 5·0 for one hour at 37°C, (b) collagenase (Sigma) 10 mg/ml at pH 7·4 for five hours at 37°C, (c) chondroitinase ABC (Boehringer) 0·01 U at pH 8 for one hour at 37°C, (d) pronase (BDH) 7000 PUK units at pH 7·6 for one hour at room temperature, (e) bovine pancreas trypsin (EC 3.4.21.4) (Sigma) about 100 000 units for one hour at room temperature or (f) in phosphate buffered saline as controls. The samples were then cooled, and repeat gel filtration using Sepharose CL-2B mini-columns was performed. The fractions were collected, mixed with scintillant, and β counted.

Incubation of [3H] Glc NAc labelled mucin with hyaluronidase, collagenase, and chondroitinase had no effect on the subsequent elution profiles, whether the mucin studied had been purified from biopsy specimen or culture medium: however, a small (<5%) loss in recovery was seen after incubation with hyaluronidase of mucin purified from culture medium. Pronase and trypsin caused a change in the elution profile of the labelled mucin (Figs 2 and 3), but with much of the radiolabel still detectable in high molecular weight components, in keeping with the known protease resistance of mucin subunits.

Glycolipid contamination was assessed by extracting Sepharose CL-2B purified mucin labelled with [3H] Glc NAc with two volumes chloroform/methanol 2:1. The chloroform and aqueous layers were separated, with scintillant, and their radioactivity counted. Washing with chloroform/methanol reduced subsequent recovery of mucin by 2·5 (3·2)% (mean (SD), n=5) showing minimal contamination by glycolipids.

In addition, aliquots of [3H] Glc NAc labelled mucin purified on Sepharose CL-2B were collected, lyophilised, and subjected to gel filtration using Superose 6. The fractions were mixed with scintillant and β counted. These were compared with the results obtained by purifying aliquots of the same sample using Superose 6 gel filtration alone. [3H] Glc NAc labelled mucin purified using Sepharose CL-2B eluted in the same position as pure mucin when subsequently subjected to gel filtration using Superose 6, 85 (9) (mean (SD)) per cent eluting between fractions 7–10 (Fig 4), which previous studies have shown to be free from contaminating non-mucin glycoproteins.

TIME COURSE OF MUCIN SYNTHESIS AND LACTATE DEHYDROGENASE RELEASE

There was a noticeable lag period of six to eight hours when the rate of incorporation of [3H] Glc NAc into mucin was low, followed by a period when the rate of mucin synthesis (total of mucin in biopsy specimen + culture medium) was roughly linear for up to 26 hours (Fig 5). Radiolabelling of mucin in the biopsy specimen homogenate mucin increased gradually to a peak at around 20 hours, whereas it continued to rise in the culture medium for the duration of the experiment. Culture of faecal fluid without biopsy specimen had no significant effect on incorporation of radiolabel into the Sepharose CL-2B column void volume.
REPRODUCIBILITY

The mean coefficient of variation for the incorporation of [3H]-N-acetyl glucosamine into mucin by different (histologically normal) colonic biopsy specimens from the same area of the colon and patient (expressed in comparison with biopsy specimen protein content) was 16% in non-treated (n=44 biopsy specimens, 8 patients), and 20% in butyrate treated specimens (n=164 specimens, 8 patients). In UC the mean coefficient of variation was 21%.

LACTATE DEHYDROGENASE RELEASE

In a further experiment, biopsy specimens were cultured as described above but without radiolabel for 24 hours, with changes of culture medium every four hours. The culture medium was kept at ~4°C for up to 24 hours, and then analysed for lactate dehydrogenase using the colourimetric method of Jones et al. Specimens were ultrasonicated in 1 ml TRIS/HCl at the end of the culture period and lactate dehydrogenase activity (after 1/50 dilution) and protein concentration assessed.

Lactate dehydrogenase release into the culture medium was high during the first four hour period (possibly because of leakage of enzymes from the cut surface). Thereafter the rate of release was steady throughout the 24 hour culture period, and was low in comparison with the biopsy specimen content at the end of the culture period (Table).

STATISTICAL ANALYSIS

Comparisons of the values obtained for mucin synthesis in the presence of different concentrations of butyrate and bromo-octanoate of the drugs studied were made by non-parametric analysis of variance (Kruskal-Wallis) followed by multiple paired comparisons. Comparisons between the left and right colon were made by the Mann-Whitney U test. Differences were considered significant at p<0.05.

Results

EFFECT OF SODIUM BUTYRATE AND SODIUM BROMO-OCTANOATE ON MUCIN PRODUCTION

Histologically normal tissue

Control biopsy specimens from the ascending colon (from the two patients with colonic inertia) showed a higher rate of total mucin synthesis than specimens from the descending colon (ascending 35.2 (11.8) (mean (SD), n=10) dpm/μg protein/h, descending 25.5 (10.6) (n=10), p<0.05). Histologically normal tissue from the ascending colon and descending colon from patients with carcinoma showed a much wider variation (ascending 26.4 (16.2) dpm/μg protein/h, descending 21.9 (15.3)).

There was an increase in mucin synthesis when specimens from the patients with colon cancer were incubated in the presence of...
sodium butyrate at all concentrations tested. The effect was most noticeable at concentrations between 0-1–1 mM; the increase was statistically significant at all concentrations tested except at the lowest (0.05 mM) and at the highest (10 mM) (Fig 6). The increases in mucin synthesis were of a similar magnitude when specimens from the ascending and descending colon of the same patients (with colonic inertia) were incubated with sodium butyrate at 0.1 mM (ascending 272 (168)% (mean (SD)), descending 306 (108), n=10); 0.5 mM (ascending 417 (257), descending 475 (349), n=10); and 1 mM (ascending 297 (178), descending 309 (197), n=10).

A concentration of 0.05 M sodium bromo-octanoate did not itself have any effect on mucin production, but abolished the butyrate induced increase in mucin production (Fig 7).

Mucin recovered from the medium accounted for 73 (12)% (mean (SD)) of mucin synthesised during the 16 hour culture period.

**Ulcerative colitis**

The rate of mucin synthesis in 10 specimens from the two ulcerative colitis patients (24-6 (12-2) dpm/µg protein/h) (mean (SD)) was marginally (not significantly) lower than the rate seen in the right colon in histologically normal tissue. When specimens from the patients with ulcerative colitis were incubated in the presence of sodium butyrate at 0-1 mM, 0.5 mM, and 1 mM there was an increase in the rate of mucin synthesis, which was less pronounced than that seen in histologically normal tissue, and was significant only at a concentration of 0.5 mM (Fig 8).

**Discussion**

This study has shown that in vitro mucin synthesis by colonic biopsy specimens occurs at a linear rate from six to 28 hours, can be stimulated over fourfold by sodium butyrate, and that this stimulatory effect is abolished by the inhibitor of β oxidation, sodium bromo-octanoate.

We found a lag period after starting the culture, during which there was little measurable incorporation of radiolabel by colonic specimens. Although some studies of glucosamine incorporation do not report a lag
period, others do, and it is probable that this represents the time taken for the radiolabel to be incorporated into completely synthesised mucus glycoprotein. Incompletely glycosylated mucins may not be excluded by Sepharose CL-2B, and it should be remembered that the time for newly formed glycoprotein to appear at the apical cell surface of human goblet cells is six to eight hours. After the lag period, specimens synthesised mucin at a constant rate.

Lactate dehydrogenase release from specimens in culture was roughly linear over 24 hours, and the rate of release was small in comparison with whole tissue lactate dehydrogenase activity. This shows that the tissue remains viable throughout the culture period and this is further supported by the linear incorporation of radiolabelled substrate into mucin throughout the culture period.

A purification step of some kind is important in studies of mucin synthesis as 50% of radiolabelled hexosamine may be incorporated into non-mucin glycoprotein by the epithelium in culture. The rapid single stage technique of mucin purification described here has the advantage that multiple samples can be purified simultaneously. The mucin thus obtained has been shown to be free from significant contamination by glycolipids and proteoglycans, and the comparative resistance to protease digestion and the high molecular weight fragments that result are strong evidence that the purified radiolabelled substance is largely mucin. Mucin does not interfere with the elution of free radiolabelled GlcNAc, and the free radiolabel and mucin are well separated.

This study shows the importance of assessing the culture medium in addition to the biopsy specimen homogenate when assessing mucin synthesis, as about three quarters of incorporated radiolabel in this study was found to be in mucin, which has been secreted into the medium. This is rather more than was found in previous studies, probably because incompletely glycosylated intracytoplasmic mucins were not excluded by the Sepharose CL-2B. Mucin purification by acid precipitatin will probably result in the co-purification of small and medium sized glycoproteins with mucin, and these smaller molecules (including incompletely glycosylated mucins) are more likely to be found in the tissue homogenate than in secretions.

Butyrate, which is formed in the colon by bacterial fermentation of dietary fibre, is the preferred energy source of colonic epithelial cells and stimulates oxygen utilisation by isolated colonocytes in culture. The luminal butyrate concentration is reduced by fasting, and is increased specifically in the proximal colon by ingestion of easily fermentable fibres (for example, oats, guar), and in the distal colon by ingestion of wheat bran, which is less easily fermentable. Butyrate reduces proliferation and differentiation in colonic cancer cell lines in vitro, but paradoxically increases proliferation in the normal colon, and a reduction in butyrate metabolism by the colonic epithelium has been proposed as a possible aetiological factor in carcinogenesis.

The concentration of butyrate that induced maximal stimulation of mucin synthesis is 100-fold less than that in the human colonic lumen, but the concentration in the portal blood is only 0.029 mM, and the concentration bathing the epithelial cell in vivo may be considerably lower than that in the lumen. Although impaired butyrate metabolism has been reported in isolated epithelial cells extracted from the mucosa in ulcerative colitis this was not confirmed in a recent study in which butyrate metabolism was studied using mucosal biopsy specimens.

Nevertheless, butyrate reduces proliferation in the mucosa in ulcerative colitis, and butyrate enemas are an effective treatment for diversion colitis and ulcerative colitis. Sodium bromo-octanoate inhibits fatty acid oxidation in perfused rat liver and in mitochondria isolated from rat liver, probably by inhibiting an enzyme of β oxidation, and experimental colitis can be induced in laboratory animals by bromo-octanoate enemas. In this study sodium bromo-octanoate completely blocked the stimulatory effect of butyrate on mucin synthesis although when tested alone did not inhibit mucin synthesis. This suggests that the colonic epithelium, at least in short term culture, can use alternative sources of energy if β oxidation is inhibited.

Goblet cell depletion of mucus is a characteristic histological feature of ulcerative colitis. Previous studies of mucin synthesis by ulcerative colitis biopsy specimens cultured in vitro have had conflicting conclusions, with reduced synthesis in inactive ulcerative colitis in one study, but normal in another. It has been argued that the increase in cell turnover seen with this disease should be accompanied by an increase in mucin synthesis, and that mucin synthesis is deficient as compared with the rate of cell proliferation. If mucin synthesis and proliferation were directly linked, however, the expected response to butyrate in ulcerative colitis would be a decrease in mucin synthesis rather than the increase described here (as butyrate enemas...
are known to reduce epithelial proliferation in ulcerative colitis). An alternative explanation is that by increasing mucin synthesis, butyrate increases mucosal protection, and this permits the rate of proliferation to reduce to a more normal rate, possibly by blocking the binding of luminal growth factors that might be dietary in origin or secreted locally. The increase in mucin synthesis induced by butyrate probably reflects an important therapeutic mode of action, as well as a physiological function.

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