Bile Salts Up-Regulate MUC5AC, Not MUC5B, Mucin Expression in Cultured Airway Goblet Cell Model HT29-MTX Cell Line

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Introduction:

The production of MUC5AC and MUC5B, the major mucins expressed by airway mucosa, could be altered by various contents of laryngopharyngeal reflux. This study used the cell line HT29-MTX as an airway goblet cell model, to analyse the possible effect of bile salts on MUC5AC and MUC5B mucin production.

Methods:

Goblet cell line derived from the human colon carcinoma cells HT29-MTX was cultured and challenged by exposure to various concentrations of a physiologic combination of bile salts. Modified ELISA was employed to measure of MUC5AC and MUC5B contents before and after challenge.

Results:

Cultured HT29-MTX cell line maintained full viability when challenged with bile salts up to 20 µmol/L. This stimulated MUC5AC production in a dose dependent manner and occurred within the first 24 hours after challenge. MUC5B production stayed at the base line.

Discussion:

Increased MUC5AC production by the bile salt challenged HT29-MTX cell line could be due to mucin upregulation or merely due to stimulated mucin release from mucin stores in the cells. This could be achieved either directly or via the release of pro-inflammatory cytokines such as IL-8. MUC5B production may need to be studied using different cell lines. Bile salts in laryngopharyngeal reflux could alter mucin production.

Conclusion:

Cultured HT29-MTX cell line can be used as a model for airway goblet cells. Bile salts challenge to these cells resulted in MUC5AC but not MUC5B release suggesting a potential rule of refluxed bile salts in modulating airway mucin expression.

regulate MUC4 expression in oesophageal cancer [8]. It had not been studied, however, whether bile salts could alter MUC5AC and/or MUC5B expression. These are the two major airway mucins produced mainly by goblet cells and secretory cells of the submucosal glands respectively [9,10]. These 2 types of cells have been found throughout the upper and lower airways and their mucin expression can be affected by various inflammatory and neoplastic conditions. Altered mucin expression and hence mucus properties can compromise mucociliary clearance and may have deleterious clinical effects [11] that could contribute in the aforementioned airway diseases.

This study investigates the potential of a physiological mixture of bile salts to stimulate mucin expression by cultured goblet cells. This was achieved by creating a physiological bile salts damage model using human goblet cells and measuring their MUC5AC and MUC5B production as makers of damage caused by exposure to bile salts.

**Methods**

**Reagents:** Unless stated otherwise reagents used were obtained from Sigma-Aldrich UK Ltd., Fisher Scientific UK Ltd. or BDH Merck Ltd.

**Goblet cell culture (HT29-MTX)**

A human goblet cell line derived from human colon carcinoma cells HT29 was cultured according to a previously described method [12]. HT29 cells were stepwise adapted to increasing concentrations of methotrexate (MTX), to produce a mucus-secreting phenotype. These HT29-MTX cells in culture form a homogeneous monolayer of polarized goblet cells which resemble characteristics of lung goblet cells (Figure 1), of which MUC5AC and MUC5B are the main secretory mucins.

![Figure 1: Goblet cell line HT29-MTX (200-fold magnification) growing in culture under normal conditions (Dulbecco’s modified eagle’s medium (DMEM) supplemented with 10% foetal calf serum (FCS), LONZA, Switzerland, at 37°C and 10% CO2) at 80-95% confluence.](image)

1 ml of Dulbecco’s modified eagle’s medium (DMEM) supplemented with 10% foetal calf serum (FCS), 50U/ml penicillin, 50mg/ml streptomycin, 50µg/ml gentamycin, and 50µg/ml amphotericin B (both LONZA Switzerland) was mixed into a set of liquid nitrogen frozen down cells. The supplemented medium and cells were further transferred to a T75 flask with 12 ml media each and placed in an incubator at 37°C and 10% CO₂. The first change of media and PBS wash was carried out after 24 hours to prevent cell death. Afterwards, media was changed every 48 hours until cells reached confluence. Confluent cells were passaged with a trypsin solution containing EDTA. Media in the T75 flask was discarded and the confluent cells incubated with trypsin for 2 to 3 minutes until they were mobilized in the flask. The flask was then washed with several changes of culture media supplemented with 10% FCS to catch any remaining cells. The cell suspension was collected in a 25ml falcon tube and centrifuged for 7 minutes at 1100 c.r.p.m. The supernatant was discarded, the pellet containing the cells re-suspended with fresh media and transferred to a T75 flask for expansion followed by a 24-well plate for experimentation (0.5ml per well). For cell count cells were passaged from the T75 flask into a falcon tube with 12 ml serum-free DMEM resting media. 0.5ml of this cell suspension was mixed with 20 ml isotonic buffer and run on a cell counter.

**Bile salt challenge of goblet cells**

Cells were seeded at approximately 50000 per well and after reaching confluence in 24-well plates the goblet cells were rested for 24 hours by incubation with 0.75 ml serum free DMEM at pH 7.4. The cells were challenged with a physiological mix of bile salts. This was composed of 50% cholic acid, 30% chenodeoxycholic acid, 15% deoxycholic acid and 5% lithocholic acid dissolved in methanol and diluted with media to give required concentrations in the proportions found in human bile [13]. We used the following concentrations: 3, 6, 9, 12, 15, 18, 20 µmol/L. Bile salts were dissolved in methanol (20 mmol/L solution), diluted down with media to the required concentrations and filtered through a sterile syringe before challenging the cells. Controls contained media free from bile salts. 0.2 ml media was collected at 24, 48 and 72 hours and assayed for MUC5AC and MUC5B by a developed ELISA as described later. Data was plotted for each bile salt concentration at 24, 48 and 72 hours.

**Cell viability assay**

Viability of the goblet cells after 72 hours bile salt challenge was measured with the CellTiter-Blue® Cell Viability Assay (Promega, USA) in a 96-well plate (Maxisorp, NUNC). 100% cell death was produced by the addition of 200 µl frozen methanol for approximately 5 minutes. A standard curve was produced using mixtures of live cells (LC) and methanol killed dead cells (DC) as follows: (100%DC, 75%DC/25%LC, 50%DC/50%LC, 25%DC/75%LC).

**Measurement of basal mucin production**

Culture media of human HT29-MTX cells were subjected to two round of Caesium Chloride equilibrium density gradient centrifugation [14] to isolate and purify polymeric mucin glycoprotein produced by HT29-MTX cells. Mucin glycoprotein...
content was measured by periodic acid Schiff’s assay [15].

Development of indirect ELISA for mucin protein measurement

We initially used slot blot ELISA described in earlier publications [15] to measure MUC5AC and MUC5B production in culture media with pig gastric mucin and human salivary mucin as a standard for MUC5AC and MUC5B respectively. The initial MUC5AC experiments showed that MUC5AC content represent 98.8% of total basal mucin recovered from the culture media (results not shown). Due to small proportion of MUC5B produced in the culture media, the initial slot blot ELISA experiments were unsatisfactory. To improve MUC5B measurement sensitivity and reproducibility, we developed an indirect ELISA that involved the use of a 96-well plate (Maxisorp, NUNC). This also allowed a higher throughput of analysis. Furthermore, due to subsequent technical difficulties with the available MUC5AC primary antibody, we decided to use basal mucin produced by the culture media as a standard to measure the MUC5AC production by bile salt-stimulated cells. The standard MUC5AC curve was linear between 0.0 and 5.0 µg (Figure 2) which could be extended up to 10 µg after which saturation was seen and MUC5B standard curve was linear between 0.0 and 0.04 µg (Figure 3) which could be extended up to 0.08 µg. Standard curve linearity was lost after these levels due to saturation with the antibody (results not shown).

Quantification of MUC5AC and MUC5B content by indirect ELISA

MUC5AC and MUC5B mucin collected from stimulated HT29-MTX goblet cells were measured in duplicate. 100 µl of sample (diluted 1 in 10 and 1 in 20) or standards (0-5 µg/ml basal MUC5AC or 0-0.04 µg/ml human salivary mucin) made up in PBS were coated onto a 96-well plate, covered and incubated at room temperature overnight. The plate was then aspirated, blotted dry and blocked with blocking agent (1% casein and 1% BSA in PBS for MUC5AC and MUC5B respectively) for 2 hours. Then, 100 µl of primary antibody [monoclonal MUC5AC 45 M1 (abcam, UK), diluted 1 in 200 with 0.1% casein in PBS, for MUC5AC, and polyclonal antibody (Tepa II) diluted 1 in 2000 with 0.1% BSA in PBS for MUC5B] was added. The plate was covered and incubated for 1.5 hours and aspirated, washed with three changes of 0.05% Tween 20 in PBS followed by two changes of PBS and blotted dry. 100 µl of secondary antibody [anti-mouse (peroxidase conjugated) for MUC5AC and polyclonal peroxidase conjugated anti-rabbit (Sigma A667) for MUC5B, each diluted 1 in 5000] was then added to each well, the plate covered and incubated for 1.5 hours. The wash step was repeated and 100 µl of substrate solution 2,2’-Azino-bis 3-ethylbenzothiazoline-6-sulfonic acid (ABTS) added, the plate covered and developed for 20 minutes and the reaction stopped with 100 µl of 1% SDS. The absorbance was measured at 405 nm (Figure 3) with a plate reader (Bio-Tek EL808). Blank controls were made by omitting the primary antibody and incubating in antibody diluent instead.

Statistical Analysis

Statistical evaluations used One-Way ANOVA (Repeated Measures ANOVA, Dunn’s Multiple Comparison post test) and non-parametric unpaired tests (Spearman’s correlation test) with Prism (GraphPad Software Inc., La Jolla, California, USA). A p-value of 0.05 or less was considered statistically significant.

Results

The cultured HT29-MTX goblet cells remained viable through all the experiment times and there was no evidence of cell death at any of the challenge conditions. The cells produced both MUC5AC and MUC5B mucins. Basal MUC5AC was the predominant mucin compared to MUC5B by a factor of 80:1. Mean basal MUC5AC mucin secretion in the controls at all time points was 40±1 µg/ml whereas basal MUC5B secretion was 0.5±0.1 µg/ml at all time points. MUC5AC production was stimulated by bile salts in a bile salt concentration dependant manner. Maximal MUC5AC secretion, 2 fold of basal secretion, was reached at...
20 µmol/L bile salts. This basal mucin secretion levels was subtracted from the experimental bile salt stimulated levels.

MUC5AC upregulation occurred in the first 24 hours and was not further stimulated by extended incubation (Figure 4). At the 3 time points sampled, MUC5AC concentration remained at the same level, e.g. at 20 µmol/L bile salts concentration at 24h 1ml sample contained 50 ± 9 µg (mean ± SD) mucin above the basal level, at 48h 1 ml sample contained 50 ± 5 µg (mean ± SD) and at 72h 1 ml sample contained 50 ± 5 µg (mean ± SD). MUC5AC release had occurred in the first 24 hours and no more was released after that. MUC5B secretion remained at its basal levels of 0.5 µg/ml and was not altered by incubation with tested bile salts concentrations up to 20 µmol/L (results not shown).

Discussion

This study aimed to develop a physiological damage model models to investigate the potential of bile acids to stimulate mucus production from goblet cells. Damage models had been carried out in the past, and studies have looked into the damage potential of bile salts in animal models and ex vivo [16-20]. However, there were many problems with the investigation carried out with bile salts i.e. use of conditions not pertaining to in vivo [18,19], supra-physiological bile salt concentrations which would kill the tested cells [20-22], disregard of interspecies variability in bile composition and physiology and incomparable methodology (differing sampling methods, reliability of detection & quantification methods). These factors have to be considered to avoid errors in extrapolating results of animal experiments to man. The rat for example, does not have a gallbladder and levels of lithocholic acid are almost undetectable [21].

Previous work carried out in our lab [22] has shown that exposure of primary bronchial epithelial cells to bile salt challenges of 10 µmol/L or more was associated with significant cell injury up to cell death. However, in our current study we found that HT29-MTX cell viability was maintained at 100% when stimulated with bile salt concentrations up to 20 µmol/L. This indicates that the cell line HT29-MTX is more resistant than airway epithelial cells to bile salt challenge. This may however be a property of goblet cells compared to other epithelial cell types. The resistance could also be explained by the fact that these cells, being derived from an environment where bile and other gastric contents are common place, have an innate resistance to damage or stimulation by such components. It could also be that the HT29-MTX cells, being an immortalized human colon cancer cell line, act atypically not only in that they secrete airway specific mucins but also have higher resistance to bile acid challenge.

Lesuffleur et al. [23] have isolated MUC5AC from HT29 cell line whereas Gosalia et al. [24] have found that this cell line expresses MUC5AC at high levels with low levels of MUC5B. These studies indicate that the cell line HT29 can be used as a suitable model of airway goblet cells. We have found the same in our study with the majority of mucin produced made of MUC5AC and MUC5B expressed only at low levels. As it is difficult to grow primary lung goblet cells, we used the HT29 cell line stepwise challenged and adapted to increasing concentrations of methotrexate (HT29MTX) to obtain a homogeneous monolayer of polarized goblet cells that behaves like lung goblet cells. Therefore the HT29-MTX cells could provide an indication of the in vivo situation and act as a satisfactory goblet cell model although these cells are coming originally from colon carcinoma line and are not true lung goblet cells.

We looked at MUC5AC and MUC5B mucin production by the bile salt challenged cell line HT29-MTX as these are the 2 major mucins expressed by airway mucosa as concluded from our previous studies [14,15]. MUC5AC is an epithelial mucin mainly expressed in goblet cells of surface epithelium and to a lesser extent in the submucosal glands (SMG) whereas MUC5B is a glandular mucin mainly expressed in mucus cells of SMG and to a lesser extent in surface epithelium [9,10]. This was confirmed in our current study.

We investigated MUC5B production by this cell line to find out if this cell line behaves exclusively like goblet cells that produce mainly MUC5AC or behave also similar to the mucus secreting cells of submucosal glands which produce MUC5B as a major mucin. MUC5B production was minimal compared to MUC5AC and bile salt challenge failed to alter the MUC5B production which remained at its low basal level. This indicated that the cell line HT29-MTX behaves exclusively similar to airway goblet cells not to mucus secreting cells of submucosal glands. Further support of this comes from previous work [25] which had shown that challenging HT29-MTX cells with IL-6 and TNF-alpha did not significantly increase their production of MUC5B. However, as this is an in vitro study dealing with one type of mucus secreting airway cells, it is not known if the in vivo concomitant
presence of goblet cells and SMG could result in different mucin production in response to a bile salt challenge. A co-culture cell model needs to be developed to study the expression of this mucin under more physiologic and pathologic conditions.

Mucin expression response of this cell line to bile salt challenge seems to develop and reach its maximum within the first 24 hours and plateau afterwards. Therefore, for further bile salt stimulation experiments at these concentrations it would be important to analyse the media at multiple time points up to 24 hours to obtain more clear information as regard to the temporal upregulation of MUC5AC production. Furthermore, as the increase in MUC5AC production was close to linear up to 20 µmol/L bile salts and the cells were still 100% viable in those conditions, the concentration of bile salts could be increased beyond 20 µmol/L until viability decreases to test if and at which bile salt concentration MUC5AC production becomes maximal.

The upregulating effect of bile salts on mucin production in these cells could be due to an indirect effect of bile salts through the release of pro-inflammatory cytokines, such as tumour necrosis factor alpha, Interleukin (IL)-6 and IL-17 [26,27]. Bautista et al. [28] have shown that IL-8 regulates mucin gene expression in lung epithelial tissues in vitro, and previous work in our lab has shown that IL-8 up-regulates the secretion of MUC5AC from goblet cells in a concentration-dependent manner, with a maximum response at an IL-8 concentration of 20 ng/ml, with the response persisting for up to 5 days [29]. It had also shown that bacterial lipopolysaccharide at concentrations up to 100 µg/ml up-regulated the expression and secretion of MUC5AC (up to 40%) and IL-8 (up to 10-fold) from HT29-MTX goblet cells in a concentration-dependant manner [30].

Given the fact that mucus hypersecretion is a key element in airway inflammation and epithelial damage [31,32] and that reflux and potential bile salt aspiration has previously been documented in paediatric and adult populations [33,34], it would be plausible to treat bile reflux as a chronic contributor to lung injury in upper airway diseases [35].

Conclusion

We looked at mucin production as marker of bile salt damage in a physiological bile salt damage model using cultured HT29-PTX cells. These cells behaved similar to airway goblet cells producing MUC5AC as the main mucin and MUC5B only as a minor mucin. MUC5AC production was stimulated within the first 24 hours, proportionate to bile salt concentrations whereas MUC5B production was not affected.

Future Areas for Research

The damage models could be widely extended in that it can be investigated whether the goblet cell line HT29-PTX produces cytokines, both constitutively and after bile salt challenge. Different individual bile salts can also be tested. Moreover, a potential uptake of bile salts into the cells could be measured, as previously shown for pepsin in laryngeal epithelial cells through receptor-mediated Endocytosis [35].

References


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