Respiratory tract diseases stemming from inhalation exposure of toxic compounds significantly contribute to the global health burden. Traditional in vitro airway models, due to their lack of physiological relevance, are often unable to provide meaningful and accurate toxicological assessments. Advanced in vitro airway models, however, promise to provide more predictive information for use in human airway health. In this report, airway models comprised of fully differentiated primary human bronchial tracheal epithelial cells (HBECs) were generated to assess toxicological response to both short and long-term exposure from either cadmium chloride or pentamidine. Additionally, the toxicological responses from short-term exposure of compounds in airway models were compared to undifferentiated primary HBECs. Here we show that self-constructed airway models may serve as useful tools for future airway toxicity research.

**ABSTRACT:**

Respiratory tract diseases stemming from inhalation exposure of toxic compounds significantly contribute to the global health burden. Traditional in vitro airway models, due to their lack of physiological relevance, are often unable to provide meaningful and accurate toxicological assessments. Advanced in vitro airway models, however, promise to provide more predictive information for use in human airway health. In these studies, airway models comprised of fully differentiated primary human bronchial tracheal epithelial cells (HBECs) incubated in 24-well plate inserts and cultured under air-liquid interface (ALI) for 4 weeks were generated. The toxicological response to short-term (24 hours) exposure to either cadmium chloride (CdCl₂) or pentamidine were evaluated and compared in both airway models and undifferentiated HBECs. In addition, the toxicological response to long-term exposure (1, 2 weeks) to either compound in airway models was also explored. Changes in viability and cytokine expression were quantified and compared in both airway models and undifferentiated HBECs. Additionally, histological imaging (H&E, alcian blue, IHC) was conducted on airway models to visually assess model disruption, inflammation, and tight junction disruption. All airway models expressed dose-dependent response to both CdCl₂ and pentamidine exposure, with increased cell death corresponding with increased compound concentrations. Additionally, airway models demonstrated higher resistivity to cell death compared to undifferentiated counterparts. Moreover, exposure to low concentrations resulted in increased cytokine expression relative to untreated controls. Finally, long-term exposure to CdCl₂ resulted in model disruption and death, whereas pentamidine exposure demonstrated limited model disruption. These results suggest that airway models may serve as useful tools for future airway toxicity research.

**INTRODUCTION:**

The human respiratory tract is an important subject of study in a variety of fields. Respiratory infections represent the most common form of infection and act as a significant focus in disease research.¹,² Moreover, inhalation toxicity is the most prominent route of...
tory exposure, causing an estimated 7 million early deaths worldwide each year according to the World Health Organization. In pharmaceutical product development, inhalation serves as a viable route of therapeutic administration, bypassing the first pass metabolism as well as serving as a more acceptable route of administration for patients relative to intravenous or intradermal delivery. Providing relevant models able to recapitulate the human respiratory tract for these diverse scientific fields is of critical importance for global health.

To better serve the needs of human respiratory research, 3-D-airway models have been generated to better emulate the physiological complexity of the bronchial tracheal tract. We seeded primary human bronchial tracheal epithelial cells (HBECs) on microporous membrane supports that were incubated while partially exposed to air, known as air liquid interface (ALI). Following this incubation period, the HBECs fully differentiate to generate airway models containing goblet, ciliated, and basal cell lines, providing a physiologically relevant representation of the bronchial tracheal tract. In contrast, traditional in vitro 2-D-airway models, comprised of undifferentiated primary cells submerged in media, are limited in offering physiologically relevant data. The process of generating airway models, while lengthy compared to traditional models, can be easily performed, providing researchers with an affordable do-it-yourself model.

In the previous work “Evaluating airway ALI model fabrication methods and comparing differentiation potential of primary and hTERT-immortalized epithelial cells,” ATCC showcased an optimal method of fabricating airway models comprised of human primary HBECs with consistent full epithelial differentiation using multiple primary cell lots from both ATCC and other commercial vendors. In this report, ATCC investigated the toxicological response from both 3-D airway models comprised of fully differentiated primary HBECs as well as traditional 2-D in vitro models comprised of freshly seeded undifferentiated primary HBECs. Two different primary cell lots were used to generate both models and were subjected to both short and long-term exposure to either cadmium chloride (CdCl₂) or pentamidine. Cadmium, a metal utilized in industrial processes, is ranked 7th on the Agency for Toxic Substances and Disease Registry substance priority list as well as categorized as a substance of very high concern by EU’s European Chemical Agency due to its toxicity and risk of inhalation exposure. In contrast, pentamidine is a potent antimicrobial agent that can be administrated via inhalation and is listed in the World Health Organization list of essential medicines, however adverse side effects from its use are commonly reported. Both compounds were chosen to serve as representative substances in the fields of environmental/industrial monitoring and pharmaceutical safety. Here, various concentrations of these two compounds were utilized to compare changes in viability and inflammation from both undifferentiated cells and airway models. These studies reveal not only the differences in exposure response in both models, but also demonstrate the utility in using advanced 3-D airway models as useful tools future airway toxicity research.

**MATERIALS AND METHODS:**

**INITIAL CELL CULTURE**

Both lots of primary (ATCC® PCS-300-010™) HBECs were cultured according to ATCC’s product sheet recommendations. Briefly, all cells were first grown in ATCC complete airway epithelial growth media, consisting of ATCC Airway Epithelial Cell Basal Medium (ATCC® PCS-300-030™) supplemented with Bronchial Epithelial Cell Growth Kit (ATCC® PCS-300-040™) and 5 mL Penicillin-Streptomycin solution (ATCC® 30-2300™). During culturing and cell proliferation, the cell passage number for all cell lines did not exceed two. Cell growth media was replaced every other day.

For undifferentiated cell models, HBECs were passaged, and collected in growth media at a concentration of 50,000 cells/mL. 96-well plates were prepared, with each well having 0.2 mL of cell solution added (10,000 cells per well). To ensure cell attachment, plates were left at room temperature for at least one hour after seeding, followed by placement in 37°C incubation. The following day, plates underwent toxicity testing. To account for the edge effect, only the interior wells in the 96-well plates underwent testing, with the outer wells filled with 0.2 mL of Dulbecco’s Phosphate Buffered Saline (DPBS) (ATCC® 30-2200™).

To fabricate differentiated airway models, permeable trans-well inserts for 24-well plates with PET membrane with 0.4 µm pores (Corning cat# 353095) were placed into Costar clear 24-well multi-well plates (Corning cat# 3524) the day before cell seeding. To prevent edge effects on airway models, inserts were added to interior wells only, with outer wells filled with 2 mL DPBS. The plate inserts were then coated on the apical side with 0.3 mg/mL Collagen solution (Stemcell Technologies cat# 04902) diluted with DPBS, with plates incubating overnight at 4°C.

On the day of seeding, the collagen-coated inserts were apically rinsed twice with 200 µL DPBS, with 0.5 mL of ATCC complete airway epithelial growth media added to the basal side of wells containing trans-well inserts. HBECs were passaged, harvested, and then resuspended in growth media at a concentration of 500,000 cells/mL. The cell solution was added apically to each insert at a volume of 0.2 mL (100,000 cells per well). Following cell seeding, plates were left at room temperature for at least one hour to allow for cell attachment, followed by placement in 37°C incubation. Cells were incubated for 2-3 days until full confluency was reached. If confluency was not reached before 48 hours, both apical and basal side media were replaced with 0.2 and 0.5 mL of complete growth media respectively.

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AIRWAY MODEL ALI CULTURING

Once reaching full confluence, both the apical and basal media were removed from each airway model, with the basal media being replaced with 0.5 mL PneumaCult ALI maintenance media (Stemcell Technologies cat# 05001) supplemented with recommended components (Stemcell Technologies cat# 07980 and 07925). In contrast, no replacement media was added to the apical side of the models, partially exposing the cells to the air. Basal media was replaced every other day with fresh complete differentiation media. Cell differentiation was observed visually via microscopy, while weekly images from airway models were collected using an EVOS FL digital microscope (Life Sciences). Following two weeks of ALI incubation, weekly apical washes with (2x) 0.2 mL DPBS were conducted to remove excess mucin from models. To confirm proper airway model formation, TEER measurements were conducted using were collected using an EVOM2 Epithelial voltmeter (World Precision Instruments). After four weeks of ALI incubation and TEER confirmation, models were considered mature and ready for toxicology testing.

UNDIFFERENTIATED CELL TOXICOLOGY TESTING

Following 24 hours after initial cell seeding in 96 well plates, undifferentiated HBECs were subjected to 24-hour exposure to either CdCl₂ or pentamidine. Here, 16 different concentrations of CdCl₂ (Sigma Aldrich cat# 20298) and pentamidine (USP cat# 1504900) were prepared in DPBS, ranging from 2.5-400 µg/mL (14-2183 µM) or 5.4-2368 µg/mL (9.3-4,000 µM) respectively. Higher pentamidine concentrations required heating solution to 37˚C and vortexing vigorously to ensure complete solubility. Cell media was removed and replaced with 200 µL of respective compound concentration. Blank controls were prepared by administering 0.2mL DPBS to selected cells. Plates were placed in incubator for 24 hours following compound solution administration. Following 24 hours incubation, 100 µL media was collected to assess IL-8 cytokine expression. Quantification of IL-8 expression was conducted using Human IL-8 ELISA kit (Invitrogen cat# KHC0081) and following the provided instructions. Viability studies were conducted by adding 100 µL CellTiter-Glo 3-D Cell Assay Reagent (Promega cat# G968B) into each sample well. Plates were incubated at room temperature for 30 minutes, followed by pipette mixing and transferring 100 µL of sample into black sided 96-well plates. Samples in wells from the black sided plates were assessed via luminescence using a SpectraMax i3x plate reader ( Molecular Devices), with increased relative luminescence units (RLUs) corresponding to increased viability. All RLU values were corrected by subtracting the media only controls, followed by converting RLU values as percent viability relative to untreated controls [% Viability = 100 × (Sample RLU / Avg. Untreated RLU)]. Viability changes in cell models was expressed by calculating the half maximal inhibitory concentration (IC₅₀) using the corrected RLU values.

FULLY DIFFERENTIATED CELL MODEL TOXICOLOGY TESTING

Following four weeks of ALI, airway models were subjected to 24-hour exposure to either CdCl₂ or pentamidine, using the same concentrations administered to undifferentiated HBECs. For 1- and 2-week administration, a reduced number of concentrations were used, focusing on intermediate and lower compound administration. Prior to compound administration, basal media was replaced with fresh 0.5 mL fresh differentiation media, followed by with careful rinsing of the apical side using (2x) 0.2 mL DPBS. To approximate inhalation toxic exposure, 200 µL of respective compound solutions was administered on the apical side of the airway model. For short-term exposure, airway models were incubated 24 hours after administration, followed by sample processing. For both 1- and 2-week exposure, basal media was replaced every 48 hours, with apical media replaced weekly. For sample processing, basal media was removed, followed by 100 µL apical media collection for cytokine expression. Afterwards, 100 µL CellTiter-Glo 3-D Cell Assay Reagent was added to the apical side of the membrane. Following 30 minutes of room temperature incubation, 100 µL of cell solution was transferred to black sided 96-well plates for luminescence measurements.

For histological examination, separate sets of airway models undergoing similar conditions and exposure times were prepared. Following 24 hour, 1- or 2-week exposure time, airway models were fixated and permeabilized using BD cytofix/cytopermin fixation and permeabilization solution (BD cat# BD 554722). Samples were stored in 10% ethanol at 4˚C until further processing. Preserved models were paraffin embedded, sectioned, and stained with either H&E or alciain blue, followed by histological examination and imaging. Additional paraffin embedded samples underwent separate ICH staining and imaging using either Rabbit ZO-1 (Fisher Scientific cat# 40-2200) or Rabbit MUC5AC (Cell Signaling Technology Cat# 61193) antibodies.

RESULTS:

HBECs seeded in 24-well plate inserts were incubated under ALI conditions for a period of four weeks to induce epithelial differentiation. During incubation, microscopy images of airway models were taken weekly. Figure 1 illustrates the morphological changes the cells in airway models undertake during ALI, with obvious morphological changes occurring at week 2 (Fig 1C), followed by more pronounced changes occurring at week 3 (Fig 1D). Under these conditions, airway models undergo epithelial differentiation, resulting in mature airway models containing basal, goblet, and ciliated cells. These changes were observed in both primary HBEC lots as well as all airway model.
replicates. Moreover, TEER measurements demonstrated acceptable resistivity values and minimal variability between replicates (data not shown).

The first set of viability studies were conducted to assess short-term (24 hour) exposure from CdCl₂ on both undifferentiated HBECs and airway models. To visualize potential airway model disruption from CdCl₂ exposure, microscopy images of mature airway models were taken 24 hours following compound administration. Figure 2 illustrates the differences in mature airway model responses from DPBS treated (blank controls), low (53.9 µM), or intermediate concentrations (795 µM) of CdCl₂. Here, short-term exposure to low concentrations of CdCl₂ results in no observable disruption of model integrity relative to DPBS controls, while intermediate exposure to the compound shows widespread cell death, greatly disrupting airway model integrity.

In addition to microscopy imaging, changes in viability were assessed by generating IC₅₀ curves from both undifferentiated HBECs and airway models using two different primary HBEC lots (Figure 3). We observed that the undifferentiated airway models demonstrated a gradual and consistent dose-dependent decrease in viability with increased CdCl₂ administration (Fig. 3A), with both primary HBEC lots exhibiting IC₅₀ values of 87.5 µM ± 10.8 and 92.5 µM ± 10.8 respectively. In contrast, airway models demonstrated higher resistivity to cell death, maintaining nearly 100% viability up to 149 µM CdCl₂ exposure. However, the drop in viability in airway models was more pronounced, relative to undifferentiated HBECs (Fig. 3B). The IC₅₀ values of airway models lots 1 and 2 were calculated to be 203.1 µM ± 7.2 and 273.7 µM ± 12.3 respectively, a 2-fold increase in IC₅₀ values relative to untreated HBECs (Fig. 3C-D).

Viability measurements from short-term pentamidine exposure was also assessed. Figure 4 illustrates both the changes in viability as well as generated IC₅₀ curves from both undifferentiated HBECs and airway models following 24-hour pentamidine administration. Similar to results from CdCl₂ administration, both lots of undifferentiated primary HBECs demonstrate dose-dependent decrease in viability with increased pentamidine administration, with lots 1 and 2 exhibiting IC₅₀ values of 60.4 µM ± 5.5 and 57.1 µM ± 13.5 respectively (Fig. 4A). However, airway models exhibited significantly higher resistance to cell death relative to undifferentiated HBECs, maintaining nearly 100% viability at pentamidine concentrations 1185 µM, as well as maintaining an average viability over 10% even at the highest exposure concentration (Fig. 4B). Here airway model lots 1 and 2 demonstrated IC₅₀ values of 2,811 µM ± 201 and 2,279 µM ± 113 respectively, with IC₅₀ values averaging over 40-fold higher than undifferentiated primary HBEC counterparts (Fig. 4C-D).

To better visualize changes in airway model structure from short-term CdCl₂ exposure, histological analysis on separate sets of exposed airway models was conducted. Figure 5 shows representative images of alcian blue stained airway models incubated in either 0 (DPBS blank controls), 53.9 (low), 147.9 (intermediate), or 2183.4 µM (high) CdCl₂ for 24 hours. DPBS blank controls demonstrated appropriate model morphology, showing the presence of goblet and ciliated cells (Fig. 5A). Short-term exposure to low concentrations of CdCl₂ resulted in no noticeable differences in model morphology (Fig. 5B). In contrast, intermediary CdCl₂ exposure clearly results in model deterioration, with decreased model thickness and absence of cells on the membrane support (Fig. 5C). Finally, higher levels of CdCl₂ exposure resulted in the complete degradation of the airway model, with one or two cells present on the entire membrane (Fig. 5D).

Histological analysis was also conducted on airway models exposed to short-term pentamidine administration. Figure 6 illustrates representative images of alcian blue stained airway models incubated in either 0 (DPBS blank controls), 46.2 (low), 1,185 (intermediate), or 4,000 µM (high) pentamidine for 24 hours. Unlike CdCl₂ treated counterparts, airway models demonstrated greater resistance in model degradation from short-term exposure to pentamidine, with both low and intermediate concentrations showing no observable difference in morphology relative to DPBS treated blank controls. However, slight model disruption was only observed in the highest treated (4,000 µM) pentamidine sample (Fig. 6D).

In addition to assessing changes in viability from short-term compound exposure, cytokine expression was also assessed to better examine toxicological response from both compounds. Figure 7 shows the changes in IL-8 expression in both undifferentiated HBECs and airway models from 24-hour exposure from to CdCl₂. In both undifferentiated HBECs and airway models, low to intermediate administration of CdCl₂ results in increased expression of the proinflammatory cytokine, compared to untreated blank controls. In contrast, models subjected to high CdCl₂ administration exhibited cytokine expression levels similar to or lower than blank controls, which was attributed to cell death. In the pentamidine administered samples, high pentamidine administration resulted in little to no detectable IL-8 in both airway models and undifferentiated HBECs, whereas samples administered with low pentamidine concentrations showed lowered cytokine expression relative to blank controls (data not shown).

Following the conclusion of the short-term viability studies, airway models were subjected to one- or two-week exposure to either CdCl₂ or pentamidine. Figure 8 displays the changes in viability and cytokine expression from continuous exposure to either compound. After 1-week of CdCl₂ exposure, both airway model lots exhibited slightly elevated viability measurements from low exposure concentrations relative to untreated controls, probably as a function of wound healing response. In contrast, moderate long-term CdCl₂ administration resulted in significant decrease in airway model viability. Additionally, 2-week exposure results exhibited similar trends, with low CdCl₂ concentrations demonstrating near equivalent viability response relative to untreated controls (Fig. 8A). Viability measurements taken
from long-term pentamidine administration resulted in similar trends to CdCl₂ exposure, with low pentamidine concentrations demonstrating comparable viability response to blank controls whereas long-term moderate pentamidine exposure resulted in decrease viability (Fig. 8B). Cytokine expression was also assessed using apical washes collected at the end of compound administration (Fig. 9C). Similar to short-term testing, airway models demonstrated elevated levels of cytokine expression to low cadmium administration following 1 week incubation relative to untreated controls, whereas moderate cadmium administration results in low cytokine expression, attributed to cell death. There was no observable trend between increased cytokine expression to increased compound administration following 2-week administration, which was attributed to suboptimal cytokine sample collection from long-term studies (data not shown).

Parallel with long-term viability measurements, histological assessments were also conducted on separate airway model samples. Figure 9 shows representative images of alcian blue stained airway models treated with either low to moderate administrations of either CdCl₂ or pentamidine for one week. Increasing CdCl₂ administration during this period results in progressively deteriorating airway model integrity. Pentamidine exposure showed a similar, albeit a more gradual trend, with model integrity being better preserved from long-term pentamidine exposure. These trends were observed in two week administered samples as well (data not shown).

Finally, separate IHC staining and imaging was conducted to visualize potential ZO-1 protein disruption from long-term CdCl₂ exposure as well as increased MUC5AC expression short-term pentamidine administration (Figure 10). Representative IHC-stained images of airway models stained for ZO-1 protein demonstrated that decreased ZO-1 expression is correlated to increased CdCl₂ administration, relative to blank controls (Fig. 10A-C). In contrast, no observable correlation was found between with increased MUC5AC expression from increasing pentamidine administration (Fig. 10D-F). This result was attributed to the preparation of histology samples, which may have resulted in the washing and removal any excess mucin from the airway model samples.

DISCUSSION:

In these studies, self-constructed airway models comprised of fully differentiated HBECs were fabricated for use in toxicological studies. Here, the toxicological response between airway models and freshly seeded undifferentiated HBECs from short-term administration of two different compounds were compared. Moreover, this study also examined the variation between models comprised of two separate primary cell production lots. Finally, long-term exposure studies of airway models were also explored.

During these studies, compounds were administered to the apical side of airway models to better approximate the physiologically relevant exposure of the human lung to toxic compounds. We observed that airway models subjected to short-term CdCl₂ or pentamidine exposure exhibited average IC₅₀ values of 238 and 2545 µM respectively. To our knowledge, no relevant study has yet been conducted to assess changes in viability in airway models from 24-hour exposure to pentamidine. In contrast, a previous study testing airway models to short-term CdCl₂ exposure estimated IC₅₀ to be within a range 100-300 µM, in agreement to our findings.¹⁵ However, the referenced study administered CdCl₂ on the basal side. This difference may not affect airway models comprised of only HBECs; however, co-culture models may not respond in a physiologically relevant manner to compound administration through basal media.

Short-term cytokine results aligned with our expectations, with low-moderate CdCl₂ administration results in higher cytokine expression relative to blank controls, whereas high compound fails to elicit higher cytokine expression due to greater cell death. In contrast, pentamidine administration failed to induce increased cytokine expression. Previous in vitro studies showed that pentamidine exposure inhibits IL-8 expression, which was observed in this study as well.¹⁶ During long-term testing no trend was observed during 2-week incubation, which was attributed to cytokine sample loss when basal media was replaced during the study. Although basal media replacement is vital in sustaining airway model viability, any media or solution incubated with airway models should be collected and assessed during the study. Long-term viability measurements were unaffected to this incident due to directly assessing primary cell metabolism.

These studies showcase how self-fabricated airway models can be utilized to provide data that’s more physiologically relevant to the human airway model. This was clearly demonstrated when comparing the response from both undifferentiated HBECs and airway models from pentamidine administration. Moreover, the seeding of additional cells, such as fibroblasts, dendritic, or endothelial cells are possible using this airway model fabrication method and may offer additional insights utility in toxicology studies.

ACKNOWLEDGEMENTS:

Histological processing and imaging were conducted by VitroVivo Biotech LLC.
**FIGURES AND TABLES:**

**Figure 1:** Representative microscopy images of primary bronchial epithelial cells under (A) 0, (B) 1, (C) 2 and (D) 3 weeks of ALI. Prolonged incubation under ALI conditions induces fully epithelial differentiation in airway models. Scale bars represent 400 µm.
Figure 2: Representative microscopy images of airway models apically treated with either (A) 0, (B) 53.9 or (C) 795.6 µM CdCl₂ for 24 Hr. No observable differences were seen between airway model blank controls and models administrated with low concentrations of CdCl₂. In contrast, intermediate CdCl₂ exposure results in high levels of cell death and model disruption (floating black spots). Scale bars represent 400 µm.
**Figure 3:** Changes in viability between (A) undifferentiated HBECs and (B) airway models from 24-hour exposure to CdCl₂. Both undifferentiated HBECs and differentiated airway models were assessed by generating (C) IC₅₀ curves and (D) comparing IC₅₀ values between the two groups. Airway models demonstrated higher resistivity to cell death and IC₅₀ values, relative to undifferentiated HBECs.

**Table D:**

<table>
<thead>
<tr>
<th>Sample Name</th>
<th>IC₅₀ Value (µM)</th>
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<tr>
<td>Undifferentiated Lot 1</td>
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</tr>
<tr>
<td>Undifferentiated Lot 2</td>
<td>92.49 ± 10.8</td>
</tr>
<tr>
<td>Differentiated Lot 1</td>
<td>203.1 ± 7.2</td>
</tr>
<tr>
<td>Differentiated Lot 2</td>
<td>273.7 ± 12.3</td>
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Figure 4: Changes in viability between (A) undifferentiated HBECs and (B) airway models from 24-hour exposure to pentamidine. Both undifferentiated HBECs and differentiated airway models were assessed by generating (C) IC₅₀ curves and (D) comparing IC₅₀ values between the two groups. Airway models demonstrated significantly higher resistance to cell death and as well as over 40-fold higher IC₅₀ values, relative to undifferentiated HBECs.
Figure 5: Representative alcian blue stained images of airway models apically treated with 24-hour exposure to either (A) DPBS blank controls, (B) 53.9, (C) 147.9, or (D) 2183.4 μM CdCl₂. Increased CdCl₂ administration results in greater airway model disruption and cell death. Scale bars represent 20 μm.
Figure 6: Representative alcian blue stained images of airway models apically treated with either (A) DPBS blank controls, (B) 46.2, (C) 1,185, or (D) 4,000 µM pentamidine for 24 Hr. Compared to CdCl₂ counterparts, airway models demonstrated no observable model degradation from short-term pentamidine exposure, except for the highest administered dosage of 4,000 µM, which exhibited moderate model disintegration. Scale bars represent 20 µm.
Figure 7: Analysis of IL-8 cytokine expression from 24-hour administration of CdCl₂ to both undifferentiated HBECs and mature airway models. Values are shown as blank subtracted. Low concentrations of CdCl₂ results in increased proinflammatory cytokine expression, relative to blank controls. In contrast, intermediate to high administration, results in decreased expression, due to cell death.
Figure 8: Changes in viability in airway models exposed to either (A) CdCl₂ or (B) pentamidine for 1 and 2 weeks, (C) as well as cytokine expression following 1 week administration of CdCl₂.
Figure 9: Representative alcian blue stained images of airway models apically treated with either (A) 0, (B) 14.0, (C) 53.9, or (D) 795.6 µM CdCl₂, or (E) 0, (F) 46.2, (G) 156, (H) 1185 µM pentamidine for 1 week. Long-term exposure from low or moderate concentrations of CdCl₂ results in airway model disruption. In contrast, airway model integrity is better preserved from long-term pentamidine exposure. Scale bars represent 20 µm.
**Figure 10:** Representative ICH-stained images of (A) untreated control airway models, airway models exposed to (B) 14.0, or (C) 147.9 µM CdCl₂ for 2 weeks. ZO-1 protein expression is shown as red with DAPI control in blue. ZO-1 protein disruption is associated with increased CdCl₂ exposure. Representative ICH-stained images of airway models treated with (D) 0, (E) 9.1, (F) 46 µM pentamidine for 24 Hr. MUC5AC expression shown as red with DAPI control in blue. There was no correlation with increased MUC5AC expression with increasing pentamidine administration.
REFERENCES:


