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Chronic Pepsin Exposure Promotes Anchorage-Independent Growth and Migration of a Hypopharyngeal Squamous Cell Line

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Sponsorships or competing interests that may be relevant to content are disclosed at the end of this article.

Abstract

Outcome Objectives. (1) Investigate the role of reflux, specifically pepsin, in laryngopharyngeal carcinogenesis. (2) Evaluate effects of chronic pepsin exposure on cell migration, apoptosis, and colony-forming ability in hypopharyngeal cells.

Study Design. Translation research.

Setting. Academic research laboratory.

Methods. Human hypopharyngeal squamous carcinoma FaDu cells were chronically exposed to nonacidic pepsin (exposed for 24 hours, 4 times over 2 weeks at the following concentrations: 0.01 mg/mL, 0.1 mg/mL, or 1 mg/mL). Precise wounds were created in confluent cell plates, and rates of cell migration into wounds were quantified. Separately, cell viability of chronic pepsin-exposed FaDu cells acutely treated with paclitaxel was measured. Finally, a clonogenic assay was performed on these cells to measure effects of chronic pepsin exposure on colony-forming ability.

Results. An increased rate of relative wound density was observed in chronic pepsin-treated (0.01 mg/mL, 0.1 mg/mL) cells compared with control (P < .001), suggesting greater rates of cell migration. Pepsin-treated (0.1 mg/mL) cells demonstrated on average greater cell viability compared with control after exposure to paclitaxel, suggesting possible apoptotic resistance; however, this was not statistically significant. Chronic pepsin exposure (0.1 mg/mL, 1 mg/mL) was associated with a dose-dependent increase in colony-forming ability relative to control (P < .001).

Conclusion. Hypopharyngeal squamous cell line chronically exposed to pepsin demonstrated increased cell migration and colony-forming ability relative to control cells. These experiments indicate that chronic pepsin exposure acts as a promoter of tumorigenesis and metastasis of airway epithelium, suggesting a role for pepsin in laryngopharyngeal carcinogenesis attributed to gastric reflux.
Initial investigations to identify the causative agents of LPR-attributed disease evaluated the effects of gastric acid on the mucosal lining of the upper aerodigestive tract. However, injury to the upper aerodigestive tract was seen despite proton pump inhibitor therapy, and esophageal and laryngeal injury was found to occur even in nonacidic conditions. Further studies found that pepsin played a significant role in laryngeal injury despite nonacidic conditions. Recent evidence has shown that, at neutral pH, pepsin is taken up by laryngeal and hypopharyngeal epithelial cells by receptor-mediated endocytosis and retained in the low-pH environment of intracellular vesicles such as endosomes and Golgi. This retention of pepsin in a low-pH environment would allow pepsin to restore its proteolytic activity and potentially cause further mucosal injury.

Clinical studies have evaluated the incidence of LPR in patients with laryngopharyngeal cancer. Some have demonstrated a higher rate of LPR in nonsmokers and nondrinkers with laryngeal cancer. While a correlation between LPR and laryngeal cancer has been demonstrated in prior studies, causality has been difficult to prove. Specifically, identifying LPR as an independent risk factor for laryngeal cancer has been difficult, as these investigations are often confounded with patients with prior tobacco or alcohol dependence. In addition, these research studies and clinicians use different methods to measure and diagnose LPR. Currently, a causal role of reflux in laryngeal cancer has not been adequately demonstrated. Understanding the molecular effects of chronic pepsin exposure in the laryngopharynx may provide further support for its role in carcinogenesis.

We previously looked at 84 genes implicated in carcinogenesis and evaluated the effects of acute pepsin exposure on expression of these genes in human hypopharyngeal squamous cell carcinoma (SCC) FaDu cells. Three genes were noted to have increased expression by more than 1.5-fold, and 24 genes, including many tumor suppressors, demonstrated reduced expression by more than 1.5-fold. In addition, growth curve assays showed that acute pepsin exposure increased the percentage of cells in the replicative S phase in both human hypopharyngeal SCC FaDu cells and primary laryngeal epithelial cells. The pepsin-induced increase in cell proliferation exhibited both a dose- and time-dependent response, suggesting that acute pepsin exposure increases cell proliferation and may cause aberrant cell growth. The effects of chronic pepsin exposure were also analyzed in a study conducted by Allen et al using a hamster buccal pouch model to demonstrate the role of pepsin as a cofactor in tumor growth. The study showed that co-treatment of the hamster buccal pouch with 7,12-dimethylbenzanthracene (DMBA), a known carcinogen, and acidified pepsin (3 times daily for 14 weeks) generated significantly larger tumors than DMBA alone, thereby demonstrating a potential role for pepsin in malignant transformation. Cumulatively, these data reveal that pepsin exposure increases cell proliferation and alters expression of genes involved in carcinogenesis, suggesting that uncontrolled chronic reflux of pepsin could cause unregulated cell growth and malignancy.

We hypothesized that reflux of nonacidic pepsin contributes to oncogenic transformation by acting as a tumor promoter. To investigate the role of reflux, specifically pepsin, in laryngopharyngeal cancer, the effects of chronic pepsin exposure on cell migration, apoptosis, and anchorage-independent growth were analyzed in a human hypopharyngeal SCC FaDu cell line. A FaDu cell line was used as these cells are technically optimal cells for this initial investigation and would provide information regarding the impact of pepsin on promoting tumor growth, which is one step in oncogenic transformation.

Materials and Methods

Cell Culture

Human hypopharyngeal SCC FaDu cells (ATCC, Manassas, VA) were cultured in Minimum Essential Medium (ATCC) supplemented with 10% fetal bovine serum (ATCC) and 1% antibiotic/antimycotic (Life Technologies, Grand Island, NY). When exposing these cells to pepsin, porcine pepsin was used, as it is commercially available and has similar activity as human pepsin. Pepsin is at a concentration of 1 mg/mL in the stomach. It presumably is diluted in aerodigestive secretions as the reflux travels proximally to the laryngopharynx. Physiological concentrations were used ranging from 0.01 mg/mL to 1 mg/mL.

Cell Migration Assay

Hypopharyngeal SCC FaDu cells were grown to 100% confluence on a 96-well Essen BioScience ImageLock Plate (Essen Bioscience, Ann Arbor, MI) at 37°C with 5% CO2. The cells were exposed to porcine pepsin (0.01 or 0.1 mg/mL; Sigma-Aldrich, St Louis, MO) at pH 7 or with media at pH 7 alone (control) for 24 hours in 6 technical replicates. Wells were washed with phosphate-buffered solution, and fresh growth media were replaced. Precise wounds were made in each well using a WoundMaker (Essen Bioscience), a tool designed to make a wound with consistent size and decreased variability among each well. The plate was placed in an IncuCyte incubator, a live-cell imaging system (Essen Bioscience), and images were collected at 1-hour time intervals.

Images of cell migration within each well were obtained while in the Incucyte, an incubator that provides real-time images of cells to document migration during the incubation process. The Incucyte software uses the parameter of relative wound density (RWD) as the most robust metric to quantify cell migration. At each time point, the RWD, which corresponds to the spatial cell density in the wound area relative to the spatial cell density outside the wound area, was calculated by the Incucyte software RWD algorithm. The RWD and its rate of change were compared between control and pepsin-treated conditions. In addition, the area under the curve was calculated for each condition and used to determine statistically significant differences.

Apoptosis Resistance Assay

This assay was performed to assess the effects of pepsin on disruption of the apoptotic pathway and cell viability.
Human hypopharyngeal SCC FaDu cells were treated for 24 hours with porcine pepsin (0.01, 0.1, or 1 mg/mL) at pH 7 or with media at pH 7 alone (control) 2 times over a 1-week time period. There were 3 technical replicates for each condition. Cells were maintained at <90% confluence and allowed to rest at least 24 hours between each treatment. Cells were then treated with 50 μM paclitaxel (Sigma-Aldrich), an apoptosis-promoting agent, or media alone (control) and incubated for 12 hours at 37°C with 5% CO2. Viable cells that were observed after treatment with paclitaxel were then counted. Poisson regression with overdispersion adjustment was performed to determine statistical difference among the conditions.

**Clonogenic Assay**

The clonogenic assay was performed to assess the effect of pepsin on anchorage-independent growth and colony-forming ability. Often, cells need a solid surface to attach to for further proliferation, and normal cells fail to grow when suspended in viscous fluid or gel (agar). As this assay is performed using a previously transformed cell line, comparison of colony growth in each condition would provide the ability of pepsin exposure to promote tumor growth. Prior to the assay, hypopharyngeal SCC FaDu cells were treated for 24 hours with porcine pepsin (0.1 or 1 mg/mL; Sigma-Aldrich) or with media at pH 7 (control) in replicates of 6, 4 times over a 2-week time period. The concentration of 0.01 mg/mL pepsin was not used, as prior investigation evaluating the impact of pepsin exposure on cell proliferation did not demonstrate a statistically significant difference at this concentration. Cells were maintained at <90% confluence and allowed to rest at least 24 hours between each treatment. Bottom agar was prepared with 0.5% agarose melted and added to 50% normal growth media. One-milliliter aliquots of the bottom agar were placed into the 6-well plates. A total of 2500 cells of each of the replicates for the 4 conditions were mixed with 500 μL of 0.3% agarose in 75% normal growth media and overlayed onto solidified bottom agar. The bottom agar served as a barrier from the solid plate and the cells suspended in the top agar. Therefore, cells that possessed only anchorage independence would grow. Cells were placed in an incubator at 37°C with 5% CO2 for 2 weeks. One hundred microliters of growth media was added to each well as needed during the incubation period. After 2 weeks of incubation, colonies were noted in the agar plates. Wells were stained with 0.005% crystal violet and washed with sterile water. Images were obtained of each well and condition (Photo DocIt Imaging System, UVP, Upland, CA). Colony number was assessed using Image J 1.47 software (imagej.nih.gov/ij/download/). The data obtained were the mean of 6 wells for each condition. Statistically significant differences were determined.

**Statistical Methodology**

Poisson regression with overdispersion adjustment was performed on the apoptosis resistance assay and the clonogenic assay. For the data from the cell migration assay, the area under the curve of the RWD (%) over 24 hours was calculated for the curve for each sample, then a linear model was fit to examine the differences of the 3 types of samples. P values were adjusted for multiple comparisons by the Tukey-Kramer method. The significance level was set at .05. All data analyses were carried out using the Statistical Analysis System, version 9.2 (SAS Institute, Cary, NC).

**Results**

**Cell Migration Assay**

The ability of nonacidic pepsin to alter cell migration was assessed by measuring the RWD over time in the wound-healing assay (Figures 1 and 2). The maximal difference in RWD was detected at 24 hours postwounding; the control group had a relative wound density of 49% at 24 hours, and cells exposed to 0.01 mg/mL pepsin and 0.1 mg/mL pepsin had an RWD of 62% and 82% at 24 hours, respectively. The area under the curve of RWD was calculated for each of the groups (control, 0.01 mg/mL pepsin, and 0.1 mg/mL pepsin). There was a significant difference when comparing the area under the curve of RWD of the 3 groups, P < .001. It was highest in the 0.1 mg/mL pepsin group and lowest in...
the control group. There were significant differences observed when comparing any of the 2 groups among the 3 ($P < .01$). No significant differences were noted in cell morphology among the 3 different conditions.

**Apoptotic Resistance Assay**

FaDu cells were treated acutely with an apoptosis-promoting agent, paclitaxel, after chronic exposure to pepsin at 3 different concentrations to assess the effects of pepsin on apoptosis. The mean number of viable cells counted following paclitaxel treatment of the 0.01 mg/mL pepsin treatment condition was 1.1-fold higher than the mean number of viable cells counted in the control condition not treated with pepsin but treated with paclitaxel. The mean number of viable cells counted following paclitaxel of the 0.1 mg/mL pepsin treatment condition was 1.75-fold higher than the mean number of viable cells counted in the control condition. The mean number of viable cells of the 1 mg/mL pepsin treatment condition that were counted following paclitaxel was 1.48-fold higher than the mean number of viable cells counted in the control condition. Results are displayed in Figure 3. There was no statistically significant difference between the 4 conditions (control, 0.01 mg/mL pepsin, 0.1 mg/mL pepsin, 1 mg/mL pepsin; $P = .078$). Moreover, it was not significantly different for the pairwise comparison of any of the 2 groups.

**Clonogenic Assay**

Colony formation was noted on pepsin-treated plates after 2 weeks of incubation, with a unique morphology compared with controls (Figures 4 and 5). Soft agar clonogenic assay shows increased colony formation in those FaDu cells exposed to pepsin. The mean of 6 replicates demonstrated that 11 colonies formed in control wells, 293 colonies in those cells treated with 0.1 mg/mL pepsin and 305 colonies in those cells treated with 1 mg/mL pepsin. A greater than 25-fold increase in colony number was noted in those wells containing cells that were exposed to chronic pepsin treatment compared with control/untreated wells, as displayed in Figure 6. There was a significant difference among the 3 conditions ($P < .001$). Among the 3 conditions, the control group was significantly different from the 0.1 mg/mL pepsin and 1 mg/mL pepsin groups, with a $P$ value <.001. However, 0.1 mg/mL pepsin was not significantly different from the 1 mg/mL pepsin group ($P = .972$).

**Discussion**

An association between laryngopharyngeal reflux and laryngeal cancer has been demonstrated in a number of previous studies. Demonstration of the ability of pepsin to cause oncogenic transformation would lend further support to the role of reflux in carcinogenesis. While our previous work focused on the effect of acute pepsin exposure on cell proliferation, the hamster buccal model used by Allen and colleagues suggested a role for chronic pepsin exposure in oncogenic transformation. The study reported herein corroborates previous findings of the tumorigenic potential of chronic pepsin exposure in the upper airway and suggests a role for pepsin in metastasis.

The results of the wound-healing assay confirmed that chronic nonacidic pepsin exposure dose dependently induced an increase in cell migration, a process that is imperative in a number of biologic pathways including chronic inflammation and tumor metastasis. Our previous work has shown that pepsin treatment induces altered expression of metastasis-related genes including extracellular matrix genes and those involved in cell adhesion, cell cycle, growth, proliferation, and apoptosis. In vivo evidence similarly supports a role for LPR in promotion of metastasis. Several authors studied the effects of refluxate on cell adhesion proteins and identified a decline in E-cadherin expression in laryngeal biopsies of patients with documented LPR. E-cadherin provides a vital structural role in cells of tissue/organ and,
with loss of expression of this protein, may decrease the adhesive function of cells and increase the possibility of tumor cell invasion and metastasis. The 5-year survival rate for laryngeal cancer is poor for advanced-stage disease. While trends in tobacco use have declined over the past few decades, the survival rates have continued to decline, specifically in patients with regional and distant disease, suggesting the influence of other factors in this disease process. Advances in our understanding of the factors that contribute to laryngeal SCC could be of fundamental importance in reducing the frequency of advanced-stage disease.

Our prior investigation of the influence of an acute pepsin exposure on gene expression using the Cancer PathwayFinder SuperArray demonstrated a reduced expression of proapoptotic genes. Accordingly, an antiapoptotic effect of chronic pepsin exposure was evaluated in our apoptotic resistance assay. These results were obtained to evaluate the effect of pepsin on the balance between cell growth and apoptosis and potential for unregulated cell growth. There were more viable cells that were repeatedly exposed to pepsin after treatment with the apoptosis-promoting agent, paclitaxel, compared with controls; however, this was not significant in our analysis.

In addition, a clonogenic assay was performed to measure the capacity of pepsin to induce anchorage independence and loss of contact inhibition. A common feature of carcinoma development and growth is the ability of transformed cells to survive under anchorage independence. The soft agar assay for colony formation, an anchorage-independent growth assay in soft agar, is considered the most stringent assay for detecting malignant transformation of cells. As expected, there were colonies noted on the control plates, as the FaDu cells used are a tumorogenic SCC-derived cell line. However, FaDu cells demonstrated significantly greater colony-forming

![Figure 4](image1.png)  
**Figure 4.** Effect of pepsin on anchorage-independent cell growth. Images of crystal violet–stained colonies of human hypopharyngeal FaDu cells chronically treated with pepsin (pH 7) or pH 7 media alone (control) grown in soft agar plates for 2 weeks. Magnification: 0.

![Figure 5](image2.png)  
**Figure 5.** Images of hypopharyngeal squamous cell carcinoma FaDu cells in soft agar assay. Magnification: 100×. (A) Control cells. (B) Treated cells with a change in morphology relative to controls. (C) Treated cells.

![Figure 6](image3.png)  
**Figure 6.** Cell clonogenic assay results. FaDu cells treated with porcine pepsin (0.1 mg/mL and 1 mg/mL) had a statistically significant increase in colony-forming ability compared with untreated cells. Error bars represent standard deviation.
ability after chronic pepsin exposure, suggesting that pepsin promotes anchorage independence. This role of chronic pepsin exposure as a tumor promoter in a hypopharyngeal squamous cell line further supports the role the pepsin in carcinogenesis of the laryngopharynx.

There were several limitations with this study. One limitation includes the use of an in vitro cell line. While in vitro experiments provide obvious benefits including simplicity and ease, they do not completely re-create the complex in vivo environment. In addition, the cell line used was an immortal, tumor-derived cell line. Further investigation using primary laryngeal epithelial cells (currently ongoing in our laboratory) and animal models will be needed to validate the role of pepsin in laryngeal carcinogenesis attributed to LPR and the potential utility of pepsin inhibitors in the prevention of LPR-attributed laryngeal cancer.

Conclusion

Pepsin in laryngopharyngeal reflux continues to be investigated as a contributor to the development of laryngopharyngeal carcinogenesis. In the study described herein, chronic pepsin exposure elicited increased cell proliferation and migration, anchorage-independent cell growth, and increased colony formation in oncogenetically transformed human hypopharyngeal cells, lending further support for the role of pepsin as an agent of oncogenic transformation.

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Author Contributions

Elizabeth A. Kelly, substantial contributions to acquisition of data, analysis and interpretations of data, drafted manuscript, provided further revision of article and final approval of version published; Tina L. Samuels, substantial contributions to conceptions and design, analysis and interpretation of data, provided critical revision of draft and final approval of version to be published; Nikki Johnston, substantial contributions to conception and design, analysis and interpretation of data, provided critical revision of draft and final approval of version to be published.

Disclosures

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