Experimental model of tracheal stenosis with submucosal resection of cartilaginous rings combined with sodium hydroxide instillations

Abstract

Objective: To experimentally develop tracheal stenosis and malacia to test new models of tracheal stents. Methods: Was resected three cartilaginous rings from the cervical trachea of dogs in group A (n = 5) and six rings in group B (n = 4) to produce malacia. The mucosa of the region with malacia then received applications of a solution of sodium hydroxide (NaOH) at 23%. Results: In histological analysis, fibrosis was found in the submucosa and adventitia, associated with granulomas in the mucosa. The average residual lumen of the segments with stenosis was 9% and 12% in groups A and B, respectively (p >0.05). Conclusion: The combination of resection of the cartilaginous rings and the application of 23% NaOH in the respiratory mucosa promoted severe tracheal stenosis, but was associated with loss of animals. Further studies are needed to verify that the isolated use of one of the techniques would be safer and more effective to develop tracheal stenosis.

Key words: Trachea. Tracheomalacia. Tracheal stenosis. Stents. Implants, experimental.

Introduction

Tracheal inflammatory stenosis and malacia are a consequence of trauma and ischemia of the respiratory mucosa, produced by the endotracheal tube and/or the excessively inflated cuff of the tube or tracheostomy tube. The result of this aggression is inflammation and fibrosis, which may be severe and compromise the entire tracheal wall, leading not only the decrease in airway lumen, but also the loss of its cartilaginous support and tracheal collapse, resulting in malacia. Surgical resection with end-to-end reconstruction (tracheoplasty) is the treatment of choice for inflammatory tracheal stenosis. On the other hand, silicone tracheal stents have become a therapeutic option for patients with increased surgical risk or too long strictures.

Currently there is no ideal model of a tracheal stent, thus several have been developed. In our setting, the high cost of imported stents prompted the authors to develop new models of silicone tracheal stents, such as the HCPA-1. In this sense, we seek an animal model of tracheal stenosis, based on published experimental studies, which mimics the pathophysiological conditions found in humans to evaluate our silicone stent. Thus, the purpose of this study is to develop an experimental model of inflammatory tracheal stenosis.

Methods

Was used nine dogs of both sexes and mixed breed, weighing between eight and 17 kg. The animals included in this study meet the criteria established by the Health Monitoring Authority, supervised by the Veterinary Service of the Animal Control Kennel Hall of Porto Alegre and the Faculty of Veterinary Medicine, Universidade Federal do Rio Grande do Sul (UFRGS), Porto Alegre, Rio Grande do Sul – RS, Brazil.

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Experimental model of tracheal stenosis with submucosal resection of cartilaginous rings combined with sodium hydroxide instillations

The experiment was conducted under the supervision of the Ethics Committee in Research of Hospital de Clínicas de Porto Alegre, which approved it under number 00-314 and 02-100. All animals received humane care in accordance with the “Guide for Care and Use of Laboratory Animals” published in “National Institutes of Health (NIH 1985:85-23, revised).” The study received financial incentive of FIPE / HCPA (Fund for the Encouragement of Research and Events of the Hospital de Clínicas de Porto Alegre – file number 00-314).

Animals with food and water fasting for eight hours underwent bronchoscopic and surgical procedures under general anesthesia. They received atropine sulfate (0.044mg/kg/SC) and xylazine (1mg/Kg/IM). Anesthesia was induced with thiopental sodium (12.5mg/kg/IV) and the animals were intubated with a 7mm endotracheal tube. Anesthesia was maintained with halothane (0.5% to 1%), and the animals spontaneously breathed a mixture of air and oxygen (3L/min).

A median cervical incision was performed from the cricoid cartilage close to the upper chest, exposing the third up to the 13th tracheal rings. To develop malacia, we performed an extra-mucosal resection of 70% of the circumference of each tracheal ring (Figure 1), as described by Korpela et al. and Marquette et al.

Two experimental groups were created according to the number of resected rings: group A (n = 5), from the sixth to the eighth tracheal rings; group B (n = 4), from the fifth to the tenth tracheal rings. The muscle and skin planes were sutured with 3-0 polyglactin and 4 0 mononylon, respectively. Soon after, rigid bronchoscopy was performed for the first application of 23% NaOH.

In the postoperative period, the animals were daily observed by a veterinarian and by researchers, with special attention to the lack of appetite, vomiting, cough, stridor, tachypnea, and intercostal retractions. We also held weekly bronchoscopies. Analgesia was performed with tramadol (1mg/Kg/SC) postoperatively as required.

The technique of inflammatory stenosis with application of 23% NaOH was previously described and published by two of the authors of this study: through a rigid bronchoscope (Karl Storz Endoskope®, Germany) we applied a cotton swab soaked in 23% NaOH solution (pH = 14) on the lining of the trachea with previously surgically induced malacia. We avoided the contact of the NaOH solution to the posterior wall of the trachea. Was subsequently repeated to application of this solution on the region of the previous scar every three (group B) or seven days (group A). The applications were repeated until there were bronchoscopic finding of stenosis greater than 50% or signs of airway obstruction.

Bronchoscopies were performed with rigid tubes, with external diameters from 6.5mm to 8.5mm. The animals were inspected at three or seven days, depending on the group, in order to access the tracheal lumen diameter before instillation of NaOH solution.

The animals were kept under daily observation. Clinical signs of airway obstruction were monitored, as well as other medical or surgical complications. They were sacrificed with a lethal dose of sodium thiopental (100mg/kg, iv) if there was any significant sign of respiratory failure or ascertained the presence of stenosis above 50% of the tracheal lumen before 50 days of follow-up. After euthanasia, the cervical tracheae were collected and fixed in 10% formalin.

The tracheal specimens were transversely cut in two levels: level 1: in the middle portion of the zone with...
greater stenosis; and level 2: in a location 2cm away from the stenosis, corresponding to the normal diameter of the trachea. The specimens were embedded in paraffin, cut with a 5mm thickness and stained with hematoxylin and eosin. The epithelium, the submucosa and inflammatory reaction were analyzed under an optical microscope with a 10X and 40X objectives.

The latero-lateral and anteroposterior diameters of the tracheal lumen of each specimen were measured using the program Sigmascan Demo-Image Analyser (Sigma®, Chicago, USA). This software allowed us to calculate the area of the tracheal lumen by using the following equation: Area (mm$^2$) = \pi \times \frac{\text{sagittal diameter} \text{ (mm)}}{2} \times \frac{\text{coronal diameter} \text{ (mm)}}{2}.

The ratio between the area of greatest stenosis (level 1) and the area of normal trachea (level 2) expressed the degree of narrowing of the tracheal lumen. Was used the Mann-Whitney test to compare the two groups in the residual lumen at level 1 (area with greater stenosis). The difference was considered statistically significant if \( p < 0.05 \).

RESULTS

The tracheas of six animals were analyzed. Three animals were excluded from the study: one animal (#1) from group A died from diarrhea two days after the first application of NaOH. Two other dogs (# 8 and #9) in group B were precociously sacrificed 24h after the first application of NaOH due to respiratory failure. Their tracheae had swollen up, but no stenosis greater than 50% had developed. The main results are shown in table 1.

In two animals (#2 and #6), there was a small laceration of the tracheal mucosa during excision of the cartilage rings. These perforations were immediately sutured with 4-0 polyglactin. On the tenth postoperative day, dog #6 developed a limited cervical subcutaneous emphysema, without other consequences. This dog had pneumonia on the eighth postoperative day, which was treated with penicillin.

At the time of the first application of NaOH, we could identify the region of malacia more clearly in group B. In all bronchoscopic examinations following the application of NaOH there was some degree of deposition of fibrinoid material in the tracheal mucosa, as well as scarring and narrowing of the area of application of the caustic solution.

On macroscopic examination, we observed stenosis and fibrosis in all tracheae. In areas of stenosis, there was distortion and blurring of the trachea rings (Figure 2). The average longitudinal extent of the stenosis was 1cm and 2cm in groups A and B, respectively.

The residual lumen of the trachea at level 1 (light stenosis) ranged from 1% to 29% (Table 1). The residual lumen level 1 in group A (9.1%) was not statistically different \(( p = 0.355)\) from the one of group B (12%). The average lumen area of level 1 in group A \((17.92 \text{mm}^2)\) was not smaller, statistically \(( p = 0.355)\) than in group B \((29.8 \text{mm}^2)\).

All tracheae displayed wall thickening by fibrosis and inflammatory infiltrate. Fibrosis and necrosis were mainly observed in the submucosa and adventitia, occupying the region of the excised cartilage rings. Granulation tissue predominated in the mucosal surface, leading to partial obstruction of the lumen. The distribution of the sequential phases of the inflammatory process in the tracheal wall was the result of applying the solution to 23% NaOH at different times (Figure 3).

The inflammatory infiltrate, consisting mainly of polymorphonuclear cells, predominated in the mucosa and submucosa. In all specimens, we observed desquamation of respiratory epithelium. Squamous metaplasia was observed in dogs #2 and #6 (Figure 3).

DISCUSSION

Although tracheoplasty is the treatment of choice for most cases of inflammatory tracheal stenosis, in some situations, as in patients without clinical conditions, non-surgical procedures are required, such as the placement of silicone tracheal stents.

<table>
<thead>
<tr>
<th>Dog</th>
<th>Group</th>
<th>Instillations of NaOH</th>
<th>Lumen at level 2 (mm$^2$)</th>
<th>Lumen at level 1 (mm$^2$)</th>
<th>Residual lumen (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>A</td>
<td>3</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>2</td>
<td>A</td>
<td>3</td>
<td>193.7</td>
<td>56.9</td>
<td>29.3</td>
</tr>
<tr>
<td>3</td>
<td>A</td>
<td>3</td>
<td>196.8</td>
<td>9.4</td>
<td>4.7</td>
</tr>
<tr>
<td>4</td>
<td>A</td>
<td>3</td>
<td>164.8</td>
<td>1.5</td>
<td>1</td>
</tr>
<tr>
<td>5</td>
<td>A</td>
<td>3</td>
<td>296.7</td>
<td>3.9</td>
<td>1.3</td>
</tr>
<tr>
<td>6</td>
<td>B</td>
<td>4</td>
<td>277</td>
<td>43.9</td>
<td>15.8</td>
</tr>
<tr>
<td>7</td>
<td>B</td>
<td>3</td>
<td>193.4</td>
<td>15.7</td>
<td>8.1</td>
</tr>
<tr>
<td>8</td>
<td>B</td>
<td>1</td>
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<tr>
<td>9</td>
<td>B</td>
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</table>

Table 1 - Number of applications 23% NaOH solution and morphometric analysis.
To develop and improve airway orthoses, some experimental models of stenosis and malacia have been described. Once the silicone stents are more satisfactorily set in stenoses of more than 50% of the lumen and greater than 1 cm in extension, we seek to adapt and combine two models of stenosis: resection of cartilage rings in a row, originally described by Mair et al., and then modified by Korpela et al. and Marquette et al., and the application of caustic solution by bronchoscopy, published by our group.

We thus promote the inflammatory changes found in post-intubation stenosis through an easy to perform and inexpensive method, consisting of application of 23% NaOH in the tracheal mucosa at intervals of seven (Group A) or three days (group B) to accelerate the development of stenosis. Additionally, we adapted, in dogs, the technique of removal of the cartilaginous rings described above for pigs and rabbits, with the goal of producing a dynamic narrowing of the airways and a greater reduction of the lumen, since the standalone application NaOH yields only superficial injury, without reaching the perichondrium cartilage.

The results of our experiments demonstrate the efficacy of combining the two methods described above for the development of an inflammatory stenosis with a reduction of at least 70% of the diameter of the airway. Additionally, there was no statistical difference in relation to the intensity of stenosis produced when comparing the number of rings resected and the interval between applications of NaOH.

Although two surviving animals in group B developed satisfactory 2cm long stenosis, two others of the same group had to be prematurely sacrificed due to airway obstructive symptoms. On the other hand, our study also showed that the resection of only three cartilage rings and the application of NaOH in a span of three rings promotes an apparently less intense stenosis, but shorter (<2 cm), perhaps rendering it inappropriate for the evaluation of orthoses.

In the article by Korpela et al., five consecutive rings were resected and only one animal died, by airway obstruction, between 32 operated. Thus, it appears that the extent of experimental malacia was not the only factor that caused respiratory dysfunction of the two early euthanized animals in group B. However, an association of six rings malacia and the edema caused by the application of NaOH possibly led to severe obstruction of the airway seen in these dogs.

Perhaps the weekly interval of applications, or even a decrease in concentration of NaOH solution, could result in a slower installation stenosis in group B. In the study by Hanauer et al., 23% NaOH was also instilled once a week (up to three applications), in the larynx and produced a stenosis of at least 50%, with no deaths associated with airway obstruction.

We justify the use of fewer animals in the study due to the early development of severe inflammatory tracheal stenosis. Also, the death of two animals in group B led us to limit the number of animals.

Recession of three or six rings, associated with applications NaOH solution, causes a significant...
narrowing of the trachea. However, future work should show whether the weekly application of NaOH alone for a length of at least 2cm of the airway, or simple resection of six or more tracheal rings, will provide a superior experimental model, without causing severe acute airway obstruction, greatly reducing the loss of animals, meaning a more safe and effective model to test new tracheal stents.

**RESUMO**

**Objetivo:** Desenvolver, experimentalmente, malácia e estenose traqueal para testar novos modelos de órteses traqueais. **Métodos:** Ressecamos três anéis cartilaginosos da traqueia cervical de cães no grupo A (n=5) e seis anéis no grupo B (n=4) para produzir malácia. Logo após, uma mucosa da região com malácia recebeu aplicações de uma solução de hidróxido de sódio (NaOH) a 23%, e os animais eram acompanhados com exames broncoscópicos para observar o desenvolvimento de estreitamento da luz da via aérea. Quando a estenose era de mais de 50% da luz, ou havia sinais mínimos de insuficiência ventilatória, os animais eram sacrificados. O segmento de via aérea estreitada foi então coletado para análise histológica e era calculada a área de luz residual do segmento traqueal com estenose e malácia. **Resultados:** Na análise histológica, foi constatada fibrose na submucosa e adventícia, associada a granulomas na mucosa. A luz residual média dos segmentos com estenose foi de 9% e 12% nos grupos A e B, respectivamente, (p>0,05). **Conclusão:** A combinação da ressecção de anéis cartilaginosos e da aplicação de NaOH 23% na mucosa respiratória promoveu uma estenose traqueal intensa, porém esteve associada à perda de animais. Novos estudos são necessários para verificar se o emprego isolado de uma das técnicas seria mais seguro e eficaz para desenvolver estenose traqueal.


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