The Effect of Fibroblast Growth Factors in Grafted Fascia into the Vocal Fold of Rabbits

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Abstract

Introduction  The human larynx is a very important organ for communication. Many conditions lead to scarring of the vocal folds, decreasing voice quality.

Objective  We aimed to determine whether fibroblast growth factors (FGFs) may influence tissue integration of grafted fascia into the vocal folds of an animal model.

Methods  This is an experimental animal study with 12 adult rabbits that were submitted to a grafting fragment obtained from superficial cervical fascia into the vocal fold lamina propria, bilaterally. The right vocal fold was injected with FGFs. The animals were sacrificed after 1 month or 12 months, depending on the group they were assigned to, and a histological analysis of their vocal folds was performed. We analyzed the histological changes (such as the presence of fibrosis and neovascularization) induced by the acute or chronic inflammatory reactions.

Results  The FGFs induced acute inflammatory changes in all animals after 1 month of the initial experiment. The presence of FGFs triggered more fibrosis than the expected due to the surgical procedure itself when compared with the control side of all animals after 12 months of the initial experiment.

Conclusions  Fibroblast growth factors alone do not represent a good therapeutic option in phonosurgery, since we observed higher levels of fibrosis in the vocal fold lamina propria. Further studies combining more substances may be necessary to elucidate the best option to be used in this kind of surgery.

Keywords  ► vocal folds  
► rabbits  
► FGFs  
► fibroblasts  
► grafting  
► fascia

Introduction

The human larynx enables interpersonal interaction. No other organ of the human body presents similar structural arrangement like the vocal folds, especially the flexibility and vibratory features, which are related to sound production.¹

Several conditions promote structural changes of the vocal fold lamina propria, with the formation of scar tissue (increased deposition of type I collagen and fibronectin, decreased elastin, decorin, and hyaluronic acid), decreasing the vibration, leading to a poorer voice quality.²–⁴ Since the 1990s, many studies focused on reducing postoperative fibrosis and scarring, and to improving the flexibility of the vocal folds.⁵–¹¹ However, many of these techniques are not yet available for clinical usage.

Fibroblast growth factors (FGFs) are potent regulators of cell proliferation, differentiation and function and are critically important to normal development, tissue maintenance, wound repair and angiogenesis. Fibroblast growth factors are also linked with several pathological conditions.¹² Therefore, we aim to determine whether FGFs may influence tissue integration of grafted fascia into rabbits' vocal folds.
Method

Ethics
The Institutional Review Board approved this study (CEP 1033–1).

Specimens
Twenty New Zealand rabbits of both genres were used in this experiment, weighing between two and three kilograms. The number of male and female animals was the same. All the animals were kept with water and food “ad libitum,” from the moment of the procedure to the moment of their sacrifice.

The rabbits were divided into two groups. Group A, with animals maintained under observation for 12 months after the surgical procedure, and group B, which was observed for 1 month. The animals were randomly assigned to each group.

Animals with congenital malformations of the larynx and female rabbits that got pregnant during the experiment were excluded from this study.

Surgical Procedure
The animals underwent intramuscular anesthesia with ketamine (25 mg/kg/dose) and xylazine (3 mg/kg/dose) and kept with spontaneous breathing.

A 4-cm cervical longitudinal incision was performed at the level of the thyroid cartilage. The subcutaneous tissue was exposed by using scissors, and a fragment of superficial fascia of the neck, of ~ 10 x 10 mm, was excised. The fascia was extended on a metal surface and left to dehydrate spontaneously.

A 2-mm longitudinal incision was made along the vocal process of the vestibular face of the right and left vocal folds, close to the glottic aperture. A 45 degree-angled dissector was introduced into the incision to create a submucosal pouch, parallel to the medial face of the vocal fold, ~ 2 mm long anteroposteriorly, in both right and left vocal folds.

After dehydration, a fragment of 1 x 1 mm of the fascia was inserted into the submucosal pouch. The right vocal fold was injected with 5 µL of FGFs solution (concentration of 1 ng/µL, diluted in sodium chloride) immediately after the fascia was implanted. The left vocal fold received no injection of FGFs and remained as a control for histological changes due to grafting the fascia.

Histological Analysis
After the animals were sacrificed, their larynxes were obtained and maintained in a 10% formalin solution for 24 hours. The vocal folds were then resected with the thyroid cartilage, the segments were histologically processed (3 μm of thickness for each section) and embedded in paraffin.

Every section contained both vocal folds for comparison. The slides were stained with hematoxylin and eosin (H and E), picrosirius red, and Masson trichrome and then examined under light microscopy (in different magnifications) to assess the following parameters: a) number and type of inflammatory cells (neutrophils, lymphocytes, and macrophages); b) presence or absence of fibrosis; c) neovascularization; d) the density of collagen within the grafted area; and e) presence or absence of foreign body reaction.

The inflammatory reaction was semiquantitatively evaluated by comparing the small portion of the surrounding of the implanted fascia with the lamina propria beyond this area. It was then classified as:

0 – absence of mononuclear cells;
I – mild (1 to 10 mature mononuclear cells were observed);
II – moderate (11 to 20 mature mononuclear cells were observed); and
III – severe (more than 20 mature mononuclear cells).

Additionally, the inflammatory reaction was classified as acute when there was presence of polymorphonuclear cells, and as chronic when there was presence of a foreign body reaction with active macrophage predominance.

Fibrosis and the density of collagen within it were analyzed using a Sony DXC-101/Trinitron CCD Camera System (Sony Electronics Inc., San Diego, CA, USA), scanned through an OCULUS TCX System (Teledyne DALSA, Waterloo, ON, Canada), and analyzed with the image editing software Image Plus 4.5 for Windows (Astra Image, Kawanishi City, Japan). Fibrosis was also semiquantitatively assessed and graded as:

0 – absence of collagen fibers;
I – mild (few foci of collagen fibers);
II – moderate (continued collagen fibers around the grafted area); or
III – severe (bigger area of continued collagen fibers around the grafted area, consisting a fibrous capsule).

Neovascularization was defined by the presence of thin wall blood vessels within the grafted area.

Statistical Analysis
We calculated and compared the data obtained from the analysis of the inflammatory reaction, presence of neovascularization, and fibrosis using the Fisher exact test. Statistical analysis obtained from the density of collagen fibers was performed by non-parametric t-test, Wilcoxon test, and Kruskal-Wallis algorithm, comparing both the left and right sides, and in both groups (A and B). Findings were considered statistically significant when p values were < 0.05.

Results
Our final groups comprised of 6 rabbits in group A (12-months survival) and 6 rabbits in group B (1-month survival). All data obtained from the histological analysis are shown in Table 1.

Histological analysis of cellular predominance in both vocal folds (injected with FGFs or control) revealed predominant lymphohistiocytic infiltration with no differences between sides or groups. Additionally, the lymphohistiocytic pattern was statistically significant when compared with the other types of cells within the inflammation (p = 0.008).

There was a statistical difference between both groups when we compared the presence of inflammatory reaction and the side of the vocal fold. The intensity of inflammatory reaction in the right vocal fold (with FGFs injection) showed frequent association with a “moderate” or “severe” reaction in group B; however, in group A, it was related to an “absent” or “mild” reaction (p = 0.011) (Figs. 1 and 2). The intensity
of inflammatory reaction in the left vocal fold (considered as a control group) showed no differences between both groups ($p = 0.573$).

The intensity of fibrosis was statistically significant when we compared the right vocal fold in both groups ($p = 0.014$). It was frequently found in group A, and considered as “mild” in most of the samples. The left vocal fold in both groups showed no fibrosis between both groups (Figs. 3 and 4).

Neovascularization was only observed in the right vocal folds (injected with FGFs) of animals from group A. However, it was not statistically significant between the groups.

None of the vocal folds among both groups showed signs of chronic inflammatory reaction, presence of giant cells, or foreign body reaction.

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<th>Table 1 Histological findings in both groups</th>
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Abbreviations: H, histiocytic pattern; L, lymphocytic pattern; LH, lymphohistiocytic pattern; LVF, left vocal fold; RVF, right vocal fold.

Note: 0, absence; I, mild; II, moderate; III, severe.

**Discussion**

Many techniques aim to restore the functional mechanism and provide a better oscillation of the vocal fold’s mucosa, such as: vocal therapy, injections of collagen or hyaluronic acid, axial incisions of the mucosa, excision of the scarring tissue; additionally, another option consists in the insertion of foreign tissue, such as fat or muscular fascia between the epithelium and the vocal ligament.13–17 Fascia graft has been used for decades in otologic and plastic surgeries without complications and with reproducible results. Fascia presents a similar consistency to that of collagen and low rate of metabolic activity. Rihkanen18 proposed the use of minced fascia in the surgery of volumetric augmentation of vocal

![Fig. 1](image1.png)  
**Fig. 1** Right vocal fold of a rabbit from group A with grafted fascia within the lamina propria and injected fibroblasts growth factor. Mild inflammatory reaction (arrow) (hematoxylin and eosin x20).

![Fig. 2](image2.png)  
**Fig. 2** Right vocal fold of a rabbit from group B with grafted fascia within the lamina propria and injected fibroblasts growth factor. Moderate inflammatory reaction (arrow) (hematoxylin and eosin x20).
The presence of vocal fold scars due to injury or inflammation can alter the ultrastructure of the lamina propria, disrupt the normal mucosal wave, and may result in glottic incompetence and dysphonia, affecting the quality of life. Several substances have been used to prevent this problem and improve glottal closure, from synthetic to biological materials: Teflon, Gelfoam (Pfizer Inc., New York, NY, USA), intralesional steroids, bovine collagen, homologous collagen, muscular fascia, and fat. Our study is the first to evaluate the fascia fragment integration into the lamina propria with the addition of FGFs.

First, the presence of a lymphohistiocytic pattern was observed in both groups (after 1 and 12 months of procedure) of grafted vocal folds (left and right). It is also important to note that the absence of macrophages among all the samples may be related to an absent activating factor, triggering macrophage activity. Reijonen et al demonstrated that the presence of macrophage is related to foreign body chronic inflammatory reaction. However, Reijonen et al identified that no intense inflammatory reaction was fostered until up to 12 months after muscular fascia injection in paralyzed canine vocal folds. The isolated presence of the fascia did not trigger inflammatory reaction, as demonstrated by Duke et al, who evaluated vocal folds that have undergone crushed fascia lata injection.

Secondly, in vitro studies showed that the FGFs led to an increased production of type I collagen and hyaluronic acid. Yet, Scapini et al demonstrated that their therapeutic interventions in an animal model of rabbits statistically increased the density of collagen around the graft. However, Akdogan et al found significant difference in a group of vocal folds from rabbits treated with vitamin A compared with an untreated group (lesser deposition of collagen and fibroblasts in the first group); and Krishna et al, studying in vitro and in vivo vocal fold scar model of rabbits with hepatocyte growth factor exposure, found a decreased collagen secretion by fibroblasts. Our study demonstrated that the group of vocal folds injected with FGFs seemed to trigger an increased inflammatory reaction within 30 days after the procedure, compared with the group of vocal folds with no FGFs injection. Furthermore, the density of collagen observed within the vocal folds is bigger in rabbits from group A than in animals from group B, in both sides. Our observations are similar to others found in the literature, since healing and scarring processes take more than 1 month, and mature collagen was also present during those studies.

Interestingly, the grafted fascia injected with FGFs seemed to be associated with an increased rate of neovascularization, which may indicate a positive effect of grafted tissue integration on the receiver site. No preliminary work evaluated the neovascularization process in grafted vocal folds.
Our findings also suggest that FGFs may have triggered an acute inflammatory reaction. Because there was a statistical difference between both groups and both sides of the vocal folds, we may assume that the presence of grafted fascia was not the only factor to trigger the inflammatory reaction. Only one study had evaluated the presence of inflammatory reaction after grafting fascia into vocal folds,² with similar findings.

This is the first study to evaluate the cervical superficial fascia fragment integration into the vocal folds of rabbits with the addition of FGFs. All specimens in the longer observation group showed frequent signs of fibrosis that may have been triggered by FGFs. This may suggest that fascia grafting, used isolatedly, may become a good option for surgical procedures because it did not induce an acute or chronic inflammatory reaction or trigger fibrosis.

Conclusion

Fibroblast growth factors may have induced fibrosis and neovascularization when injected in the vocal fold of rabbits. Fibroblast growth factors alone do not represent a good therapeutic option in phonosurgery, since we observed higher levels of fibrosis in the vocal fold lamina propria. Laryngologists should be aware of this when performing phonosurgery.

References

14 Chan RW, Titze IR. Hyaluronic acid (with fibronectin) as a bioimplant for the vocal fold mucosa. Laryngoscope 1999;109(7 Pt 1):1142–1149