Structural changes in the extracellular matrix after cross-linking of nasal polyp tissue

Cambiamenti strutturali della matrice extracellulare dopo cross-linking del tessuto polipoide nasale

Nathalia Coronel1*, Rogério Pezato1*, Jônatas Bussador do Amaral1, Leandro Ticlia2, Thiago Freire Pinto Bezerra3, Ligia Ferreira Gomes4, Juliana Dreyfuss4, Wallace Chamon5,6, Luiz Carlos Gregório1

1 ENT Research Laboratory, Department of Otolaryngology-Head and Neck Surgery, Federal University of Sao Paulo, Sao Paulo, Brazil; 2 Laboratory of Micro rheology and Molecular Physiology, Institute of Physics, University of São Paulo-USP, São Paulo, SP, Brazil; 3 Department of Otolaryngology, Federal University of Pernambuco-UFPE, Recife, PE, Brazil; 4 Department of Biochemistry, Federal University of São Paulo-Unifesp, São Paulo, SP, Brazil; 5 Department of Ophthalmology and Visual Sciences, Federal University of São Paulo-Unifesp, São Paulo, SP, Brazil; 6 Department of Ophthalmology and Visual Sciences, College of Medicine, University of Illinois at Chicago, Chicago, IL, United States

* N. Coronel and R. Pezato contributed equally to this work.

SUMMARY

Chronic rhinosinusitis with nasal polyposis is a disease characterised by a mechanical dysfunction that facilitates anomalous growth of the nasal mucosa, due to its unique remodelling process. The development of procedures capable of interfering with this process is thus of the utmost importance. This study deals with the evaluation of the effects of cross-linking upon the nasal polyp tissue extracellular matrix. Six patients undergoing surgery for polypectomy were selected. The riboflavin/UVA collagen cross-linking technique was applied to the surface of epithelialised and de-epithelialised resected polyps of the intervention group. The control group polyps were not submitted to cross-linking procedures. Ultrathin polyp tissue sections (8 μm) were processed for immunofluorescence with mouse anti-type I collagen antibody and AlexaFluor 488 conjugated secondary antibody plus DAPI counterstaining, and analysed by confocal microscopy. The effect of riboflavin/UVA collagen cross-linking was visible on confocal fluorescence microscopy. Quantitative morphology was associated with fluorescence imaging analysis, and pixel density and brightness were evaluated. The surface of treated polyps exhibited a higher density of collagen fibres compared to control polyps, as could be observed both visually and through objective measurements of the fluorescent regions. The effect was enhanced on the surface of the de-epithelialised polyps. The higher density of collagen fibres exhibited by the de-epithelialised treated polyps demonstrates the feasibility of this technique in interfering with the remodelling process and the mechanical dysfunction found in chronic rhinosinusitis with nasal polyposis.

KEY WORDS: paranasal sinus disease, chronic disease, rhinosinusitis, chronic rhinosinusitis, sinusitis

RIASSUNTO

La rhinosinuite cronica con poliposi nasale è una malattia caratterizzata da una disfunzione meccanica che facilita la crescita anomala della mucosa nasale, a causa del suo peculiare processo di rimodellamento. Questo studio si propone di valutare gli effetti del cross-linking sulla matrice extracellulare del tessuto del polipo nasale. Sono stati selezionati sei pazienti sottoposti ad intervento chirurgico di polipectomia. È stata applicata la tecnica del cross linking del collagene con riboflavina / UVA alla superficie dei polipi resecati epithelializzati e disepitelializzati del gruppo di pazienti sottoposti ad intervento. I polipi del gruppo di controllo non sono stati sottoposti a procedure di cross linking. Le sezioni ultraottitzili (8 μm) di tessuto polipoide sono state analizzate mediante immunofluorescenza con anticorpi anti collagene di tipo I di topo e anticorpi secondari coniugati AlexaFluor 488 con controcolorazione DAPI e analizzate mediante microscopia confocale. L’effetto del cross-linking è risultato visibile al microscopio confocale a fluorescenza. La morfologia

Conflict of interest

The Authors declare no conflict of interest.
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Introduction

Chronic rhinosinusitis with nasal polyposis (CRSwNP) is characterised by chronic inflammation of the nasal mucosa with an altered remodelling process, which affects extracellular matrix (ECM) formation and facilitates mechanical dysfunction and oedema formation. The pathogenesis of CRSwNP is still not fully understood; however, its foundation is the aforementioned chronic inflammatory process, characterised by a major imbalance in immunomodulation which increases inflammatory cells (such as dendritic cells, eosinophils, and T helper cells) and decreases immunoregulatory cells (such as regulatory T cells), as well as an impaired immunomodulation of mesenchymal stem cells.

Studies on the mechanical forces involved in nasal polyp development have been conducted in an attempt to at least partly elucidate the pathophysiology of CRSwNP. Changes in the mechanical response of the nasal mucosa have been observed in patients with CRSwNP compared to the healthy mucosa of the middle meatus, with a loss of effectiveness in adequately increasing the interstitial hydrostatic pressure (IHP) in response to water inflow during the inflammatory process.

This discovery provides a new perspective for biomechanical management of CRSwNP beyond treatment of the inflammatory process (corticosteroids, non-steroidal anti-inflammatory drugs, leukotriene receptor antagonists, anti-immunoglobulin antibodies and biological agents that modulate the inflammatory environment). Within this context, the study of the mechanical dysfunction of CRSwNP has become increasingly relevant. It has previously been shown that ECM composition interferes with the biomechanical response of nasal mucosa. Nasal mucosa with fibrosis (synchiae) and healthy nasal middle meatus mucosa produce similar mechanical responses, suggesting that changes in ECM composition may contribute to prevent oedema formation and, potentially, have an impact on CRSwNP treatment.

Recent studies have confirmed that increased IHP contributes to a reduction of nasal polyps. In one investigation, the authors increased this pressure by imparting continuous positive airway pressure (CPAP) in nasal polyp tissue. Such indirectly increased interstitial hydrostatic pressure was associated with a decrease in polyp volume and milder nasal-obstruction symptoms. Considering the close interdependence between the IHP response and ECM composition, the remodelling process is an essential target in approaching the mechanical dysfunction of CRSwNP. It is reasonable to speculate that not only would changes in ECM composition be capable of altering the mechanical properties of the tissue, but also that structural changes in the existing ECM could affect tissue mechanical properties. In ophthalmology, the riboflavin/UVA cross-linking technique has been used to alter ECM conformation and thus prevent the progression of corneal bulging in keratoconus, a disease characterised by derangement of ECM deposition, which facilitates change in the shape of the cornea.

In the same way, the strengthening of the structures that compose the ECM in nasal polyp mucosa might impact the remodelling process and act on the biophysics mechanisms involved in the nasal polyp development. In this context, we evaluated the effect of this cross-linking technique upon polyp stromal collagen in patients with CRSwNP.

Materials and methods

This study was approved by the University of São Paulo Research Ethics Committee, protocol number 2905802, and conducted in accordance with the provisions of the Helsinki Declaration.

Histological sections of nasal polyp tissue from 6 patients with NP, diagnosed according to the European Position Paper on Rhinosinusitis and Nasal Polyps (EPOS) criteria, were evaluated. Patients between 18 and 60 years of age, non-steroidal anti-inflammatory tolerant, non-smokers were selected. The exclusion criteria were degenerative or autoimmune diseases, cystic fibrosis, pregnancy and topical or systemic corticosteroid therapy within the 4 weeks preceding surgery for removal of nasal polyps.

Application of the cross-linking technique

Immediately after resection, polyp fragments were prepared and divided into two groups: control and intervention. The control group received no intervention. In the intervention group, riboflavin/UVA cross-linking was performed on the epithelial surface of the polyps of patients 1, 2, 3, 4 and 5.
In patient 6, the polyp was cut in half and the treatment applied to the de-epithelialised cut surface (Fig. 1). In the intervention group, a photosensitising solution containing 0.1% riboflavin-5-phosphate and 20% dextran T-500 was applied to the surface of the polyp every 5 minutes for 30 minutes to ensure saturation of the polyp stroma with riboflavin. After 30 minutes, UVA radiation (365 nm) was applied 45 mm from the polyp surface, for 30 minutes, using a solid-state device (X-Link, Opto Eletrônica, São Carlos, Brazil) with a surface irradiance output of 3 mW/cm² (Fig. 1). The energy applied was 5.375 J/cm². Surface irradiation was ensured by continuous, microprocessor-based, automated monitoring by the X-Link device, which uses an internal power meter. After application of the cross-linking technique, polyps were frozen at -80°C. Ultrathin cryostat sections (8 μm, -21°C) of treated and untreated polyps were obtained and analysed. In the first four patients, sections were made parallel to the surface exposed to riboflavin/UVA. In patients 5 and 6, sections were obtained longitudinally (perpendicular to the exposed surface), so that all slides included the UVA-exposed area and the underlying stroma.

**Eosinophil count**

Part of the polyps was fixed using 10% acetaldehyde, maintained for 24 hours at room temperature, and then preserved in 70% alcohol for analysis. The previously fixed nasal tissue portions were embedded in paraffin and 4-μm-thick sections were obtained using a microtome. All sections were affixed onto Superfrost Plus glass slides (Menzel Glaser, Braunschweig, Germany) and dried at 60°C for a few hours. For deparaffinisation, slides were washed successively in xylene (3 times for 10 minutes), 100% ethanol (2 times for 5 minutes), 90% ethanol (2 times for 5 minutes) and 70% ethanol (2 times for 5 minutes). The nuclei were stained with alum haematoxylin (Lillie-Mayer’s solution) for 5 minutes and rinsed in running tap water. Differentiation was performed with 0.3% acid alcohol, sections were rinsed again in running tap water and, subsequently, in Scott’s tap water substitute (sodium hydrogen carbonate 10 g, magnesium sulphate 100 g, distilled water 5 L). After rinsing in tap water, sections were stained with eosin solution (1% eosin Y 400 mL, 1% aqueous phloxine 40 mL, 95% alcohol 3100 mL and glacial acetic acid 16 mL) for 2 minutes, dehydrated, and cleared. Histological examination was performed by a pathology expert through a Leica DM2000 binocular microscope at 400x magnification. The absolute number of eosinophils per high-power field (HPF) was counted in an average of 10 independent fields of view selected from the most inflamed area of tissue.

**Processing for immunofluorescence**

The slides were washed with phosphate-buffered saline (PBS), covered with 2.5% PBS and albumin solution, and stored in a moisture chamber under refrigeration. After 2 hours, they were incubated with primary antibody (anti-type I collagen, Calbiochem, Darmstadt, Germany) at 1:50 concentration in PBS containing 1% bovine serum albumin and 0.1% saponin. The next day, the slides were again rinsed with PBS and incubated with secondary antibody (AlexaFluor 488 conjugated anti-mouse IgG, Molecular Probes, Carlsbad, CA) at 1:300 concentration in PBS containing 1% bovine serum albumin and 0.1% saponin. At 1:2500 DAPI (4,6-diamidino-2-phenylindole dihydrochloride, Molecular Probes) in a PBS solution containing 0.1% saponin, for 30 minutes, to ensure nuclear staining. Finally, slides were washed again with PBS and mounted with Fluoromount-G liquid medium (Electron Microscopy Sciences, Hatfield, PA).

![Figure 1. Polyp and area of riboflavin/UVA application. (A) In patients 1, 2, 3, 4, and 5: on the epithelialised surface of the polyp. (B) In patient 6: polyp cut in half and treatment applied on the de-epithelialised cut surface.](image-url)
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All binary sequences representing the same texture were equivalent because they represent the same texture pattern.

Objective assessment of images
The images obtained by confocal microscopy were also examined for two objective parameters: tissue texture, through pixel encoding by the local binary pattern (LBP) method and luminous intensity distribution.

1. Analysis of tissue texture by pixel encoding

   LBP is a method used in the field of computer vision for texture analysis of monochrome digital images. In the context of a digital image, the texture of each pixel is defined by the local variations in intensity of the nearest neighbouring pixels. When applied to an image, LBP compares the intensities of a central pixel with those of its 8 neighboring pixels, assigning a value of 1 when the intensity of a neighboring pixel is greater than or equal to the intensity of the central pixel and 0 otherwise. This generates a numeric representation that consists of a sequence of 8 binary digits (0 or 1), which represents the characteristics of the central pixel. This sequence of 8 binary digits is then expressed as its decimal equivalent. The 8-digit binary sequence can generate up to 256 different local patterns, some of which can be considered equivalent because they represent the same texture pattern.

   All binary sequences representing the same texture were grouped into the same set, known as the LBP code set. From this perspective, up to 10 sets of LBP codes could be created, which were denoted by numbers 0 to 9. Each of these codes was associated with a specific image feature. For example, codes 2 and 6 were associated with half-edges, while codes 3 and 5 were associated with dark and light edges, respectively. Half-edges, corresponding to discontinuous edges which do not exceed 2 pixels in length, represent collagen fibres that were photographed only partly or present any loss of continuity. Edges, in our context, represent the contour of whole collagen fibres.

   We applied the LBP method only to the regions of interest of the confocal photomicrographs. Data were obtained in the area juxtaposed to the surface to which the cross-linking agents were applied, at a depth of 105 μm (Figs. 2A-C), and in the control polyps, in the corresponding area also 105 μm in depth (Figs. 2D-F). Finally, a histogram of the decimal values attributed to each pixel group was generated, representing the texture pattern associated to the digital images of clinical specimens.

2. Analysis of luminous intensity distribution

   Analysis of luminous intensity distribution specifically evaluates pixels associated with greater brightness. In our photomicrographs, brighter areas (i.e., those with greater luminosity) represent collagen fibres.

   Brightness values in a monochrome image range from 0 to 255, where 0 represents the darkest hue and 255, the lightest. Bright pixels were defined as those with a hue exceeding a given threshold T, where 0 ≤ T ≤ 255. One threshold value was attributed to the whole set of digital images under study. Subsequently, it was calculated which percentage of the whole image pixels belong to the bright part of the image, revealing the extension of the image area occupied by collagen fibres. This strategy allowed collagen fibre areas of control and intervention images to be compared, as the control and intervention images were acquired under the same conditions.

Results

Eosinophils per High-Power Field (HPF)
All 6 polyps analysed presented more than 5 eosinophils per HPF, characterising the eosinophilic endotype of CRSwNP.

Tissue immunofluorescence microscopy

Microscopically, a 105 μm-thick fluorescent line was visible on the treated surface of epithelialised (Figs. 3A-B) and de-epithelialised (Figs. 3C-D) polyps, stratified into a superficial zone with more organised collagen fibres and a deeper zone with less organised collagen fibres. The differences between the control and intervention groups, concerning the organisation of collagen fibres were even more evident in the de-epithelialised polyps.

Analysis of tissue texture by pixel encoding

Among the controls of the epithelialised polyp group, the most frequent LBP profile observed was code 4 (Fig. 4A), corresponding to the edges of collagen fibres. Edge detection occurred due to the contrast of the bright collagen fi-
bres with dark areas in which fibres were absent. The presence of edges implied a reference level for the control fibre densities in the epithelialised polyp tissues. The exposure of the epithelialised polyp tissues to riboflavin/UV A induced a distinct LBP profile. The lowering of code 4 relative frequencies and the higher frequencies for code 8 corresponded to the bright hue of the labeled areas, denoting a greater concentration of collagen fibres in which individual edges were less observable.

Among the de-epithelialised polyps (Fig. 4B), code 4 was also the most frequent LBP profile among the control polyps, while the UVA-exposed de-epithelialised polyp had its highest peak on code 9.

The tissue texture analysis demonstrated that both the epithelialised and the de-epithelialised polyps increased collagen fibre concentration.

In both situations, the pattern of distribution of LBP code densities after UVA exposure has changed; density of code 4 decreased in the treated polyps, which corresponds to an increase in collagen fibre concentration.

**Luminous intensity distribution**

Figure 5 shows a plot of tissue luminous intensity distribution for individual patients. It was demonstrated a higher luminous intensity after UVA applications, which corresponds to greater collagen fibre densities in the analysed areas.

**Discussion**

Studies of tissue biomechanics in NP have led to a progressive paradigm shift by revealing that the development of
Figure 3. Photomicrograph of epithelialised nasal polyp section stained with anti-type I collagen antibody. (A) Control polyp, with rather disorganised collagen fibres (green) on the cut surface. (B) Intervention polyp exposed to riboflavin/UVA cross-linking of the epithelialised surface, with greater organisation of collagen fibres in the superficial stroma (above and to the right of the white line). Photomicrograph of de-epithelialised nasal polyp section stained with anti-type I collagen antibody. (C) Control polyp, with rather disorganised collagen fibres (green) on the cut surface. (D) Intervention polyp exposed to riboflavin/UVA cross-linking of the de-epithelialised surface, with highly organised collagen fibres in the superficial stroma (arrow). Sup. Str.: superficial stroma; Deep Str.: deep stroma.
this condition is based on two processes: not only chronic inflammation, but also mechanical dysfunction. Within this context, the present study aimed to induce changes in the structure of the polyp ECM with a view to potentially modify the mechanical dysfunction found in polyp tissue. This was the first investigation to evaluate a method capable of altering the shape and density of collagen in the ECM of nasal polyps. For this purpose, we used the riboflavin/UV A collagen cross-linking process. This method was chosen because it is already in clinical use in ophthalmology and has proven efficient in modifying the natural history of keratoconus, a disease characterised by derangement of ECM deposition, which facilitates change in the shape of the cornea  

At the cellular level, ontological genetic analysis has shown that, in mesenchymal stem cells derived from nasal polyps, genes related to the ECM remodelling process are differentially modulated when compared to those of bone marrow-derived mesenchymal stem cells. It has also been demonstrated that fibrotic tissue found in the nasal mucosa (synechiae) and the mucosa of the middle nasal meatus have similar mechanical properties. Moreover, it is suggested that the fibrotic modification of the biomechanical properties of the nasal mucosa of patients with CRSwNP, a tissue repair phenomenon in which increased collagen concentrations play a predominant role, could interfere in the altered remodelling process found in polypoid tissue.

Collagen fibres provide tensile strength to tissues. In living tissue, these fibres are stabilised through the formation of intermolecular covalent bonds, mediated by lysyl oxidase enzymes. The intensity of this stabilisation can be increased artificially by application of UVA radiation directly to tissue, without altering the composition of the underlying ECM, through the cross-linking technique.

Previous studies performed on the porcine cornea have shown that riboflavin acts as a photosensitising agent and, when irradiated by UVA, leads to the formation of free radicals that create new chemical bonds within the corneal...
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This procedure increases the diameter of the collagen fibres and establishes stronger interfibrillar attachment, increasing the biomechanical stability of the exposed tissue. Based on this method, we employed the same protocol for riboflavin/UVA exposure used in ophthalmology for keratoconus treatment in order to evaluate whether the same changes that occur in the corneal ECM can be induced in nasal polyp tissue.

In the images obtained from the epithelialised polyp, there was a slight increase in fibre density up to 105 μm below the epithelium, i.e., the depth reached by UVA radiation. Increased fibre density data were corroborated by objective analysis of the pixel patterns and brightness of images. The increase in collagen fibre density was greater in the de-epithelialised polyp, with a higher fibre density at the surface where the riboflavin and UVA were applied. This revealed that the presence of the epithelium interfered negatively with riboflavin penetration into the stroma. This phenomenon resembles that found in the cornea, where removal of the epithelium intensifies the effect of cross-linking.

Objective analysis of pixel patterns and luminosity in the irradiated region reinforces the results of visual examination and provides reliable data for comparison, showing a greater impact on fibre density in the treated polypoid tissue than in untreated tissue. These results also provide an opportunity for further investigations to replicate this pilot experiment, prompting further research into the ECM of the nasal polyp stroma and on the potential impact of structural changes to the conformation of the constituent proteins of the ECM on the mechanical properties of nasal polypoid tissue.

If the tissue changes observed after exposure to the cross-linking technique are shown to have a direct effect on the biomechanics of nasal polypoid tissue, this would open new avenues for the treatment of nasal polyposis, no longer based on anti-inflammatory agents or surgery alone, but rather by directly altering the mechanical properties of the affected tissue.

This technique has the potential for immediate application after polypectomy in patients with a history of recurrent CRSwNP, when this mucosa is already partially de-epithelialised. Once polyps have been removed and mucosal re-epithelialisation has not yet begun, one would expect for cross-linking to have the same effect observed in the de-epithelialised polyp in this study. We could also take advantage of the amplification of access to sinus cavities after sinus surgery and promote cross-linking on sinus walls.

Within this context, riboflavin/UVA cross-linking might also have utility as an outpatient office procedure to slow the growth of polypoid tissue, as our study showed that, although the method was less effective in epithelialised tissue, it still had an effect on the extracellular matrix structure of the nasal polyp. Prospective clinical studies are needed to evaluate the applicability of this procedure in the treatment of CRSwNP.

Conclusions

The results of this study show that riboflavin/UVA cross-linking promotes changes in the structural conformation of collagen fibres and increases their density on the exposed surface of nasal polyp tissue.

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