Different types of tissue composition in inflammatory or reparative upper airway disorders*

Laurence De Coster¹, Philippe Eloy², Liesbeth Ferdinande³, Jasmien Taildeeman³, Claude A. Cuvelier¹, Jean-Baptiste Watelet¹

¹ Department of Otorhinolaryngology, Ghent University, Ghent, Belgium
² Department of Otorhinolaryngology, CHU Mont-Godinne UCL, Belgium
³ Department of Pathology, Ghent University, Ghent, Belgium

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SUMMARY

Background: Composition changes of extracellular matrix (ECM) can lead to functional disorders of the upper airways (UA). The aim of this study was to systematically measure both the association patterns and the correlation degree between tissue composition parameters in UA inflammatory diseases.

Methodology: Nasal samples were obtained from patients with chronic rhinosinusitis with (CRS+NP), without nasal polyps (CRS), with post-operative adhesions (S) and normal nasal mucosa (NM). A reproducible semi-quantitative method, which takes epithelial and lamina propria damages into account was applied for haematoxylin and eosin, alpha-smooth muscle actin, reticulin, elastin, laminin and collagen type IV stainings.

Results: The most severe cases of epithelial shedding have been found in a significant higher amount in CRS+NP when compared with NM. The most severe cases of inflammatory reaction were mainly found in CRS+NP. CRS+NP had significantly more severe cases of oedema than NM. Excluding elastin, networks in other ECM proteins were found modified in fibrotic fields but to a lesser extend in oedematous regions in all conditions.

Conclusion: Although non specific, oedema in the lamina propria is a key-feature of CRS+NP, while fibrosis, massively present in CRS and S, affects profoundly the distribution of ECM proteins in these areas.

Key words: upper airways, chronic rhinosinusitis, nasal polyps, epithelium, extracellular matrix

Introduction

An adequate interaction between all anatomical and histological components of the nasal and paranasal cavities is necessary for ensuring the protective properties of the nose (1,2). The nasal pseudostratified columnar epithelium (respiratory epithelium) is separated from the lamina propria by a continuous basement membrane and a contingent of resident, inflammatory cells. The subepithelial region contains seromucous glands, blood vessels and a supporting connective tissue of loose type, which plays an essential role in inflammatory reactions but can also be the site of structural changes during chronic inflammations.

Remodelling is defined as a process leading to transient or permanent changes in tissue architecture, which involves breakdown of tissue structures as well as repair (3,4). During normal wound healing of the upper airways, it has been established that extensive oedema and fibrosis in nasal lamina propria appear and diminish in a well-coordinated sequential manner (5). However, changes in the composition of extracellular matrix (ECM) can lead to functional disorders and are suspected to
more, tissue changes interfere significantly with the responsiv-
ness to medical therapies, making the protective or deleterious
role of changes in airway tissue structure largely debated

In human and animal models, mucosal damages, such as epithe-
lial shedding, thickening of basement membrane, fibrosis, have
been regarded as important histomorphological features of
chronic rhinosinusitis  

Several ECM proteins have been the scope of recent publica-
tions on tissue composition in airway chronic inflammation.
Elastic fibres are ensuring tissue elasticity. Type-IV collagen is
primarily found in the basal lamina and could function as filtra-
tion system. Reticulin or reticular fibres is a type of ECM fibre
closely related to type-III collagen in composition. The laminins
are an important and biologically active part of the basal lamina,
influencing cell differentiation, migration, adhesion as well as
phenotype and survival  

Finally, besides the ECM proteins, the expression of alpha-Smooth Muscle Actin (α-SMA) correlates
with the activation of myofibroblasts, which are known to play a
key role in the development of fibrotic responses

The respective proportion of the specific epithelial and ECM
composition features remains largely unknown in chronic upper
airway diseases. To have an indirect view on the dynamic and
long-during remodelling process, the aim of this study was to
systematically describe its end results through the analysis of
major histomorphological damages, fibroplasia and neoan-
giogenesis in tissue samples from chronic rhinosinusitis with
(CRS+NP) and without nasal polyps (CRS), in post-operative
adhesions (or synechia) (S) in comparison with normal mucosa.
Finally, the expression and distribution of laminin, reticulin,
estatin, alpha-smooth muscle actin and type IV-collagen was
analysed on all tissue samples.

Materials and methods

Materials

Nasal biopsies were obtained in patients operated by FESS at
the department of Otorhinolaryngology, University Hospital
Mont-Godinne UCL, Belgium. Nasal samples were obtained
from patients with CRS+NP (n = 10) and CRS (n = 10) and from
patients with post-operative S (n = 10) during routine nasal and
sinus surgery. Samples from CRS and CRS+NP were obtained at
the level of ethmoid sinus cavities, while samples from synechia
were obtained from the resection piece. These patients were
operated after failure of appropriate medical treatment. Medical
systemic and local treatment was stopped two weeks before
surgery. Ten subjects with normal nasal mucosa were consid-
ered as controls. The normal mucosa obtained in controls, was
biopied at the level of the inferior turbinate. At the moment of
surgery, they were free of any acute or chronic nasal or sinus dis-
eases and were operated for snoring or lachrymal duct patholo-
gies. All patients and controls were non-smokers.
Allergy was explored in all cases by use of Skin Prick Test. If these
tests were negative and if the history was suggestive for this
condition, complementary RAST were performed. However, the
patients presenting positive allergy tests were not considered as
normal controls.
With regard to NSAID intolerance, the identification was done
on history basis. No systematic provocation test was performed.

Methods

Preparation of sections. Fresh surgical resection specimens of
human nasal tissues were collected and immediately fixed in 4%
buffered formalin (Labonord, Templemars, France). After embed-
ding in paraffin, five-micron sections were cut and mounted on

Histochemistry. Next sections were dewaxed and rehydrated
in graded alcohols. For the haematoxilin-eosin staining, slides
were immersed in Gills haematoxylin solution (Merck, Darm-
stadt, Germany) for 2 x 4 minutes. After washing in running tap-
water, sections were dipped in a HCl solution and submerged
for 3 minutes in a 1% eosin solution (Merck). Sections were then
dehydrated and mounted (Sakura, Zoeterwoude, The Nether-
lands).
The stainings for elastin and reticulin were performed manually
according to Van Gieson and Gomori, respectively. For elastin
staining, sections were stained using the Elastica van Gieson
staining kit (Merck). Briefly, sections were immersed in an elastin
solution for 10 minutes. After washing in running tapwater,
sections were submerged for 5 minutes in a haematoxilin –
iron nitrate solution followed by 2 minutes in a picrofuchsin
solution after washing. Sections were then dehydrated and
mounted with Tissue-Tek (Sakura). For reticulin staining, slides
were dipped in a kalium permanganate solution for 12 minutes
followed by washing in running tap water for 3 minutes. Then,
slices were submerged for 4 minutes in a kalium metabisulfit
solution and washed again after which sections were immersed
for 6 minutes in an iron aluin solution. Next, a silver solution was
applied (1 min) followed by formaldehyde (5 min), gold chloride
(10 min), kalium metabisulfit (4 min), natrium thiosulphate (1
min) and kernechtred for 5 minutes. Afterwards, sections were
dehydrated and mounted.

Immunohistochemistry. The immunohistochemical stainings
were performed on the Ventana automated immunostainer with
the Ventana iView DAB detection kit (Ventana Medical Systems,
Strasbourg, France) using the peroxidase anti-peroxidase techni-
que. Briefly, tissue sections were deparaffinized, rehydrated, and
antigen retrieval was performed for laminin (protease pretreat-
ment) and collagen type IV (0,01 M citrate (pH 6) pre-treatment)
staining. After blocking of aspecific peroxidase staining, primary antibodies were applied for 30 minutes (α SMA: 1/40, Biogenex MU128-UC, San Raman, California, USA; Laminin: 1/15, Biogenex PU078-UP; Collagen type IV: 1/20, DAKO M0785, Glostrup, Denmark). Next, biotinylated secondary antibodies were applied, followed by HRP-conjugated streptavidin. Immunostaining was finally visualized using 3,3'- diaminobenzidine as a chromogen (DAKO), and haematoxylin as a counter stain.

Quantification. A reproducible semi-quantitative method, which takes both staining percentage and intensity into account, was applied in haematoxylin and eosin, alpha-smooth muscle actin, reticulin, elastin, laminin and collagen type IV stainings. All parameters were scored semi-quantitatively on a four-point scale by a senior pathologist (CC), who was blinded for diagnosis and clinical data. Zero represented the lowest (absent) and three the highest score (severe). As some histological markers were more abundant than others in normal nasal mucosa, the scoring system was calibrated for each marker independently by examining a representative number of samples. Epithelial shedding was characterized in function of the length of basal cells or basement membrane uncovered by epithelial cells. The thickness of the basement membrane was determined as average of measurements in sections where the epithelium was properly oriented to avoid artifacts in measurement. Cells and ECM deposition were analyzed separately in different compartments, in epithelium, around glands, blood vessels and in lamina propria. The connective tissue, when possible, oedematous and fibrotic regions, fibroplastic and endothelial proliferation were also considered independently. The analysis included all areas of the biopsies and both a specific and global score were given for each parameter. Both the distribution and degree of severity of the histomorphological parameters were studied.

Statistical analysis
Statistical analysis was performed with SPSS 11.0 (SPSS Inc., Chicago, IL, USA). In function of the parameters, the non-parametric Mann Whitney U test or Chi-square were used to analyse the difference between groups. Correlations were described with their Spearman correlation coefficient r and their p value. A post-hoc adjusted statistical difference of p < 0.05 was regarded as significant.

This study was approved by the ethics committee of the faculty of Medicine and Health Sciences of University Hospital Mont-Godinne UCL (nr S8-2004).

Results
The mean age was 51.2 years with range from 18 to 79 years. The gender proportion was 15 males and 25 females. In the disease groups, 20% of patients were found sensitized for the most frequent aeroallergens (house dust mites, tree and grass pollens, animal danders) and 20% suffered from asthma. Two patients suffered from COPD and one had aspirin intolerance. Sensitization to aeroallergens and lower airway diseases were not found in the control group.

1. Histomorphological findings by Haematoxylin Eosin
1.a. Structural tissue. Epithelial shedding was found in all pathologic conditions and, in normal mucosa, only mild cases were observed. Moderate and severe cases were mainly found in CRS+NP and other pathologic conditions when compared to normal mucosa (Chi-square = 14.400, p < 0.0001) (Figure 1a). The most severe cases of epithelial shedding have been found in a significant higher amount in CRS+NP when compared with normal mucosa (p = 0.029). However, the thickness of the basement membrane was not found significantly different between pathologic conditions.

Gland atrophy was found to vary significantly between the disease conditions (Chi-square = 6.400, p = 0.011) but was not found in normal mucosa. When compared with normal mucosa, gland atrophy was found mainly increased in samples from CRS+NP (p = 0.029) and from S (p = 0.012) (Figure 1b). Signs of neoangiogenesis were only found in abnormal amount in the different disease conditions but not in normal mucosa. Even present in variable amounts between samples (Chi-square = 22.500, p < 0.0001), it was mainly found in S (Figure 1c). On the other hand, samples with extensive network of capillaries in the lamina propria were mainly observed in CRS+NP and CRS (Chi-square = 6.400, p = 0.011)

1.b. Inflammatory reaction. The severity of inflammatory reaction was found significantly different between the different conditions (Chi-square = 6.350, p = 0.042) and the most severe cases were mainly found in CRS+NP. On HE staining, eosinophils, neutrophils, lymphocytes, and mast cells were not found to be significantly different between groups but plasma cells were found in significantly higher number in CRS+NP when compared to normal mucosa (p = 0.029). The presence of lymphoid follicles in the lamina propria was found different between pathological samples (Chi-square = 16.900, p < 0.0001) and was mainly detected in CRS (Figure 1d).

1.c. Connective tissue. Chronic rhinosinusitis with nasal polyps had significantly more severe cases of oedema than NM (p = 0.009) (Figure 2a). CRS+NP were found to have less cases presenting fibrosis in lamina propria (Chi-square = 14.400, p < 0.0001) than other inflammatory conditions. Most severe cases of fibrosis were mainly found in S but this increase did not reach statistical significance when compared with the other conditions (Figure 2b). Fibroplasia was found significantly different between the disease groups, with more severe cases in CRS+NP and S (Chi-square = 20.600, p < 0.0001).
Table 1. Major histomorphological findings.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Nature</th>
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<td>- Gland atrophy</td>
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<td>Overall inflammation</td>
<td>Accumulation of plasma cells</td>
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<td>Fibroplasia</td>
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<td>- Fibroplasia</td>
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<td>Less cases of:</td>
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<td>- Decreased density of</td>
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Legend: CRS-NP: chronic rhinosinusitis without nasal polyps; CRS+NP: chronic rhinosinusitis with nasal polyps; S: synechiae (adhesions); ECM: extracellular matrix; aSMA: alpha-smooth muscle actin

Figure 1. Structural changes of nasal mucosa in normal and abnormal conditions.
NM: normal mucosa; CRS: chronic rhinosinusitis without nasal polyps; CRS+NP: chronic rhinosinusitis with nasal polyps; S: synechiae. Severity of a) Epithelial shedding, b) Gland atrophy, c) Neoangiogenesis, d) Presence of lymphoid follicles.
Table 1 shows a summary of the histomorphological findings in the different groups found in this study.

2. Histochemistry
2.a. Reticulin. Changes in distribution or intensity of reticulin network were not correlated with the type of upper airway disease. However, there was a non-significant trend in CRS+NP and S to present reduced or absent reticulin network in the subepithelial region (Chi-square = 5.150, p = 0.071).

2.b. Elastin. Elastin was found in all the walls of blood vessels. Elastin deposition was not found changed in fibrotic or oedematous regions. Elastin network was found respected in all pathologic conditions, without any significant differences between them.

3. Immuno-histochemistry
3.a. Alpha smooth muscle actin. No difference between diseases was observed in expression of aSMA in vessels. In normal mucosa, the majority of samples had a non-significant tendency to present moderate or high amounts of aSMA-positive cells while, in CRS+NP or CRS, none or few aSMA-positive cells were found (Chi-square = 3.600, p = 0.058).

3.b. Laminin. Laminin was present in blood vessels of all samples. In regions presenting fibrosis, staining of laminin was observable but not in oedematous field. The different diseases conditions did not show significant differences.

3.c. Type IV collagen. Staining for type IV collagen was mainly found in the basement membrane, blood vessels, and fibrotic fields. No significant difference was found between the groups. In oedematous regions, type IV collagen was poorly distributed.

Figures 3 illustrates tissue composition characteristics of CRS and CRS+NP, while Figure 4 reports ECM protein expression in fibrotic and oedematous fields.

Discussion
Even if CRS+NP and S present predominant oedematous and fibrotic patterns, respectively, this study demonstrated that many aspects of changes in tissue composition are not specific of these upper airway diseases and can present a wide range of severity.

In this cross-sectional analysis, we explored only the end results of the tissue remodelling process, which, for being completely addressed, should benefit of a longitudinal study design. The first innovative observation is that tissue changes can affect nearly all types of chronic inflammatory diseases in upper airways, although with different degree of severity and many similarities with ECM changes in asthma have been observed (13,14). In vasomotoric and allergic rhinitis, changes in tissue composition appears to be more limited (15-17). Epithelial shedding can accompany clinical response to cold, dry air or to urban pollution (18).

A second important finding is the confirmation of the close link between inflammatory reaction and extensive variations in ECM composition, which has been confirmed in this study. Lymphoid follicles are regularly found in chronic inflammatory processes in upper airways (19). Oedema is regularly found associated with a severe local inflammatory reaction, probably through a local activation of fibroblast matrix metalloproteinases by inflammatory cells (20). Finally, as demonstrated for the first time in this study, lymphoid follicles were mainly found in lamina propria of CRS samples and absent in normal tissue samples.

The third essential observation of this study is the confirmation that all histomorphological changes are not specific of a particu-
lar chronic upper airway disease. With regard to the epithelial changes, Ponikau et al., have reported that epithelial damage was observed in tissue from refractory CRS (21). The basement membrane thickening and the mean grade of sub-epithelial collagen deposition are significantly higher in CRS as compared to controls (22). It has been recently suggested that the increased basement membrane was correlated with prolonged duration of sinus symptoms and the coincidence of asthma (23, 24). Regarding the respiratory epithelium, nasal polyps show a diversity of histological changes including hyperplasia, atrophy or squamous metaplasia, and goblet cell hypertrophy (25,26). On the other hand, basement membrane thickening and epithelial damage, has been shown correlated to the local infiltration of eosinophils (27) and IL-17A-positive cells (28). In our study, the thickening of the basement membrane was not found specifically linked to a sub-group of chronic inflammatory diseases of the upper airways, independently of their allergic or asthmatic background. Another important observation is that, in contrast with the epithelial changes, the major damages observed at the subepithelial level seem to be more closely linked to specific inflammatory nasal or paranasal pathologies (29) extensive oedema in CRS+NP, extensive fibrosis in S and a combination of oedema and fibrosis in CRS. As already demonstrated in earlier histomorphological studies, in CRS+NP with or without cystic fibrosis, the lamina propria presents major oedema (30), which is considered as discriminative of CRS+NP against other types of CRS (31) and which is probably mediated by a large amount of cytokines and enzymes (32-34). Interestingly, the absence of neoangiogenesis in CRS+NP described in our study supports recent data (35). As demonstrated here, all components of the ECM, but elastin are

Figure 3. Comparison between chronic rhinosinusitis with or without nasal polyps.

a) CRS: lymphoid follicles (HES X 100); b) CRS+NP: severe subepithelial inflammatory reaction and oedematous lamina propria (HES X 100); c) CRS+NP: intact elastin network in oedematous fields (Elastin X 100); d) CRS+NP: reduced subepithelial network of reticulin (Reticulin X 200).
affected by the plasma exudation and pseudocyst formation. The distribution of elastin seems to be less affected by oedema probably because of its physico-chemical properties. The oedematous regions in both CRS and CRS+NP were found poor in αSMA-positive cells. This finding confirms the observation of Nishijima et al. showing that the distribution of αSMA-positive cells tended to be more remarkable in pedicle areas than in central and more oedematous areas.

For the first time, we demonstrated that, in post-operative adhesions, reticulin was also found affected in extremely fibrotic areas. Regarding the ECM composition, laminin and type IV collagen were found in fibrotic regions of S and CRS but no significant difference with other diseases was found. Interestingly, in contrast with observations in asthma, elastin seems also to resist better to fibrosis in upper airways and no major changes in the elastin network has been observed in S. This suggests probable differences in protective actions supported by elastin in upper or lower airway diseases.

Finally, although upper airways can also be subject to major ECM reorganization, this study has demonstrated that, in human, the gland integrity, absence of lymphoid follicles and limited neoangiogenesis should be considered as histomorphological markers of a healthy upper airway mucosa.

The study was limited to the pathological analysis and did not consider symptom severity, because all patients, after medical failure, were still presenting symptoms justifying surgery. For ethical reasons, we were not able to sample ethmoid tissue in controls: we considered nasal mucosa from the inferior turbinate

Figure 4. Fibrotic and oedematous fields.

a) Adhesion: fibrosis, fibroplasia and neoangiogenesis (HES X 200); b) Adhesion: type IV collagen deposition (Type collagen staining X 200); c) CRS: laminin in fibrotic regions (Laminin X 200); d) CRS+NP: absence of laminin in oedematous regions (Laminin X 200).
as representative of the paranasal mucosa. The processes leading to these changes and the time sequence of development of these different remodelling aspects were not considered in this study. For example, oedema can rapidly be induced; the tissue development of fibrosis takes more time. This complex relationship was not specifically explored in this study. As for all mechanistic analysis, these in vivo results must be compared with caution to in vitro findings.

Another limit of this study is the poor specificity of HE staining for describing sub-populations of inflammatory cells but this analysis was not considered as a primary objective of the study.

Conclusion

Changes in ECM structures are common findings in chronic inflammatory diseases of upper airways, but rarely specific. Chronic rhinosinusitis with nasal polyps and post-operative adhesions present more oedematous and a trend to more fibrotic matrices, respectively. Epithelial shedding and thickening of basement membrane were not found different between the sub-groups of diseases.

Authorship contribution

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Conflict of interest

None

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ERS- Clinical-Research Grant 2013/2014

In co-operation with the Foundation of Rhinology and Facial Plastic Surgery (RhiPla-Stifung Ulm, Germany) the European Rhinologic Society offers financial support for clinical research in the field of rhinology. There is an increasing demand for evidence-based results of diagnostic and therapeutic procedures, in particular surgical interventions. This is why this specific grant aims to support controlled clinical studies in all fields of rhinology including surgery of the nose, sinuses and anterior skull base.

The support amounts to € 10,000,- in a 2-years-period.

It can be used for equipment, material or personnel. A preliminary report is asked after 1 year, a final report after 2 years. An oral report should be presented at the biannual ERS-Congresses.

Applicants should be ENT-specialists (age-limit 45 years) and members of ERS. Applications should be made on-line and include:

- CV and scientific background including previous publications and grants
- Affiliation and members of the working group
- Letter of recommendation from own head of department
- Title, concept and goals of the planned research-project
- Detailed description of the financial costing of the grant

Please send applications with the documents by email to: rhinologysecretary@amc.uva.nl

Deadline: January 1st, 2013