ASTHMA AIRWAY REMODELING

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➢ Describe what is asthma remodeling
➢ Describe the evidence of asthma remodeling
➢ Describe the pathophysiology of asthma remodeling
➢ Describe some treatment options
First described in 1922 by Hubert and Koessler in cases of fatal asthma

Histological abnormalities in the airways due to constant inflammation

Long term inflammation leads structural changes of the airways

Impacts all cases of asthma in small and large airways (not just severe cases)
Determining asthma severity and airway remodeling

- FEV1
- Symptom scores (intermittent, mild, moderate, and severe persistent asthma)
- Fixed airway obstruction
- High resolution CT reveals thickened airways that increase in proportion to disease severity
- Bronchial biopsies

Other noninvasive measures may be useful for serial monitoring in severe asthma
COMPONENTS OF AIRWAY REMODELING

- Structural changes include:
  - Loss of epithelial integrity
  - Thickening of basement membrane
  - Subepithelial fibrosis
  - Goblet cell hyperplasia
  - Sub mucosal gland enlargement
  - Increased smooth muscle mass
  - Decreased cartilage integrity
  - Increased airway vascularity
Evidence for remodeling

Peter K. Jeffrey, MD

- Reviewed the histologic and electron microscopic images
- Epithelial damage, thickening of the reticular basement membrane, increases in glands and goblet cells, increases in the number and volume of vessels, and increases in airway smooth muscle
- Reported showing an 18% increase in myofibroblasts 24 hours after allergen challenge
  - New muscle growth by this mechanism
  - Convincing evidence for injury, inflammation, and remodeling, with increases in wall thickness, and important changes in myofibroblasts and airway smooth muscle

<table>
<thead>
<tr>
<th>Features of Airway Remodeling</th>
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<tr>
<td><strong>Goblet and mucous gland hyperplasia:</strong></td>
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<tr>
<td>• Increased sputum</td>
</tr>
<tr>
<td>• Airway narrowing</td>
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<tr>
<td>• Airway wall thickness</td>
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<tr>
<td><strong>Increased smooth muscle mass:</strong></td>
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<tr>
<td>• Inflammation/fibrosis products</td>
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<tr>
<td>• Asthma severity</td>
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<tr>
<td>• Airway narrowing</td>
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<tr>
<td>• AHR</td>
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<tr>
<td><strong>Angiogenesis:</strong></td>
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<tr>
<td>• Airway wall oedema</td>
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<tr>
<td>• Mediators delivery</td>
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<tr>
<td><strong>Subepithelial fibrosis:</strong></td>
</tr>
<tr>
<td>• Severity of asthma</td>
</tr>
<tr>
<td>• AHR</td>
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<tr>
<td>• Protective against BC (?)</td>
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<tr>
<td><strong>Epithelial alteration:</strong></td>
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<tr>
<td>• AHR</td>
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<tr>
<td>• Decrease protective barrier</td>
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Why does this occur?

- Duration of asthma
- Greater use of medications
- Chronic inflammatory process which leads to activation of
  - Eosinophils
  - Neutrophils
  - Mast cells
EPITHELIAL ALTERATION AND CLINICAL IMPACT

- Destruction of ciliated cells
- Epithelial shedding
- Goblet hyperplasia
  - Increased sputum
- Clinical implications
  - Loss of physical barrier against inhaled allergens
  - Loss of epithelial surface area
  - Airway narrowing
  - Airway wall thickness
Subepithelial fibrosis

- Reported in all cases of asthma
- Basement membrane thickening
- Fibrosis resulting in increased deposition of extra-cellular matrix proteins by fibroblasts
- Associated with asthma severity and increased airway wall thickness

SUBEPITHELIAL FIBROSIS AND CLINICAL IMPACT
Respiratory airway smooth muscle is the critical modulator of airway tone

Asthmatic airways
- Smooth muscle mass is increased due to increase in the size (hypertrophy) and number (hyperplasia)
- Smooth muscle cells migrate to the subepithelial area of the asthmatic airways increasing in size
- Smooth muscle cells participate in the inflammatory and remodelling processes
  - Release of proinflammatory mediators such as cytokines and chemokines
- Chemokines have the ability to induce human airway smooth muscle cell migration and to increase their contractility
- Contribute to the overall airflow obstruction in these patients
Vascular alterations occur leading to increased size of airway wall vessels and angiogenesis

Airway wall edema
- Changes in the airway wall
- Greater expression of vascular endothelial growth factor

Clinical consequences of airway wall angiogenesis
- Reduced airway caliber via airway wall edema
- Increased inflammatory response and remodeling

ANGIOGENESIS AND CLINICAL IMPACT
In asthmatic adults and children

- Goblet cell hyperplasia and submucosal gland hyperplasia are seen
- Consequences of these abnormalities mostly result
  - Increased sputum production
  - Airway narrowing due to sputum secretion
  - Increased airway wall thickness

GOBLET CELL AND MUCOUS GLAND HYPERPLASIA AND CLINICAL IMPACT
• Airway cartilage is an important determinant of airway wall stiffness and integrity

• Decreased cartilage volume and increased cartilage proteoglycan degradation

• Reduced cartilage integrity results in a more bronchoconstriction

LOSS OF CARTILAGE INTEGRITY AND CLINICAL IMPACT
Asthmatic individuals are characterized by airway inflammation:

- Eosinophils, mast cells and lymphocytes migrate to airways
- Release of mediators
  - Bronchoconstriction, mucous secretion and remodeling
- Correlates with airway hyper responsiveness (AHR)
- Mediators such as IL-11 and IL-17 have potent remodeling properties
- Histamine participates in airway remodeling through increased fibroblast
Effect of airway remodeling on lung function

- Structural alterations and inflammatory process are related to the magnitude of functional abnormalities in asthma.
- Airway remodelling has significant effects on lung function and is believed to be responsible for the chronicity of asthma.
Allergic rhinitis is a risk factor for asthma and airway remodelling

Patients with rhinitis have a three fold chance of getting asthma

Rhinitis control improves asthma control

Montelukast and antihistamines are the mainstay therapy in rhinitic subjects with improving asthmatic severity and management
Immunotherapy is reserved for moderate to severe allergic rhinitic subjects

- Immunotherapy reduces inflammatory cell recruitment and activation
- Reducing allergen sensitivity leads not only to relief of rhinitis but also helps control asthma
- Beneficial effects of rhinitis treatment on asthma is reduction of airway inflammation
- Whether reduced inflammation leads to diminished airway remodeling and less asthma expression is still unknown
- More research is required
Corticosteroids

- Control of persistent asthma is best achieved early with ICS use
- ICS have the potential to influence remodeling of airways
- Inflammatory mediators can be reduced by use of corticosteroids
- Antiproliferative effect of corticosteroids
  - Benefit asthmatic patients by reducing smooth muscle mass

ICS form the basis of asthma therapy and the best way to control airway inflammation

More studies are to be performed as there isn’t quite convincing evidence of its effect on remodelling
Montelukast is a leukotriene receptor antagonist is commonly used in asthma therapy and rhinitis

Asthmatic patients with nasal polyposis treated with Montelukast
- 70% improvement in nasal symptoms
- 60% to 90% improvement in clinical asthma scores

Montelukast decreases sputum eosinophils after allergen challenge

Anti-inflammatory effects may play an important role in the pathogenesis of airway remodeling

ANTAGONIST LEUKOTRIENES (VERY PROMISING)
Montelukast has been shown to significantly inhibit:
- Smooth muscle hyperplasia
- Mucus gland hyperplasia
- Subepithelial fibrosis

Decreased lymphocyte and myofibroblasts count in the airways after only eight weeks of Montelukast treatment.

Animal and human studies indicate that anti leukotrienes may prevent airway remodeling of goblet and smooth muscle cell hyperplasia and subepithelial fibrosis.

Long-term studies are needed to confirm the clinical outcomes of the anti remodeling effect of leukotriene receptor antagonist.

ANTAGONIST LEUKOTRIENES (VERY PROMISING)
Treatment with anti-immunoglobulin (Ig) E or (Omalizumab)
- Reduces blood Ig E and decreases asthma symptoms
- Treatment with anti-IgE (e.g. omalizumab) significantly reduced both sputum and tissue eosinophilia as well as IgE-positive T and B cells
- Had very little effect on AHR in asthmatic patients
- Anti-IgE treatment has been shown to reduce levels of the circulating cytokines IL-5 and IL-13
- Improve lung function with moderate to severe allergic asthma requiring daily administration of corticosteroids
- Currently, little effect of anti-IgE treatment on the airway remodeling process has been reported
- Beta-2 agonists (Albuterol) reduces airway muscular tone and improve expiratory flows

- There is little evidence that beta-2 agonists affect airway remodeling

- One study by Orsida shown that Salmeterol with ICS can effectively reduce the number of vessels in their lamina propria compared with patients treated with corticosteroids alone

- Theophylline has no evidence supporting its influence on airway remodeling

BETA-2 AGONIST AND THEOPHYLLINE
Inflammatory process remains the primary target of new drugs used in the future prevention and treatment of asthma.

Current therapies are not designed to specifically treat the underlying remodeling process.

Four treatment options are under investigation:

- **Bacille Calmette-Guérin vaccines (vaccine for tuberculous)**
  - Capable of switching the immune response thus reducing inflammation and remodeling.
- **IL-5 antibodies** target eosinophil-mediated inflammation.
- **Rapamycin**, a macrolide analogue
  - Immunosuppressive effects and may influence inflammation and remodeling in experimental mouse models of asthma.
- **Phosphodiesterase (PDE) inhibitors**
  - Reported to have bronchodilatory, anti-inflammatory and potential anti-remodeling properties.
Bronchothermoplasty is a novel mode of intervention that consists of applying an electric current to segmental and subsegmental bronchi. The goal of destroying the smooth muscle reducing its capacity to contract. Further studies are needed to determine the utility of this treatment but it has been shown to alter airway structure in a possibly beneficial way.
Bronchial thermoplasty for severe refractory asthma

Bronchial thermoplasty involves delivery of radiofrequency energy to the airway wall, which ablates the smooth muscle layer, lessening bronchoconstriction and improving symptoms.

Treatments are done in three separate procedures, with meticulous mapping of the areas treated. The right lower lobe is treated in the first procedure, the left lower lobe in the second, and the two upper lobes in the third. The right middle lobe is not treated.

Cross-sectional photograph of a treated airway from a patient with lung cancer resected 20 days after treatment. Magnification: 40x.

Trichrome-stained section at higher magnification (400x). Airway smooth muscle is largely absent to the left of the arrow.

The thermoplasty device within the airway with the electrodes deployed.

Medical Illustrator: Mark Sabo
The AIR2 trial: Effect of bronchial thermoplasty and sham thermoplasty on health care utilization. Health care utilization in the 12 months after real or sham thermoplasty.

