Chronic inflammation and frequent exacerbations in patients with COPD: the role of PDE4 inhibitors

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Summary

Inflammation represents an important hallmark of several diseases. In COPD the inflammatory lung response may be chronic and dysregulated with accumulation of inflammatory cells that causes lung parenchyma damage and airway remodeling. The inflammatory cascade involves different cells, mediators, proteins, oxygen species that leads the local inflammation to become systemic and promotes comorbidities and complications.

Different stimuli may worsen the COPD pathology, including the well known COPD exacerbation, a peak of local and systemic inflammation. Also hypoxia may stimulate inflammatory genes worsening local and systemic level of inflammation.

Understanding the inflammation can allow us to differentiate COPD phenotype and lead us to personalize the therapy. Several have the specific target of inflammation, particularly PDE4 inhibitors may have a role in certain COPD phenotype.

KEY WORDS: chronic obstructive pulmonary disease, COPD, inflammation, phosphodiesterase-4, roflumilast, acute exacerbation.

Dysregulated and prolonged inflammation as a pathologic process in COPD

Inflammation, a response triggered by damage to living tissues, represents the hallmark of several lung diseases. A correct inflammatory response is a defense mechanism to protect individuals from infections and injuries. Nevertheless, the inflammatory response may be dysregulated and prolonged becoming itself a cause of disease, as we can find in several chronic lung diseases. It is the case of chronic obstructive pulmonary disease (COPD) where the airways, in particular the peripheral ones, lung parenchyma and pulmonary blood vessels are common places of altered prolonged inflammatory response (1). Chronic inhalation of irritants leads to the recruitment and accumulation of inflammatory cells, such as neutrophils, T-lymphocytes (CD8+) and macrophages. (2) have been demonstrated in airways of COPD patients and this producing as main effects airway remodeling and lung parenchyma destruction. Both innate immunity with neutrophils, macrophages, eosinophils, NK cells, gamma delta cells and dendritic cells and adaptive immunity with T and B cells have roles in COPD together with the activation of structural cells as epithelial cells, endothelial cells and fibroblasts (2).

Increased number of macrophages, have been found in COPD patients airways, up to 5-10 fold and the number of macrophages has been related to emphysema entity (2). Once activated by cigarette smoke, macrophages may orchestrate the chronic inflammatory response with release of both inflammatory mediators and chemokines that amplify inflammation, of reactive oxygen species, of elastolytic enzymes, of smooth muscle constrictors, of mucus gland activators and of matrix metalloprotease enzymes (MMPs). Neutrophils are the most represented cells, their number has been related to COPD severity, and they are found mostly in patients sputum and BAL probably because of their rapid transit and less in the airway walls. Smoking has direct stimulatory effect on granulocyte production, release from bone marrow and on their survival. Neutrophils are directed to airways because of several chemokines secreted by different cells as macrophages, T cells, epithelial cells and the same neutrophils activated in the airways of COPD patients. These cells are source of reactive oxygen metabolites, inflammatory cytokines, lipid mediators, antibacterial peptides and tissue damaging enzymes as serine proteases, cathepsine G, proteinase-3 and matrix metalloprotease (MMP-8, MMP-9). Neutrophil products also induce mucus hypersecretion by both an acute secretagogue effect and by augmentation of the bronchial mucus producing apparatus. Also lymphocytes, mostly CD8+ cell, eosinophils and mast cells found in both the large and small airway walls (2).
All these cells cooperate to the inflammatory cascade that triggers the release of inflammatory mediators such as tumor necrosis factor-α (TNF-α), interferon-γ (IFN-γ), interleukins (IL-1, IL-6, IL-8) and fibrinogen (3). These mediators sustain the inflammatory process and lead to tissue damage and different systemic effects.

The oxidative agents are implicated in both the generation of mucus metaplasia in chronic bronchitis and in the destruction of the lung tissue in emphysema. Local inflammation pathway is composed by reactive oxygen species (ROS), a decreased level of alpha 1 antitrypsin, increased H2O2, NO and S-nitrosothiol level and several damages to lipid membrane of epithelial cells. ROS may cause injury to the epithelial and endothelial cells through membrane lipid peroxidation, mitochondrial damage, and endoplasmic reticulum stress resulting in unfolded protein response (UPR). Chronic inflammation leads to various structural changes in the lung, which further perpetuate airflow limitation (Figure 1). The disequilibrium between the oxidant agents and antioxidant agents leads to inflammation and causes the typical airway disease of COPD patients (4). In fact, inflammation of the airways and emphysema with alveolar attachments disruptions in small airways increase airway resistance, rise the flow obstruction and dynamic hyperinflation, increases the work of the diaphragm and all this factors producing the clinical symptoms of dyspnea.

The role of PDE4

The inflammatory cascade involves different messenger proteins that regulate the inflammatory process. In the lung, the second messenger cyclic AMP (cAMP) is involved in the regulation of inflammatory cells, mucociliary clearance and pulmonary vascular remodeling (6). It suppresses immune and inflammatory cell activity (in inflammatory cells such as neutrophils, T-lymphocytes and macrophages), relaxes airway smooth muscle and modulates pulmonary nerve activity (7). In turn, an elevated concentration of intracellular cAMP activates protein kinase A, which enhances phosphorylation of proteins may inhibit numerous inflammatory cell functions, e.g. proliferation and release of cytokines and chemokines, reactive oxygen species, arachidonic acid metabolites, chemotaxis and proteases. cAMP modulates human airway epithelial cells that might support mucociliary clearance, human lung fibroblasts to potentially prevent a fibrotic response and the proliferation of pulmonary artery smooth muscle cells thereby reducing pulmonary vascular remodelling (8).

The enzyme phosphodiesterase 4 (PDE4) degrades cAMP into its inactive form, thereby stimulating the activity of inflammatory cells.

The role of local inflammation in COPD

The local inflammation of the lung generates a systemic inflammatory response that subsequently becomes chronic, leading to comorbidities and complications with an increase of cardiovascular risk, loss of muscle mass, decrease of bone density. An important effect of systemic inflammation is endothelial dysfunction. The main function of the endothelial cells is to maintain homeostasis of the vascular wall by adjusting the tone, permeability, adhesion of inflammatory cells, platelet aggregation and to prevent the activation of the coagulation cascade (Figure 3). Nitric oxide has a central role in endothelial function through the powerful vasodilatory, anti-inflammatory, anti-platelet and anti-proliferative effect, while to endothelin should be given the role of the most potent endogenous vasoconstrictor. An imbalance between these two factors with the possible in-
volvement of other factors, determines the endothelial dysfunction (5). This condition is associated with atherosclerosis and several cardiovascular diseases such as hypertension, coronary syndromes, diabetes and chronic renal failure. A number of comorbidities can be explained with this mechanism. In fact to describe the comorbidities of COPD we could use the term “chronic inflammation”.

Figure 2 - Effect of PDE4 on inflammatory cells.

Figure 3 - Endothelial dysfunction in COPD.
The role of hypoxia

Hypoxia is an important consequence of pulmonary disease. Results of chronic hypoxia are well known, but also the intermittent hypoxia has been recently shown to play an important role. In fact intermittent hypoxia reduces the resistance and endurance of the exercise, acts on systemic inflammation and on kidney function reducing renal blood flow and producing sodium retention (27). Based on studies of sleep apnea syndrome, the intermittent hypoxia shows an increase of two main factors: HIP and NF-kB (Figure 4).

A down-regulation by hypoxia is seen in genes involved in cytoskeleton maintenance (Rho kinase), mRNA processing (heterogeneous nuclear ribonucleoprotein H1 and splicing factor) and DNA repair (REV3).

Hypoxia-inducible factor (HIF) -1 is a heterodimeric transcription factor, that can bind and regulate the expression of genes involved in metabolism (adenylate kinase 4, galactokinase) and apoptosis (galectin-3 and gelsolin). A down-regulation by hypoxia is seen in genes involved in cytoskeleton maintenance (Rho kinase), mRNA processing (heterogeneous nuclear ribonucleoprotein H1 and splicing factor) and DNA repair (REV3). All these genetic regulations cause a proinflammatory response induced by hypoxia and promote enhanced survival of myeloid inflammatory cells, such as granulocytes, monocytes and macrophages, resulting in their functional longevity and inflammatory response.

Inflammation and acute exacerbations of COPD

COPD exacerbations is defined generally as changes in respiratory symptoms. Focusing on inflammation, COPD exacerbation could be defined as a peak of inflammation during the course of the disease. Serum biomarkers such as various cytokines, adipokines, C-reactive protein and coagulation factors are elevated during exacerbations.

An increase of lung inflammation, as during the exacerbation, worsens the systemic inflammation that leads to an increase in major acute events, especially cardiovascular events. The same happens in pneumonia in the post-acute period (9). Exacerbations also are related to means functional decline, symptoms increase, quality of life worsening, increased risk of hospitalization and mortality, and exposes the patient him/herself to relapsing further exacerbations. The prognosis becomes unfavorable in most patients who have exacerbations especially if they require hospitalization. The detection of higher blood biomarkers during exacerbation compared with baseline, such as CRP, IL-8, TNF-α, leptin, endothelin-1, eosinophil cationic protein, myeloperoxidase, fibrinogen, IL-6, α1-antitrypsin, leukotrienes E4 and B4 (Table 1) confirm that exacerbation deals with a systemic peak of inflammation.

Figure 4 - Intermittent hypoxia and inflammation.
All these molecules bind and trigger the activation of vascular endothelium on luminal surface of the vessels and promote endothelial dysfunction and atherosclerosis (10). This leads to a destabilization of atherosclerotic plaque and cardiovascular complications.

Because of the pivotal role of inflammation in the genesis of systemic comorbidities we should act on it and particularly on exacerbations. Several studies have been done to find solutions to improve the outcomes. The bronchial epithelium and endothelium are a target reached by inhalation therapy. The bronchodilators have an important role in exacerbations prevention as the fixed combination β-2 agonists and steroids (TORCH study), the tiotropium (UPLIFT study), the fixed combination with tiotropium (INSPIRE study). On inflammation is also acting the inhibitor of phosphodiesterase-4 Roflumilast, an available drug which acts either on CD8 cells, macrophages, neutrophils, epithelial cells and smooth muscle. The mechanism of this drug is to prevent the conversion of 3'-5' cyclic AMP to AMP. It increases the amount of 3'-5' cyclic AMP in the cell and modulates the phosphorylation of the inflammatory proteins (Figure 5).

The rationale for PDE4 inhibitors in COPD patients with chronic inflammation

Differently from asthma, inhaled corticosteroids (ICS) shows a limited response in COPD (11). The low number of glucocorticoids receptors (GR) in neutrophils may explain, in part, why inflammation in COPD is insensitive to glucocorticoids as monotherapy. Furthermore, COPD inflammation is different from asthma one making steroid-resistance much more frequent (12). The reduced effect of ICS in COPD seems due to the marked reduction in histone deacetylase 2 (HDAC2), the nuclear enzyme that corticosteroids require to switch off activated inflammatory genes (13), rendering these patients resistant to the effects of ICS. The reduction in HDAC2 is thought to be secondary to oxidative stress, both independent and by activation of phosphoinositide-3-kinase-α (PI3Kα). Inhibition of PI3Kα has recently shown to restore corticosteroid sensitivity in mice and may hold therapeutic promise (14, 15). Marwick et al. (16) have shown that formoterol reverses oxidative stress-induced corticosteroid insensitivity via PI3Kα. Low-dose theophylline has shown to enhance the antiinflammatory effects of steroids during exacerbations of COPD (17) and seems to have the capacity to restore the reduced HDAC2 activity in COPD macrophages (18). Roflumilast has also been shown to increase the ability of formoterol to enhance glucocorticoid-dependent gene transcription in human airway epithelial cells (19). A large part of pre-clinical research has shown that PDE4 inhibition has the potential to target the three main components of COPD: bronchoconstriction, mucus hypersecretion and airway remodeling. As PDE4 is the major cAMP-metabolizing enzyme, the inhibition of PDE4 suppresses the inflammatory response and the epidermal growth factor receptor-induced Mucin 5AC over-expression, that directly inhibits mucus production. Inhibition of PDE4 may also lead to minimization of airway remodeling by suppressing the release of TNF-α (20).

The PDE4 inhibitor Roflumilast is now indicated for maintenance treatment of severe COPD (FEV1 post-bronchodilator less than 50% predicted) associated with chronic bronchitis in adult patients with a history of frequent exacerbations as add on to bronchodilator treatment. In this patient population, Roflumilast 500µg once-daily may reduce exacerbation rate and improves lung function.

Are all COPD a same phenotype based on inflammation?

Several guidelines define COPD as an inflammatory disease, but there are few or no specific recommendations on the assessment and targets for reduction in inflammation, as these are not established (21-23). Nevertheless, there is for sure a COPD phenotype where the inflammation is predominant and a potential target of therapy (24). A phenotype which needs more attention for its burden and that represents a population that completely cross the GOLD classification is the so-called “frequent exacerbator” of the ECLIPSE study.
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flammation with higher levels of sputum interleukin IL-6 and IL-8 even in the stable state (26). We have to take in account this population and identify them, because of the worst prognosis and quality of life. The aim of future investigations will be the characterization of COPD phenotypes physiopathologically and therapeutically for a better prognostic assessment and therapeutic management (24, 27).

References

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