Transepithelial exit of leucocytes: inflicting, reflecting or resolving airway inflammation?

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ABSTRACT
The passage of infiltrated tissue granulocytes across airway epithelium into airway lumen is generally considered to be a pathogenic process in asthma and chronic obstructive pulmonary disease (COPD). An alternative hypothesis is proposed here—namely that the transepithelial egression of infiltrated leucocytes acts to rid diseased airway tissues of proinflammatory cells efficiently and non-injuriousively. Several clinical observations previously discussed as ‘unexpected’ and ‘puzzling’ support this hypothesis. In acutely resolving allergen challenge-induced inflammation, in patients with mild asthma, airway wall eosinophils disappear without evidence of apoptosis but with evidence of a developing airway lumen eosinophilia. In the same postchallenge resolution phase, lymphocytes, neutrophils and mast cells exhibit peak numbers in airway lumen of individuals with asthma. In severe asthma requiring intubation, clinical improvement is similarly paralleled by a marked increase over several days in airway lumen neutrophils. Increased numbers of sputum neutrophils and lymphocytes also occur as symptoms improve in COPD over several months after smoking cessation.

Conversely, when the transepithelial exit of leucocytes has been inhibited in inflamed animal airways the inflammation in the airway wall has been much aggravated. Finally, transepithelial egression of numerous granulocytes and lymphocytes clearly can occur without any harm to airway epithelial barriers. The present hypothesis of ‘resolution through egression’ provides a novel interpretation of common airway lumen data, cautions against administration of agents that impede leucocyte egression in inflammatory airway diseases and infers new approaches in disease resolution research.

INTRODUCTION
Although both the airway lumen and airway wall are designated as in vivo locations they differ dramatically. Only the wall is under the dynamic influence of nerves and microcirculation. Indeed, the human airway mucosa harbours a particularly profuse network of capillaries and venules. Throughout the nasal passages and stretching all the way from the trachea to the smallest bronchioli these microvessels occupy the area just beneath the epithelial lining. When required, the subepithelial microcirculation promptly delivers plasma proteins and leucocytes locally anywhere along the tracheobronchial tree. After extravasation the cells remain in the airway wall for variable lengths of time. Here we discuss the further fates of the airway wall leucocytes especially in association with resolution of inflammatory airways diseases.

The accessible leucocytes in the airway lumen have been extensively studied in diseases such as asthma and chronic obstructive pulmonary disease (COPD). The lumen cells are generally considered to reflect the intensity of cell-mediated inflammatory processes in the diseased airway wall. However, caution is advised in interpretation of airway lumen findings since airway wall and lumen may differ as to which granulocyte, eosinophil or neutrophil,1 and which T lymphocyte, especially Tc1 or Tc2,2 is predominant. A further caveat concerns the possibility that leucocytes rise and fall in an opposite direction in the airway wall compared with the airway lumen. Here we discuss data suggesting that exit of leucocytes into the airway lumen is not the pathogenic process it is assumed to be. We argue that increased lumen leucocytes simply reflect the fact that the cells have been cleared from the diseased airway wall as inflammation resolved. This idea is not novel. In ancient medicine therapeutic purging of the airways was advocated, and authors active in ~19003 saw profuse discharge of eosinophils into sputum as a sign of recovery from severe asthma episodes.

CLINICAL IMPROVEMENT AND TRANSEPITHelial ELIMINATION OF AIRWAY WALL LEUCOCyTES
To some extent transepithelial migration of leucocytes into the airway lumen (‘luminal entry’, ‘transepithelial egression’, ‘transmigration’) is a continuous process. Several types of cells thus migrate to the airway mucosal surface to carry out immune surveillance, sentinel, defence and repair duties. For this overview we were looking for observations indicating that the transepithelial migration could also act to rid diseased airway tissues of proinflammatory cells. During development of airway inflammation newly recruited cells may accumulate in the mucosal tissue rather than moving on, into the airway lumen. But what happens when inflammation resolves? Allergen challenge in allergic subjects with asthma produces bronchial inflammation and symptoms that may resolve within 24 h of challenge. In such allergen-challenged subjects,4–6 and in animal models,7 8 airway lumen eosinophils have increased markedly during resolution as eosinophils evidently have disappeared from the airway wall.4–6 Hence, a negative correlation between airway wall and airway lumen cells was observed.4 Similarly, other leucocytes including lymphocytes and neutrophils have peaked in the airway lumen during the resolution phase after allergen challenge in human asthma.9 Further, increased numbers of mast cells in the airway lumen correlated with the magnitude
of the allergen challenge-induced late phase asthma reactions recorded several hours earlier. There are no known data demonstrating that the airway wall inflammatory cells disappeared by mechanisms other than transmigration into the airway lumen. Thus, observations in human and animal asthma in vivo models support a resolving role of transepithelial migration of inflammatory cells.

Severe asthma is characterised by airway wall neutrophilia. Of interest here is that in severe asthma, requiring intubation, the number of airway lumen neutrophils increased markedly over several days as the asthma improved prior to extubation. Further support for a role for transepithelial egression as a mode of ridding diseased bronchial tissues of inflammatory cells comes from smoking cessation studies. Thus, symptom improvement over several months following smoking cessation in COPD was associated with sputum neutrophilia and increased sputum lymphocytes. Also, sputum eosinophilia has been associated with improved lung function in patients with asthma treated with β-agonists. Increased leucocyte numbers in the airway lumen might not be such a bad sign after all.

**ACCELERATED TRANSEPITHELIAL MIGRATION IN CHRONIC DISEASE**

Neutrophilia in the airway lumen in stable COPD can be associated with lack of neutrophilia in the airway wall. Similarly, patients with COPD with chronic bronchitis have been characterised by low airway wall eosinophil counts but high lumen eosinophil numbers. Chronic conditions such as these may thus be characterised by a degree of accelerated transepithelial migration of neutrophils and eosinophils, respectively. Stopping de novo recruitment of leucocytes to the airway wall may be an effective therapeutic intervention in these patients. In severe as compared with mild COPD, airway wall mast cells were reduced. This occurred without increased apoptosis of these cells but with increased numbers of mast cells in the airway lumen. These data, together with findings in asthma, suggest that airway wall mast cells are eliminated through transepithelial migration.

**DEATH OF EOSINOPHILS AND LYMPHOCYTES IN THE AIRWAY WALL, AND STEROID TREATMENT**

Indirect support for the role of egression comes from the lack of any compelling evidence to show that airway wall leucocytes are eliminated through other mechanisms. We and others have thus failed to demonstrate a role for apoptosis, the so-called ‘silent death’, in resolution of airway wall eosinophilic inflammation. Airway wall eosinophils have increased and declined without occurrence of detectible apoptotic eosinophils, whether phagocytosed or not. Instead, we observed that primary lysis, without prior apoptosis, is a major cause of eosinophil death in vivo. This fate led to clusters of free eosinophil granules being scattered in human diseased airway mucosa. Hence, eosinophil death in vivo, through cytolysis, causes rather than resolves airway inflammation. In agreement with this, its occurrence is reduced by steroid treatment. In vitro, corticosteroids can increase apoptosis of eosinophils and lymphocytes and they may stimulate phagocytic elimination of apoptotic leucocytes. However, such acknowledged ‘proresolution’ effects remain to be demonstrated in the diseased airway wall. The reputed eosinophil apoptosis-inducing effect of glucocorticoids has so far not been borne out in vivo studies on airway tissues, nor could steroid-induced T cell apoptosis be consistently demonstrated in biopsies obtained from patients with asthma and COPD. Apoptosis of leucocytes in the airway lumen is a different matter. However, as with the findings in the airway wall, airway lumen data may not establish the resolving role of apoptosis that we and others have been seeking (see above).

**IMPEDING TRANSEPITHELIAL EXIT OF LEUCOCYTES AGGRAVATES AIRWAY WALL INFLAMMATION**

It has been suggested that the induction of eosinophil apoptosis may be of benefit in allergic airway diseases. A seminal study in this field examined Fas-induced eosinophil apoptosis in the airway lumen in allergic mice. Later studies demonstrated that impeded transepithelial egression of granulocytes could explain the ‘promising’ reduction in airway lumen eosinophilia found in these Fas-treated animals. Many apoptotic inflammatory cells that accumulated in the airway wall were not engulfed and underwent necrosis. Indeed, airway wall inflammation was not improved by the induced apoptosis but was much aggravated. Inhibitors of intercellular adhesion molecule-2 (ICAM-2) or matrix metalloproteinase-2 (MMP-2) and MMP-9 have also reduced transepithelial migration of leucocytes and caused aggravation, and even lethal, pulmonary effects in allergic mice. Viral infection causes exacerbations of asthma and COPD. Hence, it is of interest that in rats with pre-existing virus-induced inflammation allergen challenges produced persistent eosinophilia in the airway wall but not in the airway lumen, and was associated with loss of lung elastic recoil. In contrast, allergen-exposed control rats had fewer eosinophils in the airway wall but more in the airway lumen and no loss of elastic recoil. Hence, viral infections may impede transepithelial egression of inflammatory cells, thus worsening the effects of allergen exposure. Virus-induced inhibition of egression, a major component of host defence, would also have promoted the virus infection.

Giving an antibody to block interleukin 5 (IL-5), a regulator of eosinophilopoiesis, eosinophil migration and eosinophil survival, eliminated both blood and airway lumen eosinophilia but had little effect on airway wall eosinophil numbers. Since no proapoptotic effect has been demonstrated, the persistant airway wall eosinophilia could well reflect attenuated egression. Recent attempts to produce beneficial effects in COPD and asthma by reducing the traffic of granulocytes include the use of chemokine receptor (CCR) inhibitors, but the possibility that such drugs may fail due to reduced egression of leucocytes from the diseased airway wall probably needs to be considered.

**MUCOSAL TISSUE CHEMOTRACTION REDUCES EGRESSION BY RETAINING LEUCOCYTES IN THE AIRWAY WALL**

Airway wall neutrophilia occurs following steroid treatment of COPD and asthma. Based on findings in vitro in 1996, this frequent observation has been attributed to inhibition of neutrophil apoptosis. Morphological features best define apoptosis, so it is significant that Gicyzcki et al were unable to find any effect of steroid treatment on neutrophil apoptosis compared with placebo treatment when studying the ultra-structure of neutrophils in COPD biopsy tissues. These authors concluded that ‘the functional significance of the potential for steroids to reduce the clearance of neutrophils by their effect on apoptosis is unclear in vivo and requires further study’. Rather than modulating neutrophil apoptosis, we suggest the published data indicate that steroids reduce transepithelial egression of neutrophils (in contrast, steroid treatment permits transepithelial exit of eosinophils). In mild asthma, steroid
treatment has caused upregulation of neutrophil-attracting chemokines such as IL-8 and interferon γ-inducible protein 10 in bronchial tissues. In severe exacerbations of asthma, CXCL5-dependent mechanisms, possibly induced by steroid treatment, may also contribute to recruiting and retaining the neutrophils in the airway wall. Finally, in contrast to the steroid-induced airway wall neutrophilia, reduced numbers of airway lumen neutrophils have been demonstrated following steroid treatment of neutrophilic airway conditions. One study involving steroid-treated patients with COPD has actually demonstrated both reduced neutrophils in the airway lumen and increased neutrophils in the airway wall. Together these clinical findings suggest that steroid treatment reduces transepithelial egression of neutrophils and that this occurs through upregulation of mucosal tissue chemoattractants. Severe exacerbations of COPD exhibit up to a 100-fold upregulation of neutrophil chemokines in bronchial mucosal tissues. Neutrophil chemotraction in the airway lumen in COPD may be low and neutrophils in severe COPD exhibit reduced chemotaxis compared with neutrophils in mild COPD. Thus, several factors may act in concert to modulate transepithelial egression of neutrophils in severe COPD.

Although inhaled steroids have broad anti-inflammatory effects and may in some cases worsen respiratory infections, they appear to spare several innate immune responses of airway microcirculation and epithelium. Steroids may also help maintain innate immunity by not reducing the profuse traffic of neutrophils to the airway epithelium and surface at viral infection and associated epithelial injury—repair processes. Nitrogen dioxide, a major air pollutant, is associated with an increased susceptibility to respiratory infection as well as with loss of airway wall neutrophils. The latter effect may reflect increased transepithelial egression since neutrophils in bronchial wash fluids were increased whereas neutrophils in bronchial epithelium were reduced after exposures to nitrogen dioxide. Hence, effects on airway neutrophils are difficult to interpret in simple terms of inflammation. Further studies are warranted to determine the significance of steroid-induced airway wall neutrophilia in asthma and COPD.

**EGRESSION OF LEUCOCYTES WITHOUT HARMING THE EPITHELIAL LINING**

The passage of granulocytes across the epithelium in asthma and COPD is thought to cause epithelial derangement of pathogenic importance. This view is debatable. For example, reduced bioelectrical resistance of epithelial cultures, commonly considered evidence of transmigration-induced barrier derangement, cannot be equated with physiologically important barrier functions in vivo. Also, epithelial transmembrane junction proteins, including occludin and E-cadherin, can be lost at transmigration of neutrophils without causing increased permeability. Finally, association between granulocytes and epithelial injury may not indicate cause and effect but simply reflect the potent stimuli that damaged cells provide for recruiting activated neutrophils and eosinophils to the repair site and to the airway lumen.

The airway epithelium favours the passage of leucocytes in the physiological basal to apical direction and there is ample evidence of non-injurious transepithelial migration of numerous leucocytes. In anaesthetised guinea-pigs, 35 000 eosinophils/min/cm² surface area thus traversed a normal pseudostratified tracheal epithelium without evidence of damage even at the ultrastructural level. Bronchial instillation of chemoattractants such as leukotriene B4 and bacterial lipopolysaccharide in humans and sheep was associated with marked movement of neutrophils into the airway lumen without epithelial injury. The yielding plasticity of paracellular junctions clearly allows entry of cells and macromolecules into the airway lumen without increasing epithelial permeability in the reverse direction. Lymphatic transport of infiltrated leucocytes and extravasated plasma proteins occurs but is quantitatively of less importance than the egression—exudation processes.

**MECHANISMS INVOLVED IN TRANSEPITHELIAL EGRESS**

Little is known about the molecular regulation of transepithelial loss of leucocytes during the resolution of airway inflammation. Extrapolating the extensive knowledge of mechanisms concerning the passage of leucocytes across the venular endothelial lining to mechanisms regulating the transepithelial egression of leucocytes is unwise since the two crossings are in opposite directions: the venular paraendothelial exit is apical to basal and the epithelial passage is basal to apical. Based on interactions between neutrophils and intestinal epithelial cells and focusing on pathobiology, Chin and Parkos have reviewed molecular mechanisms of epithelial transmigration of leucocytes. Many different adhesive interactions between neutrophils and epithelial basolateral receptors and intercellular junction proteins occur during transepithelial migration. However, details regarding neutrophil transmigration may not apply to other leucocytes. Also, intestinal epithelial mechanisms may not be entirely valid for airway epithelium. Transmigration of neutrophils may thus harm intestinal but not airway epithelial monolayers. Some general features of transepithelial migration of leucocytes are outlined, however, in figure 1.

MMPs may produce transepithelial loss of parenchymal granulocytes in mice by creating transepithelial chemokine gradients. Lumen levels of CCL11 may contribute to non-injurious loss of airway wall eosinophils, and tumour necrosis factor α (TNFα) may promote transmigration of both eosinophils and neutrophils. It is widely assumed that the clearance of CCR7+ memory and effector lymphocytes from inflamed airway tissues involves apoptosis, thus ‘ignoring a potentially very important exit pathway through the bronchial epithelial barrier’ according to Porter and colleagues. They suggested that polarised localisation of CXCL10 and CXCL11 towards the apex of epithelial cells could be key chemokines for elimination of T lymphocytes through egression in COPD. Apical epithelial CCL5 and reduced tissue expression of CCL5 may regulate transepithelial migration of leucocytes in allergic airways. ‘Chemorepellents’ could also contribute to moving leucocytes away from sites of airway wall inflammation.

**ELIMINATION OF LEUCOCYTES FROM THE AIRWAY LUMEN**

The potential role of egression as a means of reducing airway inflammation underscores the importance of cell clearance from the bronchial lumen. Death through apoptosis, followed by phagocytosis, is the topical mechanism of cell elimination. Apoptosis of cells in the airway lumen has been much studied during the last two decades with the assumption that observation on lumen cells would be valid for cells in the airway wall. However, apoptosis of lumen cells cannot reflect apoptosis in the airway wall (‘dead cells do not walk’). In this overview we discuss cell apoptosis in the airway lumen merely to see if there are data indicating involvement of this mechanism in final elimination of the cells already eliminated from the airway wall. A widely quoted, uncontrolled study from 1996 reported that steroid
increase to aid resolution of infection and inflammation'. Matute-Bello and Martin,66 who originally discovered an antiapoptotic action of bronchoalveolar lavage (BAL) fluid in adult respiratory distress syndrome, argued that neutrophil apoptosis may have little to do with outcome. A study testing the hypothesis that the airway lumen milieu in COPD would promote neutrophil survival failed to do so.62 In summary, the challenge of demonstrating a role for apoptosis and phagocytosis of leucocytes remains, whether in airway lumen or in the airway wall.

Leucocyte-driven airway inflammation primarily concerns the activity of leucocytes in the airway wall. Whereas clearing cells across the nasal and bronchial epithelium is likely to be beneficial for most patients, this may not be true for ingress across the alveolar epithelium. However, in studies of lung inflammation in mice it appears that egression of parenchymal leucocytes into the alveolar air space was beneficial and, when trans-epithelial egression was prevented, severe asphyxia resulted.26 Cells can also move from the airway lumen and across the alveolar, but not bronchial, epithelium to enter the lung circulations.67 This escape route for lumen leucocytes may be important for alveolar clearance where mucociliary transport is lacking and coughing ineffective. Egression may not be desirable in diseases where mucociliary clearance is severely impaired. It is particularly important that human peripheral airways are not overwhelmed by leucocyte- and plasma protein-rich exudates that could ruin an already compromised ciliary clearance mechanism and cause small airway closure.45 Employment of non-pharmacological and pharmacological methods to improve airway clearance may be a major component of treatments aimed at resolution of airway inflammation.

CONCLUSION

We argue that the finding of eosinophils, neutrophils, lymphocytes and mast cells in the airway lumen may reflect their successful and non-injurious elimination away from cell-mediated disease areas in the airway wall. Experimental evidence and many clinical observations support the importance of trans-epithelial egression as a mode of ridding diseased airway tissues of inflammatory cells. This possibility needs consideration in studies of airway diseases and when assessing the effects of drug interventions where data on airway lumen leucocytes alone may lead to paradoxical conclusions. Egression-promoting drugs, combined with improved mucociliary clearance, may well have a valuable role in resolving airway inflammation in asthma and COPD.

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