Airway epithelial repair: breathtakingly quick and multipotentially pathogenic

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Abstract
Epithelial shedding, even to the point of airway denudation, had already been described as a common and unifying feature of asthma by the latter half of the 19th century. However, the repair processes that specifically follow the shedding-like loss of epithelial cells have only recently been examined in vivo. This paper discusses the exceedingly fast epithelial restitution and the potential pathogenic sequelae to epithelial shedding alone that have been unravelled. Epithelial cytoprotection emerges as an important property of future therapeutic drugs for the treatment of airways inflammatory conditions.

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The centennial paradigm of epithelial shedding in asthma is based on findings of exfoliated epithelial cells in sputum and bronchoalveolar lavage (BAL) fluid and on microscopic observations of epithelial loss in necropsy and bronchial biopsy specimens. The occurrence of epithelial denudation has provided attractive explanations to phenomena such as non-specific airway hyperresponsiveness. However, such causal links between epithelial damage and the pathophysiology of asthma are far from clear – for example, we still lack information about the extent and distribution of epithelial damage. Indeed, biopsy studies displaying up to 50% epithelial denudation have failed to demonstrate differences between asthmatic and healthy control subjects. Thus, the typical picture of denuded asthmatic bronchi probably involves artifacts inflicted during sampling and/or sectioning. It has been suggested that the airways in patients with allergic diseases harbour a particularly fragile epithelium, based on exaggerated denudation at routine tissue handling. The maintenance of mucosal tightness in patients with eosinophilic airway disease further indicates that a manifest epithelial denudation is too simplistic a description of the mucosa in desquamative diseases. In this paper we discuss recent observations which indicate that epithelial restitution starts instantaneously and proceeds at very high speeds after shedding and denudation. Furthermore, such shedding-repair processes may, by themselves, be partly responsible for the pathophysiology, the leucocyte pathology, and the remodelling that occur in desquamative airway diseases.

Scientific basis
HIGH SPEED RESTITUTION AFTER SELECTIVE LOSS OF COLUMNAR EPITHELIAL CELLS: A NEW ROLE FOR BASAL CELLS
Berkart suggested in 1889 that exfoliation of mainly columnar cells is common in asthma. However, the capacity of basal cells to take over the barrier properties of the epithelium after a selective loss of columnar cells was only established quite recently. In intact human and animal airways it took less than 20 minutes for basal cells that remained after selective removal of the columnar epithelium to become extensively flattened and to cover the basement membrane. At the cell borders “seals” consisting of overlapping cytoplasmic protrusions were formed. The same type of “seals” have also been observed in vivo at sites of epithelial damage induced by allergen challenge. The extreme readiness of basal cells to assume a barrier structure suggests that sacrifice of columnar cells may be regarded as a fully functioning airway mucosal defence mechanism. Releasing columnar cells and covering the basement membrane would thus be important roles for basal cells in airway defence and repair.
Aiway epithelial repair

Figure 1  (A) Whole mount preparation, histochemically stained for epithelial cell borders, showing an overall view of sites of patchy epithelial damage (arrowheads). The detailed morphology of (B) small and (C) large damage sites is shown by scanning electron microscopy. (D) At sites of inflammation induced epithelial damage repair processes are immediately initiated; columnar cells (lightly shaded) and basal cells (darkly shaded) are involved in a continuous sealing of the epithelial lining. The restitution processes are aided by defence and growth promoting tissue responses—for example, extravasation of plasma and recruitment of leucocytes (thick arrows). Epithelial repair processes may also cause hypersecretion, proliferation of resident tissue cells, and increased local lymph node activity (thin arrows).

Thus, the local extracellular milieu will promptly be endowed with plasma-derived molecules with defence and repair-promoting properties—for example, growth factors, cytokines, proteases, and antiproteases. Within a few minutes high numbers of leucocytes also accumulate in the denuded area. Interestingly, the epithelial damage-repair processes activate both eosinophils and neutrophils. Hence, tissue responses evoked by epithelial shedding alone may contribute significantly to the airway pathology in inflammatory diseases of the airway.

TISSUE RESPONSES INDUCED BY EPITHELIAL SHEDDING

The epithelial denudation-restitution events are associated with many tissue responses including hypersecretion, plasma exudation, recruitment and activation of leucocytes, and increased proliferation of fibroblasts and smooth muscle cells. The mucosal microcirculation constitutes potentially the most important response organ during the early acute repair phase. Within minutes after denudation endothelial gaps are formed and a pronounced plasma exudation response is initiated. A gel-like network of fibrin and other attached plasma proteins is formed on the denuded basement membrane.

EPITHELIAL DAMAGE-RESTITUTION IN ALLERGIC INFLAMMATION

Using cell border staining on whole mount preparations to get a good overall image we have recently examined the epithelial changes that occur during allergic inflammation in the large airways of guinea pigs (fig 1). The significant features to emerge from these studies are (1) the patchy distribution of epithelial damage, (2) instantaneous start of epithelial repair processes, and (3) focal localisation of inflammation to the injury-repair sites.
Flattened epithelial repair cells were always present at the patches of injury, irrespective of the stage of epithelial damage. Hence, the sites of epithelial damage induced by inflammatory stimuli are exceedingly dynamic foci where epithelial shedding and repair processes occur simultaneously and where the basement membrane may be kept covered by poorly differentiated restitution cells (fig 1). The prompt start of epithelial repair also agrees with previous findings of a maintained airway absorption barrier in allergic airway diseases. Furthermore, sites of epithelial injury-repair in allergen-challenged airways are associated with a dense accumulation of plasma proteins and leucocytes including an abundance of extracellular eosinophil granules (the latter derived from cytolysis of eosinophils).  

**Therapeutic potential**

The notion that epithelial shedding-restitution processes alone are responsible for pathological changes in asthmatic airways offers new treatment options. Epithelial cytoprotective activity of future drugs would thus be expected to have beneficial effects on the exudative pathophysiology, granulocyte activation, and epithelial and subepithelial remodelling in patients with airway diseases.

**Conclusions**

The epithelial injury-repair processes discussed above suggest that the airway mucosa is well equipped for quick and efficient restitution, irrespective of the severity of epithelial shedding. Thus, in health and disease epithelial desquamation may occur without seriously impairing the mucosal barrier. Epithelial damage-restitution processes involve in vivo specific tissue responses which, besides promoting defence and repair, may contribute to several facets of the airway pathology observed in patients with inflammatory airway diseases. The prevention of epithelial damage may therefore become an important strategy in the development of new treatment options for allergic and inflammatory airway diseases.

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