Effects of topical capsaicin in seasonal allergic rhinitis

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Abstract

Background – Mucosal exudation (luminal entry) of bulk plasma is a key feature of airway defence and inflammation. In guinea pig and rat airways this response is readily produced by neurogenic irritants, notably capsaicin. Thus “neurogenic airway inflammation” has become an established concept. The present study examines whether capsaicin also produces mucosal exudation of plasma in human nasal airways both in health and disease (seasonal allergic rhinitis).

Methods – Pain-producing concentrations of capsaicin (30–300 ng/ml) were applied to the nasal mucosal surface both before and late into the pollen season. Levels of albumin in nasal lavage fluid were measured as an index of mucosal exudation of plasma. In a separate group of patients with seasonal allergic rhinitis nasal challenge with an exudative concentration of histamine was carried out before the birch pollen season and concentrations of albumin in lavage fluid were measured.

Results – Pollen counts and symptom scores revealed a mild pollen season. Capsaicin produced considerable nasal pain and this response was augmented late into the season when capsaicin also produced nasal blockage. However, capsaicin failed to produce any mucosal exudation of plasma either before or late into the pollen season. The exudative effect of histamine was confirmed.

Conclusions – The augmented pain response to capsaicin suggests that a sensory nerve hyperresponsiveness may characterise allergic airways disease. In contrast to the effects on animal airways, capsaicin failed to produce mucosal exudation of plasma in the human nasal airway. The animal based neurogenic inflammation concept is therefore not valid for the human nasal airway, not even in inflamed airways when a neural hyperresponsiveness has developed.

(Keywords: allergy, rhinitis, neurogenic inflammation, capsaicin.)

Luminal entry or mucosal exudation of bulk plasma is a key feature of airway inflammation and plasma exudation has been seen in allergic and infectious airway diseases. Nasal and bronchial airways appear to respond equally to exudative challenges. The epithelial lining, even when intact, allows rapid entry of plasma into the airway lumen through valve-like para-cellular pathways. Plasma proteins in airway mucosal surface liquids may thus directly reflect the intensity of the subepithelial inflammation.

In rat and guinea pig airways mucosal exudation of plasma is readily induced by irritants such as capsaicin and nicotine. It has been shown that irritants release neuronal tachykinins which increase airway microvascular permeability in these animals. Kröll et al have further shown that the release in guinea pig airways and lung may be through a local axon reflex.

Based largely on these animal observations it has been suggested that neurogenic inflammation may be a feature of human airway disease. Capsaicin has been shown to produce pain and atropine-dependent rhinorrhea in the human nasal airway, and its major effect in the lower airways appears to be cough. No observations have been made so far to indicate that irritants can produce airway inflammation and mucosal exudation of plasma in human airways.

The human nose may be used for studies of inflammation that are also relevant to the lower airways. In a study in healthy subjects we have shown that nasal challenge with nicotine produces intense nasal pain and dose-dependent secretion of mucin, but fails to produce any mucosal exudation of plasma. Similarly, in healthy subjects Bascom et al found no mucosal exudation of albumin in nasal airways challenged with capsaicin. Ignoring the fact that luminal entry of plasma reflects the permeability of the subepithelial microcirculation, these authors did not emphasise that their observations could question the relevance of neurogenic inflammation in humans. Neither capsaicin nor nicotine has been evaluated in human airways when inflammation is already present.

In the present study we have examined
whether capsaicin, given topically in doses that can be tolerated acutely, induces mucosal exudation of plasma in patients with seasonal allergic rhinitis. Capsaicin challenges were performed before and late into the birch pollen season and concentrations of albumin in the lavage fluid and any induced nasal symptoms including pain were determined. To compare the effects of capsaicin with those of an exudative inflammatory mediator a histamine challenge test was performed before the birch pollen season in a separate group of patients with seasonal allergic rhinitis using an exudative concentration of histamine in the middle part of the concentration-response curve.15

Methods

Subjects

Eleven patients (eight men) with seasonal allergic rhinitis aged 22–28 years received capsaicin challenge before and late into the birch pollen season. A separate group of 10 patients (nine men) with allergic rhinitis aged 19–47 years received histamine challenge outside the birch pollen season. All patients had a history of strictly seasonal allergic rhinitis and a positive skin prick test to birch pollen allergen. No drugs were allowed during and for one month before the study. The study was approved by the local ethics committee and informed consent was obtained.

Pollen Count and Symptom Score

The daily birch pollen count was registered using a pollen trap located in the study region. The pollen count was expressed as mean daily number of pollen grains/m³ of air. Overall nasal symptoms – that is, sneezes, blockage, and rhinorrhea – were scored by the patients once daily using a symptom score where 0 = no symptoms, 1 = mild, 2 = moderate, and 3 = severe. After capsaicin challenge nasal blockage and nasal pain were scored as 0 = none, 1 = mild, 2 = moderate, and 3 = severe, and sneezes as 0 = 0 sneezes, 1 = 1–4 sneezes, 2 = 5–9 sneezes, and 3 = ≥ 10 sneezes. No attempt was made to estimate secretion after capsaicin challenge because of difficulties in distinguishing between lavage fluids and actual rhinorrhea. Capsaicin challenges were carried out before the birch pollen season (early in March) and late into the pollen season (study day 30, see fig 1). Study day 1 was 15 April 1992.

Nasal Challenges and Lavages

Isotonic saline and capsaicin (30 and 300 ng/ml) dissolved in isotonic saline were introduced in sequence into the nasal cavity using the nasal pool technique. The fluid was maintained in the nasal cavity for 10 minutes and two minutes elapsed between each instillation. Before each 10 minute instillation the mucosal surface was irrigated by two 30 second saline lavages to remove albumin that might have accumulated on the mucosal surface. Furthermore, to prevent capsaicin from being retained in the nasal airway the mucosal surface was irrigated by a 30 second saline lavage using the nasal pool technique, immediately after each 10 minute challenge. (These brief lavages were not collected.) In the separate group of patients with allergic rhinitis isotonic saline and histamine (400 μg/ml) in isotonic saline were each introduced into the nasal cavity for 10 minutes using the nasal pool technique. Two minutes elapsed between the instillations and the mucosal surface was irrigated by two 30 second saline lavages before each 10 minute instillation. The recovered fluids were centrifuged (105 g, 10 minutes, 4°C) and samples were obtained from the supernatant and frozen (−20°C) awaiting analysis.

Albumin Analysis

The concentrations of albumin in the lavage fluid were measured using a radioimmunoassay sensitive to 6.25 ng/ml. Rabbit anti-human albumin (Dakopatts, Copenhagen, Denmark) and standard (Calbiochem, San Diego, California, USA) were used. Iodination was performed by the lactoperoxidase method to a specific activity of 2.0 mCi/nmol.16 Tracer and standard or sample were mixed with antiserum before adding goat anti-rabbit antiserum (Astra Draco, Lund, Sweden). The bound fraction was measured by a gamma camera (Pharmacia, Uppsala, Sweden). The intraassay and interassay coefficients of variation were 5% and 10%, respectively.

Data Analysis

The Friedmann test and Wilcoxon signed rank test were used to examine differences in pollen counts and overall nasal symptoms during the pollen season, and to examine differences in symptoms and lavage fluid levels of albumin in capsaicin challenged subjects before and late into the pollen season, respectively. The Wilcoxon signed rank test was used to examine differences in lavage fluid levels of albumin between patients challenged with isotonic saline and histamine before the pollen season. p values <0.05 were considered significant. Data are presented as means (SE).
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Results
The pollen counts and symptom scores registered during the birch pollen season revealed a mild but significant pollen season (fig 1). The nasal symptom scores were significantly increased on study days 15 and 19–35 (p<0.05–0.001) compared with study day 1 (fig 1B).

Capsaicin challenges produced dose-dependent nasal pain (burning sensation) both before and late into the birch pollen season (Friedman test, p<0.01) compared with the saline control. Late into the birch pollen season the pain response to capsaicin 300 ng/ml was significantly greater than before the season (Wilcoxon signed rank test, p<0.05). Capsaicin failed to produce significant sneezes either before or late into the pollen season, but produced significant nasal blockage late into the season (Friedman test, p<0.01; Wilcoxon signed rank test, p<0.05, comparison between control saline and capsaicin 300 ng/ml) (fig 2).

Capsaicin challenges failed to produce mucosal exudation of albumin either before or late into the birch pollen season (fig 3). Indeed, repeated administrations of increasing concentrations of capsaicin produced successively smaller levels of albumin in the lavage fluid.

This effect of the repeated lavages was significant late into the season (Friedman test, p<0.01; Wilcoxon signed rank test, p<0.05 isotonic saline versus capsaicin 30 ng/ml, p<0.01 isotonic saline versus capsaicin 300 ng/ml). In contrast, the histamine challenge carried out in the separate group of patients with seasonal allergic rhinitis produced considerable mucosal exudation of albumin (p<0.01) (fig 3).

Discussion
In this study we have examined patients with seasonal allergic rhinitis both before and late into a mild but significant birch pollen season. We have previously shown increased levels of plasma proteins in nasal lavage fluid in mild seasonal allergic rhinitis.4 Furthermore, we have shown that the airway micro-circulation is hyperresponsive to exudative mediator challenge late into the pollen season.17 Hence, the presence of allergic disease may offer particular opportunities to detect plasma exudation responses. Repeated lavages with isotonic saline were carried out before the cap-

![Graph](http://thorax.bmj.com/)

Figure 1. (A) Daily pollen counts and (B) nasal symptom scores revealed a mild but significant birch pollen season. Capsaicin challenges were performed before the start of the season (early in March) and late into the season (study day 30). Study day 1 is 15 April 1992.

![Graph](http://thorax.bmj.com/)

Figure 2. (A) Capsaicin produced nasal pain (burning sensation) both before (open columns) and late into (filled columns) the birch pollen season. This effect was increased late in the season (*p<0.05). (B) Capsaicin produced significant nasal blockage late into (filled columns) but not before (open columns) the birch pollen season. There were no significant differences between the capsaicin-induced nasal blockage before and late into the season.
Figure 3 Eff ects of capsaicin on lavage fluid levels of albumin before (open columns) and late into (filled columns) the birch pollen season, and effects of histamine in a separate group of patients out of the season. Capsaicin failed to produce mucosal exudation of albumin both before and late into the birch pollen season. In contrast, the effect of histamine on plasma exudation was confirmed (**p<0.01).

Capsaicin challenge series to remove plasma that may have accumulated on the mucosal surface, thus further increasing the possibility of detecting any mucosal exudation of plasma.

Our results show that capsaicin, in a concentration that produces intense nasal pain, does not produce mucosal exudation of plasma. This negative outcome is equally clear both before and late into the birch pollen season. The capsaicin-induced pain was significantly greater during the season than before the start of the season. Hence, both in healthy conditions and in inflammation, in the presence of a sensory nerve hyperresponsiveness, capsaicin completely failed to produce any inflammatory exudation of plasma in the human airways.

These data add significantly to the previous findings with nicotine \(^1\) and capsaicin. \(^4\) Taken together, the findings in the human airway suggest that the animal-based concept of neurogenic inflammation may have no bearing on the human nasal airway.

The airways respond to inflammatory challenges such as histamine and allergen with extravasation of plasma from post-capillary venules of the subepithelial microrcirculation. \(^1\) Non-sieved bulk plasma is thus extravasated into the lamina propria along a hydrostatic pressure gradient. The extravasated plasma moves up between the epithelial cells and may exert a hydrostatic pressure load on the basolateral aspects of the epithelial cells from beneath. \(^1\) In vitro observations suggest that the pressure load produces a transient separation of the right junctions between the epithelial cells so that paracellular pathways are readily created for the clearance of extravasated plasma into the airway lumen. \(^17\) \(^18\) We have shown that the plasma extravasation process of the airways can be monitored by analysing the concentrations of plasma proteins in airway mucosal surface liquids. This luminal index – that is, mucosal exudation of plasma – may directly reflect the intensity of the airway inflammation. \(^1\) The luminal entry of bulk plasma extends to threshold inflammatory challenges. \(^1\)

This is an important consideration in the present study where surface liquids have been sampled. It is unlikely that capsaicin could have produced extravasation of plasma into the lamina propria without incurring a detectable increase in lavage fluid levels of plasma proteins. The mechanism of the epithelial passage of extravasated plasma thus strengthens the present conclusion that capsaicin may not produce plasma exudation in human airways.

Challenges with histamine type mediators or allergen readily produce 100-fold increases in exudative indices on the surface of the human nasal mucosa. In guinea pig airways the tachykinins and capsaicin are approximately equally effective to these challenges to produce plasma exudation. \(^2\) \(^3\) Large topical doses of substance P on the human nasal mucosa may produce significant systemic cardiovascular actions but only marginal or small secretory effects. \(^2\) \(^3\) By far the largest effects have been found by Braunstein et al who reported that albumin and total protein increased 4–10 times in nasal lavage fluids obtained after large topical doses of either substance P or neurokinin A had been given. \(^3\) It is not known to what degree these data represent secretion of albumin. Further studies involving measurement of specific indices such as α,α-macroglobulin are needed to establish whether or not exogenous tachykinins may produce any plasma exudation responses in human airways. Indeed, so far only guinea pigs and rats have been found to exhibit neurogenic exudative inflammation in the airways. Considering the lack of supportive data from other species including rabbits, cats, dogs, pigs, and monkeys, it is not surprising that the human airways also lack this mechanism, most clearly shown by the absence of capsaicin-induced exudative actions. Whether the airways of humans and other species contain nerves that might release tachykinins is another matter for investigation.

It is difficult to examine mechanisms of the plasma exudation response in the human bronchial airways with great accuracy. The concentration and distribution of challenge agents and lavage fluids on the mucosal surface may not be well controlled, and it may be difficult to distinguish between lavages of the airway and the alveolar surface. It is fortunate, therefore, that the upper and lower airways may exhibit similar responses to inflammatory challenges, particularly plasma exudation responses. \(^1\) We suggest that the present findings in the nasal airways are also valid for the bronchial airways.

Non-specific airway hyperresponsiveness to topical challenges is a feature of rhinitis and asthma. In the present study we have shown that capsaicin-induced pain was significantly greater late in the season than outside the season. This suggests that sustained airway inflammation may be associated with an increased responsiveness of sensory nerves. We have previously shown a microvascular hyperresponsiveness in allergic rhinitis in which histamine produced greater plasma exudation responses late in the season than outside the
Effects are which plasma in human airways, either that human inflammatory conditions in way this study may not be explained by increased mucosal penetration of capsaicin because mucosal absorption in patients with ongoing seasonal allergic rhinitis may actually be reduced.

The previous animal data have clearly shown that capsaicin produces inflammatory exudation in healthy airways. We have now shown that this particular response is not evoked in human airways, either in health or in disease. There are, of course, many inflammatory airway conditions in addition to seasonal allergic rhinitis in which the effect of capsaicin could be evaluated. The present observations do not therefore exclude the possibility that neurogenic inflammatory exudation may be evoked in human airways.

The results of this study indicate that capsaicin does not produce mucosal exudation of plasma in the human nasal airway of patients with allergic rhinitis, neither before nor late into a mild birch pollen season. We suggest that the neurogenic inflammatory responses which are readily produced by capsaicin in guinea pig and rat airways have no relevance to human airways.

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