Infection with Rhinovirus Facilitates Allergen Penetration Across a Respiratory Epithelial Cell Layer

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Abstract
Background: Rhinovirus infections are a major risk factor for asthma exacerbations. We sought to investigate in an in vitro system whether infection with human rhinovirus reduces the integrity and barrier function of a respiratory epithelial cell layer and thus may influence allergen penetration. Methods: We cultured the human bronchial epithelial cell line 16HBE14o- in a transwell culture system as a surrogate of respiratory epithelium. The cell monolayer was infected with human rhinovirus 14 at 2 different doses. The extent and effects of transepithelial allergen penetration were assessed using transepithelial resistance measurements and a panel of 125I-labeled purified recombinant respiratory allergens (rBet v 1, rBet v 2, and rPhl p 5). Results: Infection of respiratory cell monolayers with human rhinovirus decreased transepithelial resistance and induced a pronounced increase in allergen penetration. Conclusions: Our results indicate that infection with rhinovirus damages the respiratory epithelial barrier and allows allergens to penetrate more efficiently into the subepithelial tissues where they may cause increased allergic inflammation.

Introduction
In recent years, a clear and significant association between upper respiratory tract infections with rhinovirus and an increased frequency of asthma exacerbations in children has been shown [1, 2]. In adults, asthma exacerbations are also associated to a substantial degree with upper respiratory tract viral infections [3]. Furthermore, asthma patients with allergic sensitization to aeroallergens have an increased risk of hospital admission for exacerbation of their disease in the presence of rhinovirus in patients’ airways [4]. However, possible synergistic interactions of allergic sensitization and infection with rhinovirus have not been studied in detail. Allergen exposure via the respiratory mucosa induces strong rises of allergen-specific IgE levels, which in turn lead to increased allergen sensitivity in the target organs of allergy.
A growing body of evidence supports the view that epithelial defects at mucosal surfaces facilitate allergen contact with underlying immune competent cells, thus contributing to the initiation or exacerbation of allergic disease and asthma [6]. Infection with rhinovirus has been shown to affect airway barrier function [7], and allergic sensitization in children has been positively linked to rhinovirus- but not other virus-associated wheezing [8]. Furthermore, high titers of IgE antibodies to house dust mite allergens have been found to be common and increase the risk of acute wheezing provoked in asthmatic children by rhinovirus [9].

In this study we aimed to investigate whether rhinovirus infection impairs airway epithelial barrier function for defined recombinant allergen molecules with a view to allergic sensitization or exacerbation of allergic disease.

**Methods**

**Culture of Human Bronchial Epithelial Cell Layers**

We cultured the epithelial cell line 16HBE14o–, which is derived from human bronchial epithelial cells [10, 11]. This cell line retains properties of differentiated airway epithelial cells as it grows in a polarized monolayer forming tight junctions, apical cil ia, and microvilli and exhibits regulated ion transport. It can therefore be used as a surrogate of human respiratory epithelium. Cells were cultured in minimal essential medium containing 10% fetal calf serum, 100 U/ml penicillin, 100 μg/ml streptavidin, and 2 mM glutamine. Cells were passaged at 80% confluence. For transwell experiments, 2 × 10⁵ cells were seeded in the upper chamber of the transwell system and TER was measured 11 and 30 h after the infection. Experiments were conducted 2 times in duplicate and 1 time in triplicate wells. Data are presented as the means and SD of these experiments.

**Rhinovirus Infection**

Human rhinovirus 14 (HRV14) was obtained from ATCC (Manassas, Va., USA) and used for infection after growth in suspension cultures of HeLa cells (Ohio strain; Flow Laboratories, McLean, Va., USA) and used for infection after growth in suspension cultures of HeLa cells (Ohio strain; Flow Laboratories, McLean, Va., USA) for 40 h, preparation by polyethylene glycol precipitation, and resuspension in PBS as described previously [12]. Apical infection of epithelial monolayers with HRV14 was performed on confluent 16HBE14o– cells. Cell confluence was ascertained by phase contrast microscopy (×4 objective, Olympus IX73; Tokio, Japan) and images were recorded (Olympus DP73; Tokio, Japan). Confluent cell monolayers were apically infected in minimal essential medium containing 2% fetal calf serum, 100 U/ml penicillin, 100 μg/ml streptavidin, and 2 mM glutamine (all from Gibco Life Technologies) at 34°C. One hour after infection the uninfected virus was removed by washing with minimal essential medium containing 10% fetal calf serum, 100 U/ml penicillin, 100 μg/ml streptavidin, and 2 mM glutamine and incubation was continued at 37°C. In control wells, mock infection was performed by adding PBS.

**Assessment of Cytopathogenicity**

Cells were inspected visually by phase contrast microscopy (4× objective, Olympus IX73; Tokio, Japan) and images were recorded (Olympus DP73; Tokio, Japan). Cytopathogenicity was assessed with a crystal violet assay. 16HBE14o– cells were seeded in 96-well microtiter plates at 2 × 10⁵ cells/well and incubated for 11 h, 30 h, and 5 days with 100× or 1,000× TCID50/cell of HRV14 according to the above-described infection protocol, or mock infection was performed. Cells were stained with 150 μl/well 0.1% crystal violet in acetic acid/dH₂O for 10 min at 21°C. The crystal violet solution was then removed and wells were gently washed with dH₂O. Microtiter plates were inspected visually and photographed. Viable cells are stained blue by crystal violet.

**Measurement of Transepithelial Resistance under the Influence of Different Concentrations of Rhinovirus**

Epithelial barrier function was assessed by measuring TER using an ohm voltmeter (Millipore, Bedford, Mass., USA), as described previously [13]. Baseline resistance values across polyester membranes without cellular monolayers were 100 Ω·cm² on average and were subtracted from TER measurements. Experiments investigating the effect of rhinovirus infection on epithelial barrier function were initiated when the TER reached at least 300 Ω·cm². HRV14 (100× and 1,000× TCID50/cell) was added to the upper chamber of the transwell system and TER was measured 11 and 30 h after the infection. Experiments were conducted 2 times in duplicate and 1 time in triplicate wells. Data are presented as the means and SD of these experiments.

**Incubation of an Epithelial Cell Layer with Allergen before and during Infection with Rhinovirus**

Recombinant (r) Bet v 1 (17 kDa), the major birch pollen allergen, and rPhl p 5 (28 kDa), a major Timothy grass pollen allergen, were obtained from Biomay (Vienna, Austria). A dose of 0.2 μg/ml allergen was added to apical chambers of the transwell system when cells reached TER values of 200 Ω-cm² and cell layers were incubated for 2 h at 37°C. HRV14 (100× and 1,000× TCID50/cell) was used to infect 16HBE14o– cell monolayers and TER was measured 11 and 30 h after the infection. In a variation of the experiment, amounts of allergen similar to those described above were added 1 h after infection of the cells with rhinovirus.

**Quantification of Transepithelial Allergen Migration**

rBet v 1 (17 kDa), rBet v 2 (14 kDa), and rPhl p 5 (28 kDa) (all from Biomay) were 125I-labeled using the chloramin T method [14]. 125I-labeled allergens (10⁶ counts/min (cpm)/well) were added to the upper chamber of the transwell system 1 h after infection of cells with HRV14 (100× and 1,000× TCID50/cell), or mock infection was performed. Three hours thereafter, equivalent aliquots (50 and 150 μl, respectively) from the upper and the lower chamber were analyzed in a gamma counter (Wallac, Turku, Finland). Cells were removed from the membrane and lysed for determination of intracellular radioactivity. Counts per minute were determined for both chambers and the cell layer. All experiments and measurements were repeated at different time points.
SDS-PAGE and Autoradiography

Equivalent aliquots from the upper and lower chambers of the transwell system containing 125I-labeled allergens were separated by electrophoresis on a 12.5% sodium dodecyl sulfate-polyacrylamide gel, which also contained a molecular weight marker. Gels were sealed and directly autoradiographed using Kodak X-Omat S films and intensifying screens.

Statistics

TER measurements after 30 h of HRV14 infection were analyzed for comparison of the two HRV14 concentrations with mock infection by one-way analysis of variance (ANOVA) and subsequent t tests with a Bonferroni correction for post hoc multiple testing (Excel 2010; Microsoft). p < 0.05 was considered statistically significant for ANOVA testing. For the subsequent 2 post hoc t tests (mock infection vs. 100× TCID50 and mock infection vs. 1,000× TCID50), p < 0.025 was considered statistically significant for corrected testing.

Radioactive penetration experiments were considered as results from an explorative study. Mean values were therefore analyzed using t tests to compare the mock infection control with each HRV14 concentration. p < 0.05 was considered statistically significant.

Results

Exposure of an Epithelial Cell Monolayer to Rhinovirus Induces a Decrease in TER

We investigated whether apical infection of confluent 16HBE14o– cell monolayers with 2 different concentrations of HRV14 (100× and 1,000× TCID50/cell) could affect TER. The development of TER before and after infection is shown in figure 1. The TER in mock infected wells increased from 374 ± 88 to 499 ± 117 Ω·cm² after 11 h and to 648 ± 138 Ω·cm² after 30 h. Infection with 1,000× TCID50/cell led to a marked drop in TER from 403 ± 63 to 269 ± 225 Ω·cm² after 11 h, which further intensified at 30 h after infection to 60 ± 48 Ω·cm² (30 h: p = 0.008). Changes in wells which were infected with 100× TCID50/cell were less pronounced, decreasing from 383 ± 80 to 325 ± 162 Ω·cm² at 30 h after the infection (30 h: p = 0.003).

Table 1. Lack of CPE during the first 30 h of HRV14 infection

<table>
<thead>
<tr>
<th>Time after infection (h)</th>
<th>11 h</th>
<th>30 h</th>
<th>5 days</th>
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<tr>
<td>Mock infection</td>
<td>+</td>
<td>+</td>
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<tr>
<td>100× TCID50</td>
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<td>1,000× TCID50</td>
<td>+</td>
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16HBE14o– cells were seeded at a density of 2 × 10⁵ cells/well in 96-well microtiter plates and infected with 2 concentrations of HRV14 (100× and 1,000× TCID50/cell) or by mock infection and incubated for 11 h, 30 h, and 5 days. The respective CPE observed are indicated. + = Viable cells; – = complete CPE.

No Cytopathogenic Events Were Observed 11 and 30 h after Infection

When cells were monitored visually during HRV14 infection with 100× and 1,000× TCID50/cell, no cytopathogenic events (CPE) were seen 11 and 30 h after infection. In crystal violet assays no CPE were seen at any time point in uninfected cells or cells infected with 100× TCID50/cell. At 1,000× TCID50/cell of HRV14 no CPE were seen 11 and 30 h after infection; however, 5 days after infection complete CPE were seen in infected wells (table 1).

Incubation with Allergen before and during Infection of an Epithelial Cell Monolayer with Rhinovirus Does Not Further Decrease TER

16HBE14o– monolayers cultured on a semipermeable membrane in a transwell system were incubated apically for 2 h with 0.2 μg/ml rBet v1 or rPhl p 5, or no incubation with allergen was performed. The allergen amount (0.2 μg/ml) was chosen because it corresponds to the amount of allergen a patient is exposed to during a day of high pollen season [15, 16]. Cell monolayers were then...
infected apically with 100× or 1,000× TCID50/cell of HRV14, or mock infection was performed.

Thirty hours after infection, only minimal changes in TER were observed in cells which had been preincubated with allergens compared to cells preincubated with medium (preincubation with rBet v 1: +0.6% (HRV14; 100× TCID50) and +4% (HRV14; 1,000× TCID50); preincubation with rPhl p 5: –4% (HRV14; 100× TCID50) and –5% (HRV14; 1,000× TCID50)) Furthermore, allergen exposure without rhinovirus did not alter TER compared to the control medium (~2 to +4% difference in TER between allergen exposure and the control medium).

A variation of this experiment, where cell monolayers were infected first with 100× or 1,000× TCID50/cell of HRV14, with addition of rPhl p 5 or rBet v 1 one hour after infection yielded similar results of TER after 30 h of incubation, i.e. rPhl p 5: –9% (HRV14; 100× TCID50) and –15% (HRV14; 1,000× TCID50), and rBet v 1: +7% (HRV14; 100× TCID50) and +5% (HRV14; 1,000× TCID50).

Rhinovirus Exposure Facilitates Allergen Penetration across Epithelial Cell Layers

We measured the permeability of the 16HBE14o– cell layer for 3 different unrelated 125I-labeled allergens from different allergen sources (rBet v 1 and rBet v 2: birch pollen; rPhl p 5: grass pollen) after infection of cell monolayers with 100× and 1,000× TCID50/cell of HRV14. In figure 2 results are displayed as cpm lower chamber/cpm upper chamber in percent. The permeability for Phl p 5 started to increase after infection with 100× TCID50/cell of HRV14 (triangles in fig. 2) and was strongly augmented for all allergens at 1,000× TCID50/cell of HRV14. In figure 2 results are displayed as cpm lower chamber/cpm upper chamber in percent. The permeability for Phl p 5 started to increase after infection with 100× TCID50/cell of HRV14 (triangles in fig. 2) and was strongly augmented for all allergens at 1,000× TCID50/cell of HRV14 (circles in fig. 2) compared to the wells where mock infection had been performed (asterisks in fig. 2) (1,000× TCID50: p < 0.05). Less than 5% of radioactively labeled allergen was detected in the lower chambers, when the epithelial layer was intact, as observed previously [13]. Similar effects on allergen permeability were seen when the infection time and the penetration time were increased (data not shown). No radio-labeled allergens were detected within the cell layer, indicating paracellular penetration, as already observed previously [13].

Allergens Remain Intact during Transepithelial Penetration

To visualize the extent of allergen penetration and its impact on allergen integrity, aliquots of supernatants of the upper and lower chambers were removed and analyzed by SDS-PAGE. An autoradiography of a representative penetration experiment with rBet v 1 is displayed in figure 3. After infection with 1,000× TCID50/cell of HRV14 and a penetration time of 6 h, a considerable amount of rBet v 1 penetrated to the lower chamber of the transwell system, whereas almost no rBet v 1 was found in the lower chamber of a transwell system where a mock infection had been performed. rBet v 1 appeared at 17 kDa. Similar results were obtained with rBet v 2 and rPhl
These allergens are rapidly re-mediated such as nasal secretions increase the risk of acute wheezing induced by rhinovirus of house dust mite allergen-specific IgE are common and increased allergen penetration may lead to increased stimulation of mast cells, cells producing allergen-specific IgE, and T cells. In fact, it was recently reported that high levels of allergen and rhinovirus did not increase the observed effects in our experiments.

Our observation may explain why rhinovirus-induced asthma exacerbations in allergic individuals are more severe when the allergen is present at the time of viral infection [4]. It is well known that respiratory allergen contact strongly boosts the secondary IgE response to allergens and allergen-specific sensitivity [5]. Rhinovirus-induced increased allergen penetration may lead to increased stimulation of mast cells, cells producing allergen-specific IgE, and T cells. In fact, it was recently reported that high levels of house dust mite allergen-specific IgE are common and increase the risk of acute wheezing induced by rhinovirus in asthmatic children [9]. rBet v 1 and rPhl p 5, which were used in our experiments, are equivalent to their natural counterparts in terms of biochemical and immunological characteristics [17, 18]. These allergens are rapidly released when pollen grains come into contact with isotonic media such as nasal secretions [19, 20]. One can therefore assume that they are indeed present at the epithelial barrier and that the recombinant proteins correspond to the natural counterparts. Moreover, by using pure recombinant allergens without intrinsic protease activity, we made sure that increased allergen penetration could be imputed neither to extraneous protease activity from other components, which might be present in allergen extracts, nor to the synergistic invasive effect of bacteria, e.g. *Hemophilus influenzae*, which has been described to be greatly increased by rhinovirus infection [7].

It emerges that infection with rhinovirus may be an important risk factor in allergic asthma patients and needs to be prevented. Other viruses such as respiratory syncytial virus and influenza virus were not studied in our model, although they are also known to cause asthma exacerbations in children and adults [21]. It would thus be interesting to conduct further studies in our model with these viruses.

In summary, we have shown that infection with rhinovirus impairs the barrier function of a respiratory epithelial cell layer and facilitates transepithelial penetration of allergens, thereby potentially increasing submucosal allergen concentrations and aggravating allergic disease.

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**References**


