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Survival of rhinoviruses on human fingers

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Abstract

Rhinovirus is the main cause of the common cold, which remains the most frequent infection worldwide among humans. Knowledge and understanding of the rhinovirus transmission route is important to reduce morbidity as only preventive measures are effective. In this study, we investigated the potential of rhinovirus to survive on fingers. Rhinovirus-B14 was deposited on fingers for 30, 60, 90 and 120 min. Survival was defined as the ability of the virus to grow after 7 days, confirmed by immunofluorescence. Rhinovirus survival was not dependent on incubation time on fingers. Droplet disruption had no influence on survival. Survival was frequent with high rhinovirus concentrations, but rare with low-concentration droplets, which corresponded to the usual rhinovirus concentrations in mucus observed in children and adults, respectively. Our study confirms that rhinovirus infectiousness is related to the viral concentration in droplets and suggests that children represent the main transmission source, which occurs only rarely via adults. It confirms also that rhinovirus hand-related transmission is possible and supports hand hygiene as a key prevention measure.

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Introduction

Rhinoviruses are non-enveloped, positive-stranded RNA viruses belonging to the Enterovirus genus within the Picornaviridae family and the main causative agent of the common cold [1], the most frequent infection worldwide. Although usually a self-limited viral disease, it remains a source of significant morbidity in the community. Rhinovirus is associated also with asthma/wheezing and chronic obstructive pulmonary disease exacerbations, as well as several complications, such as acute otitis media, sinusitis, bronchitis and, in some cases, lower respiratory tract diseases. Pre-school children seem to be the main reservoir [2], as approximately six rhinovirus infections are observed per year and per child [3]. There are more than 150 different rhinovirus types with almost no cross-protection, which explains the frequency of rhinovirus infections and the absence of an effective vaccine or antiviral treatment. Only preventive measures are currently effective against these highly prevalent viruses and understanding their mode of transmission is important to reduce the number of infected patients.

The nasal mucosa and posterior nasopharynx have been documented as the main sites of viral replication and therefore the main shedding site [4,5]. It is reported that person-to-person transmission is most likely due to the contamination of hands by the nasal secretions of the infected person passed to a susceptible individual, either directly to the fingers or via an environmental intermediary; infection then follows from self-inoculation to the upper nasal airways or eyes [6–9]. The required infecting virus dose is below one median tissue culture infectious dose/mL (TCID50) [8,10]. Three possible transmission routes have been described: via aerosols of respiratory droplets, direct contact by hands, or indirect contact with environmental objects (fomites).
Aerosols produced by coughing or sneezing originate mainly from saliva [11] in which the viral load is approximately 30 times lower than in nasal secretions [4,8]. As rhinovirus transmission depends on the concentration of virus in secretions [4], this supports expert opinion that aerosol or oral transmission is a rare event [12–14]. Direct contact appears to play a major role in transmission. Rhinoviruses have been shown to transiently survive on human skin [4,6,13,15], leading to the hypothesis that hand-related transmission is the main transmission mechanism [6,13,16]. Although less frequently than on skin [13,17,18], rhinovirus has been shown to survive on fomites. In an experimental study, 50% of volunteers who touched their nasal mucosa or conjunctiva after handling a contaminated fomite developed infection [15]. However, many authors consider that indirect transmission is unlikely because of the important loss of infectivity during the process [17,19,20]. Our study was designed to test rhinovirus stability on fingers under experimental conditions, which aimed to reproduce natural conditions as far as possible.

Materials and methods

We conducted a series of experiments to assess the duration of human rhinovirus infectiousness duration on fingers, as well as the impact of viral concentration on survival rates. Survival was defined as the ability of the virus to grow on HeLaOH cells after 7 days, confirmed by immunofluorescence. Experimental conditions aimed to reproduce natural conditions as far as possible.

Viral suspensions and cell lines

All experiments were performed using the RV-B14 strain and HeLaOH cells (kindly provided by F.H. Hayden, University of Virginia, Charlottesville, VA, USA) for viral culture. RV-B14 stock (1 × 10^8 TCID50/mL) was diluted with respiratory mucus to obtain three different concentrations: 1 × 10^5 TCID50/mL (high concentration (HC)); 1 × 10^4 TCID50/mL (average concentration (AC)); and 1 × 10^2 TCID50/mL (low concentration (LC)). Each HC and AC droplet contained 1.1 × 10^5 viral copies for AC, and <100 copies for LC). After 1 h of adsorption at 33°C, 1 mL of McCoy’s 5A medium (1 ×) with 2% serum (Gibco, New York, NY, USA) for 60 seconds. Then, 400 μL of this eluate was used to immediately inoculate HeLaOH cells. This represents an additional 2.5-fold dilution of the viral load present in droplets before inoculation onto cell cultures (4.4 × 10^4 viral copies for HC, 1.1 × 10^4 viral copies for AC, and <100 copies for LC). After 1 h of adsorption at 33°C, 1 mL of McCoy’s 5A medium (1 ×) with 2% serum (Gibco) was added and cells were incubated in 5% CO2 at 33°C for 7 days. For each 24-well plate, a negative control (mucus only). Each contaminated finger was untouched for a defined period of time at room temperature before testing for the presence of infectious rhinovirus. Participants’ fingers were then immersed in wells (Becton Dickinson and Co., Franklin Lakes, NJ, USA) containing 1 mL of McCoy’s 5A medium (1 ×) with 2% serum (Gibco, New York, NY, USA) for 60 seconds. Then, 400 μL of this eluate was used to immediately inoculate HeLaOH cells. This represents an additional 2.5-fold dilution of the viral load present in droplets before inoculation onto cell cultures (4.4 × 10^4 viral copies for HC, 1.1 × 10^4 viral copies for AC, and <100 copies for LC). After 1 h of adsorption at 33°C, 1 mL of McCoy’s 5A medium (1 ×) with 2% serum (Gibco) was added and cells were incubated in 5% CO2 at 33°C for 7 days. For each 24-well plate, a negative control as well as a mock-infected control finger was included. The cytopathic effect was read daily until day 7. Cells were collected after 7 days and submitted to an immunofluorescence assay.

Immunofluorescence

A J2 mouse monoclonal antibody [23] that recognizes double-stranded RNA and an anti-mouse monoclonal IgG fluorescein isothiocyanate-conjugated antibody were used to confirm the presence of viral infection (Chemicon-Millipore, Zug, Switzerland).

Results

Based on preliminary pilot experiments, we determined that rhinovirus survival on fingertips was equivalent across different respiratory viruses (influenza virus A/B, human metapneumovirus, coronavirus 229E/HKU1/OC43/NL63, respiratory syncytial virus A/B, picornavirus and parainfluenza virus 1/2/3). A further 20 min of ultraviolet radiation ensured inactivation of putative undetected viruses. To guarantee optimal growth, only mucus with a pH between 6.5 and 7 was retained.

Participants and finger contamination procedure

Six specialized laboratory collaborators (technicians and MD/PhD graduates) were recruited on a voluntary basis as previously described [22]. The protocol was approved by the institutional review board of the University Hospitals of Geneva.

Determination of infectiousness

A 2-μL drop of viral suspension of human RV-B14 mixed with respiratory secretions was deposited on the fingertips of each participant. This volume represents the mean size of a large respiratory droplet and can be easily reproduced [22]. For each subject, nine drops containing rhinovirus at different concentrations (three HC, three AC, three LC) were deposited and one negative control (mucus only). Each contaminated finger was kept untouched for a defined period of time at room temperature before testing for the presence of infectious rhinovirus. Participants’ fingers were then immersed in wells (Becton Dickinson and Co., Franklin Lakes, NJ, USA) containing 1 mL of McCoy’s 5A medium (1 ×) with 2% serum (Gibco, New York, NY, USA) for 60 seconds. Then, 400 μL of this eluate was used to immediately inoculate HeLaOH cells. This represents an additional 2.5-fold dilution of the viral load present in droplets before inoculation onto cell cultures (4.4 × 10^4 viral copies for HC, 1.1 × 10^4 viral copies for AC, and <100 copies for LC). After 1 h of adsorption at 33°C, 1 mL of McCoy’s 5A medium (1 ×) with 2% serum (Gibco) was added and cells were incubated in 5% CO2 at 33°C for 7 days. For each 24-well plate, a negative control as well as a mock-infected control finger was included. The cytopathic effect was read daily until day 7. Cells were collected after 7 days and submitted to an immunofluorescence assay.
incubation times as it remained infectious on all fingers after 30, 60, 90 and 120 min. Immediately after deposition, half of the droplets were disrupted and spread on the surface of the fingertip using a pipette tip to determine whether disrupting the integrity and environment of the droplet decreased virus survival. As all intact and disrupted droplets yielded positive culture results, we decided to continue experiments with disrupted droplets only so as to reproduce real-life conditions as much as possible.

One hour after the deposit of disrupted droplets on the fingers of the six volunteers, infectious viruses could be detected by culture in all subjects contaminated with HC droplets (6/6), in four of the six volunteers with AC droplets, and none of the six volunteers with LC droplets, which confirmed the influence of concentration on survival (Fig. 1). Of note, when droplets were directly incubated without a passage on fingers, the virus survived in 100% (4/4) of tested fingers at HC compared with 25% (1/4) at LC, despite being below the limit of detection by PCR (data not shown). Overall, the proportion of fingers with detectable viruses was 16/18 fingers at HC, compared with 6/18 and 0/18 for the AC and LC droplets, respectively (Fig. 1). Laboratory room (mean ± standard deviation 24.6 ± 0.7°C) and hood (26.4 ± 1.8°C) temperature, as well as humidity (44.5 ± 5.6%), were similar for all experiments with all subjects.

Discussion

We aimed to investigate rhinovirus transmission by person-to-person contact, the main transmission route for the most prevalent human respiratory infection worldwide. Experiments were designed to reproduce, as much as possible, conditions that could lead to rhinovirus contamination of fingertips in the community. Our study showed that rhinovirus can survive on hands for several hours, similar to previous reports of virus survival on human skin [4,6,13,15,17], emphasizing that hand-related transmission is the main transmission route. There was no influence of drying time on virus survival under 2 h, in contrast to the study of Ansari et al. where virus survival decreased during the first hour [24].

Our study showed that virus survival, and therefore infectiousness, was related to the viral concentration in droplets. This correlates well with D’Alessio et al. who found that the secondary attack rate was related to the viral concentration in the nose [4]. Inoculum seems to be a restrictive factor for transmission, with infectiousness rapidly dropping below a given concentration. As infected children appear to have a higher viral load than adults, this may explain why children are considered to be the main transmission vector. The fact that the viral load in LC droplets was below the level of detection explains why the virus could not be recovered at these concentrations, except in one case without a passage on fingers. LC droplets correspond to the viral concentration recovered in rhinovirus-infected adults and this suggests that transmission via adults occurs rarely. A recent study investigating the transmission of cold-like illnesses between siblings showed that younger children tended to become infected first in most cases. However, the secondary attack rate was greater for older siblings, probably because of a higher viral load in younger siblings’ secretions. As younger children tend also to touch nasal secretions directly with their fingers, it is probable
that this enhances transmission. The viral load in the mucus of more than 1000 rhinovirus-infected children below 1 year of age was $5.79 \times 10^{6} \text{TCID}_{50}/mL$, which is 10 to 100 times higher than our HC of $1 \times 10^{5}$ (Regamey et al., private communication). It is very probable that the difference in virus survival between adults and children is even higher than in our results.

We showed that virus survival increased at LC when there was no passage on hands. The loss of infectiousness during interhuman contact or fomite manipulation has already been described [20,24], highlighting again the importance of viral load for transmission. Similarly, rhinovirus was more frequently recovered on fingers from subjects with a high nasal viral load compared with a low nasal viral load [17].

Our study confirmed that droplet disruption had no influence on survival at a given concentration. We have previously shown that influenza virus survival on fingers was not related to virus concentration in a study using a similar methodology to the present experiments [22]. Survival of influenza virus on fingers declined rapidly, with less than 15% of the fingers remaining positive after 30 min [22]. The influenza envelope, which is known to be a determinant factor decreasing virus survival, may explain why survival was shorter and affected by droplet disruption, which was not the case for rhinovirus [22]. In a similar study, RV-14 in 10-μL droplets at $2.9 \times 10^{4}$ to $1.4 \times 10^{5} \text{TCID}_{50}/mL$ survived for 1 h on almost 40% of fingers [24]. The fact that virus survival was lower compared with our results, despite the use of bigger droplets and higher viral concentrations, may be explained by the fact that our volunteers did not wash their hands or use an alcohol-based hand rub before experiments as hand rubbing has been shown to decrease virus survival even several hours after use [25]. The fact that disinfecting hands or objects with an iodine or alcohol-based solution reduces the secondary illness rate of rhinovirus [22].

In conclusion, these laboratory results confirm that hand-related transmission of rhinovirus is possible and support hand hygiene as a key measure to prevent transmission, particularly in children who represent the main transmission source.

### Transparency declaration

The authors declare that they have no conflicts of interest.

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