Modes of Transmission of Influenza B Virus in Households

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Abstract

Introduction: While influenza A and B viruses can be transmitted via respiratory droplets, the importance of small droplet nuclei “aerosols” in transmission is controversial.

Methods and Findings: In Hong Kong and Bangkok, in 2008–11, subjects were recruited from outpatient clinics if they had recent onset of acute respiratory illness and none of their household contacts were ill. Following a positive rapid influenza diagnostic test result, subjects were randomly allocated to one of three household-based interventions: hand hygiene, hand hygiene plus face masks, and a control group. Index cases plus their household contacts were followed for 7–10 days to identify secondary infections by reverse transcription polymerase chain reaction (RT-PCR) testing of respiratory specimens. Index cases with RT-PCR-confirmed influenza B were included in the present analyses. We used a mathematical model to make inferences on the modes of transmission, facilitated by apparent differences in clinical presentation of secondary infections resulting from aerosol transmission. We estimated that approximately 37% and 26% of influenza B virus transmission was via the aerosol mode in households in Hong Kong and Bangkok, respectively. In the fitted model, influenza B virus infections were associated with a 56%–72% risk of fever plus cough if infected via aerosol route, and a 23%–31% risk of fever plus cough if infected via the other two modes of transmission.

Conclusions: Aerosol transmission may be an important mode of spread of influenza B virus. The point estimates of aerosol transmission were slightly lower for influenza B virus compared to previously published estimates for influenza A virus in both Hong Kong and Bangkok. Caution should be taken in interpreting these findings because of the multiple assumptions inherent in the model, including that there is limited biological evidence to date supporting a difference in the clinical features of influenza B virus infection by different modes.

Introduction

Influenza viruses are believed to be spread between humans through a number of modes of transmission, including primarily through inhalation of respiratory droplets containing infectious virus, and possible contact of respiratory secretions containing infectious virus with mucous membranes. A distinction is

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Data Availability: The authors confirm that all data underlying the findings are fully available without restriction. The Hong Kong dataset is freely available online. The Hong Kong study data are at http://web.hku.hk/~bcowling/influenza/HK_NPI_study.htm The Bangkok dataset is available upon request, and cannot be included in a public repository because it is owned by a third party. The data can be obtained from Dr Kim Lindblad (kl@cdc.gov).

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Competing Interests: BJC has received research funding from MedImmune Inc. and Sanofi Pasteur, and consults for Crucell NV. DKMI has received research funding from F. Hoffmann-La Roche Ltd. JSMP receives research funding from Crucell NV and serves as an ad hoc consultant for GlaxoSmithKline and Sanofi Pasteur. JMS has retired from the US CDC and now works with Sanofi Pasteur. BJC is a Section Editor for PLOS ONE. The other authors report no other potential conflicts of interest. This does not alter the authors’ adherence to all PLOS policies on sharing data and materials.

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Infectious influenza B virus can be detected in the aerosol fraction of infections, specifically that the mode of exposure leading to an infection may affect the pattern in subsequent signs and symptoms [21], and we use the same modeling framework to infer the proportion of household transmission of influenza B virus that occurs via the aerosol route.

### Methods

#### Sources of Data

During 2008–2011, large randomized controlled trials were conducted in Hong Kong and Bangkok to study the efficacy of hand hygiene and surgical face masks in reducing influenza virus transmission in households [22,23]. In each study, local residents who had acute respiratory illness and living in a household with at least 2 other people of whom none had reported acute respiratory illness in the preceding 14 days were enrolled. Pooled nasal and throat swab specimens were collected from each participant for testing with the QuickVue Influenza A+B rapid diagnostic test (Quidel, San Diego, California). Participants with a positive rapid influenza test result were further followed up along with their household contacts. Households were randomly allocated in equal proportions into one of three intervention groups: (1) a control intervention, (2) control plus hand hygiene intervention, and (3) control plus surgical face mask intervention.

### Table 1. Minor differences between the study designs in Hong Kong and Bangkok.

| Study component                          | Hong Kong                                                                 | Bangkok                                                                 |
|-----------------------------------------|---------------------------------------------------------------------------|-------------------------------------------------------------------------|
| Recruitment locations                   | 45 public and private outpatient clinics across Hong Kong (population 7 million). | Outpatient department of a large pediatric public hospital in Bangkok (population 8 million). |
| Study period                            | January 2008–June 2009                                                    | April 2008–February 2011                                                |
| Age of index cases                      | Any age                                                                   | Children 1 m to 15 y of age                                             |
| Eligibility of index case (symptoms)    | Presenting with at least two of: fever $\geq$ 37.8°C, cough, sore throat, headache, runny nose, phlegm, and myalgia; living with at least two other people. | For $<$ 2 years: fever $>$ 38°C and one or more of the following symptoms: nasal congestion, cough, conjunctivitis, respiratory distress, sore throat, new seizure. For $>$ 2 years: Presenting with influenza-like illness (fever plus cough or sore throat); living with at least two other people. |
| Exclusion criteria                      | Recent (within 14 d) acute respiratory illness in any household member    | Recent (within 7 d) influenza-like illness in any household member; recent (within 12 m) influenza vaccination in any household member. |
| Hand hygiene intervention               | Distribution of alcohol hand rub to each household member in addition to liquid hand soap to the household | Distribution of liquid hand soap to the household                      |
| Measurement of body temperature         | All households were provided and instructed in the use of a free tympanic thermometer and asked to record their body temperature daily. | Thermometers were not provided to households, and participants recorded either measured body temperature or ‘feverishness’. |

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control plus facemasks and hand hygiene interventions. A home visit was scheduled as soon as possible after randomization to implement the intervention, collect baseline demographic data and NTS specimens from all household contacts aged ≥2 years, and to describe the information to be recorded in daily symptom diaries. Further home visits were scheduled at 3 and 6 days after the first home visit to monitor adherence to intervention and to collect further NTS specimens from all household contacts regardless of illness. The two study protocols were very similar, and notable differences are summarized in Table 1.

All NTS specimens were tested by reverse-transcription polymerase chain reaction (RT-PCR) for influenza A and B virus infection.

**Table 2.** Characteristics of index cases with confirmed influenza B virus infection and their household contacts in Hong Kong, by intervention group.

| Characteristics                                      | Control | Hand hygiene | Face mask+hand hygiene |
|-------------------------------------------------------|---------|--------------|------------------------|
|                                                       | n (%)   | n (%)        | n (%)                  |
| Index cases                                           | 35 (100) | 36 (100)    | 33 (100)               |
| Age group                                             |         |              |                        |
| ≤5 y                                                  | 5 (14%) | 3 (8%)       | 4 (12%)                |
| 6–15 y                                                | 25 (71%)| 21 (58%)     | 21 (64%)               |
| >16 y                                                 | 5 (14%) | 12 (33%)     | 8 (24%)                |
| Male                                                  | 16 (46%)| 19 (53%)     | 10 (30%)               |
| Median household size (IQR)                          | 4 (3, 5)| 4 (3, 4)     | 4 (3, 5)               |
| Household contacts                                   | 112     | 101          | 106                    |
| Age group                                             |         |              |                        |
| ≤5 y                                                  | 6 (5%)  | 1 (1%)       | 5 (5%)                 |
| 6–15 y                                                | 13 (12%)| 12 (12%)     | 12 (11%)               |
| 16–30 y                                               | 21 (19%)| 17 (17%)     | 17 (16%)               |
| 31–50 y                                               | 58 (52%)| 48 (48%)     | 51 (48%)               |
| >50 y                                                 | 14 (12%)| 23 (23%)     | 21 (20%)               |
| Male                                                  | 39 (35%)| 40 (40%)     | 46 (43%)               |
| Received seasonal influenza vaccination in the previous 12 m | 15 (13%)| 12 (12%)     | 14 (13%)               |

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**Table 3.** Characteristics of index cases with confirmed influenza B virus infection and their household contacts in Bangkok, by intervention group.

| Characteristics                                      | Control | Hand hygiene | Face mask+hand hygiene |
|-------------------------------------------------------|---------|--------------|------------------------|
|                                                       | n (%)   | n (%)        | n (%)                  |
| Index cases                                           | 37 (100)| 38 (100)    | 38 (100)               |
| Age group                                             |         |              |                        |
| ≤5 y                                                  | 12 (32%)| 14 (37%)     | 10 (26%)               |
| 6–15 y                                                | 25 (68%)| 24 (63%)     | 28 (74%)               |
| >16 y                                                 | 0 (0%)  | 0 (0%)       | 0 (0%)                 |
| Male                                                  | 24 (65%)| 23 (61%)     | 23 (61%)               |
| Median household size (IQR)                          | 2 (2, 3)| 3 (2, 3)     | 3 (2, 5)               |
| Household contacts                                   | 84      | 91           | 89                     |
| Age group                                             |         |              |                        |
| ≤5 y                                                  | 1 (1%)  | 5 (5%)       | 4 (4%)                 |
| 6–15 y                                                | 10 (12%)| 14 (15%)     | 10 (11%)               |
| 16–30 y                                               | 13 (15%)| 18 (20%)     | 15 (17%)               |
| 31–50 y                                               | 49 (58%)| 41 (45%)     | 37 (42%)               |
| >50 y                                                 | 11 (13%)| 13 (14%)     | 23 (26%)               |
| Male                                                  | 35 (42%)| 39 (43%)     | 33 (37%)               |
| Received seasonal influenza vaccination in the previous 12 m | 0 (0%)  | 0 (0%)       | 0 (0%)                 |

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viruses using standard methods as described elsewhere [22,23]. In
the present analyses only the households of index cases with RT-
PCR-confirmed influenza B virus infection are included; results for
index cases with influenza A were reported elsewhere [9].

In the present analyses, we used data on influenza B virus
transmission in families from the studies in Hong Kong and
Bangkok. Specifically, we identified all index cases with confirmed
influenza B virus infection, and their household contacts. We then
determined which household contacts had RT-PCR confirmed
infection, the corresponding times of illness onset, and whether
fever and cough were reported. In the analyses we also used the
allocated intervention group for each household, and the age of
each household contact.

Ethics Statement

All subjects 18 years of age and older gave written informed
consent, and proxy written consent was obtained from parents or
legal guardians for children aged 17 years old or younger. The
protocols for the studies in Hong Kong and in Bangkok were
approved by Institutional Review Board of the University of Hong
Kong, and the Institutional Review Board of Queen Sirikit
Hospital Bangkok, respectively [22,23].

Statistical Analysis

We used the Nelson-Aalen non-parametric estimator of the
cumulative hazards of infection with or without febrile disease plus
cough in each intervention group [24]. We constructed a
competing risks survival analysis model that accounted for the
alternative modes of transmission and used it to infer the relative
importance of alternative modes of transmission assuming that the
risk of fever plus cough higher in aerosol transmission, compared
with the other two modes. We assumed independent hazards over
time of influenza transmission in households with one or more
secondary cases. The cause-specific probability of aerosol trans-

Figure 1. Cumulative hazards of RT-PCR-confirmed influenza B virus infections presenting with fever plus cough or not presenting
with fever plus cough, among the household contacts in 104 and 113 households of index cases with RT-PCR-confirmed influenza
B virus infection in Hong Kong and Bangkok, respectively.
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Modes of Influenza B Virus Transmission
mission was estimated to measure the relative contribution of aerosol transmission among all three modes.

The mixture model was used to allow for a certain proportion (\( \theta \)) of subjects to be immune or not exposed, with the density of infection described as \( f(t) = (1 - \theta)f_{a}(t) \), where \( f_{a}(t) \) is the probability density function for the exposed and susceptible group.

The time to infection (\( T \)) for each of three modes of transmission was assumed to follow a Weibull distribution with an identical shared shape parameter (\( \lambda \)). The sub-hazards for modes of transmission, \( j = 1, 2, 3 \) for aerosol, contact, and droplet transmission respectively were written as follows:

\[
\begin{align*}
 h_{a1}(T, X_{hi}, X_{mi}) &= \phi \lambda_1 \lambda_2^{-1} T^{\lambda_2 - 1} \exp(\beta_1 X_{hi}), \\
 h_{a2}(T, X_{hi}, X_{mi}) &= \phi \lambda_2 T^{\lambda_2 - 1} \exp(\beta_2 X_{mi}), \\
 h_{a3}(T, X_{hi}, X_{mi}) &= \phi \lambda_3 T^{\lambda_3 - 1},
\end{align*}
\]

where \( X_{hi} / X_{mi} \) are the dichotomous indicator variables representing the allocation of hand hygiene/surgical mask interventions respectively to individual \( i \), and \( r_1 / r_2 \) represent the relative risk reductions in contact/large droplet transmission by hand hygiene/surgical masks respectively. We assumed that the risk of fever plus cough caused by infections follows a Bernoulli distribution with mean parameter \( \pi(j), j = 1, 2, 3 \) for three arms, respectively. We estimated \( \phi, \lambda_1, \lambda_2, \lambda_3, \pi_1, \pi_2, \pi_3, \beta_1, \beta_2, \beta_3 \). We were unable to estimate \( r_1 \) and \( r_2 \) so we examined the estimates of the other parameters for a range of values of \( r_1 \) and \( r_2 \). Further technical details of the model are provided in an earlier publication [9].

We performed statistical inference under a Bayesian framework, using Markov chain Monte Carlo (MCMC) to obtain parameter estimates from the posterior distributions [25]. We specified flat priors for each parameter. For each MCMC chain we ran 120,000 iterations, discarding the first 20,000 iterations as burn-in, and drawing every tenth subsequent value to compose the posterior distribution. All the statistical analyses were conducted in R version 2.15.1 (R Foundation for Statistical Computing, Vienna, Austria).

### Results

In Hong Kong and Bangkok there were 104 and 113 households, respectively, with an index case with RT-PCR-confirmed influenza B virus infection. The characteristics of index cases and their household contacts are shown in Tables 2 and 3 for Hong Kong and Bangkok respectively. We examined the cumulative hazard of RT-PCR-confirmed influenza B virus infections for household contacts, and found increases in the risk of infection with fever plus cough, and decreases in the risk of infection without fever plus cough, in the intervention arms compared to the control arm. The change was particularly apparent in the households in Bangkok (Figure 1). To be more specific, we found a statistically significant decrease in the risk of infection without fever plus cough, in the hand hygiene plus face mask arm compared to the control arm in the households in Bangkok.

Under the scenario where randomization to the hand hygiene intervention reduced contact transmission by 50% while randomization to face mask and hand hygiene interventions reduced both contact and droplet transmission by 50%, we fitted the transmission model to the Hong Kong and Bangkok data. We estimated that in the absence of interventions, aerosol transmission was responsible for 37% and 26% of secondary infections in Hong Kong and Bangkok, respectively (Table 4). We also varied the assumed efficacy of hand hygiene and face masks from 0% to 100% and estimated the relative importance of aerosol transmission in the absence of interventions, which ranged from approximately 20% to 80% in Hong Kong and 20% to 32% in Bangkok (Figure 2).

| Parameters | Hong Kong (104 households with 319 contacts) | Bangkok (113 households with 264 contacts) |
|------------|---------------------------------------------|-------------------------------------------|
| \( \beta \) Shape of the Weibull distribution | 2.16 (1.30, 3.12) | 0.77 (0.39, 1.28) |
| \( \lambda_1 \) Force of contact transmission* | 0.18 (0.01, 0.40) | 0.16 (0.01, 0.48) |
| \( \lambda_2 \) Force of droplet transmission* | 0.20 (0.01, 0.40) | 0.07 (0.00, 0.24) |
| \( \lambda_3 \) Force of aerosol transmission* | 0.22 (0.02, 0.38) | 0.08 (0.00, 0.25) |
| \( \gamma_1 \) Risk of fever plus cough for infections by contact route | 23% (1%, 66%) | 25% (1%, 63%) |
| \( \gamma_2 \) Risk of fever plus cough for infections by droplet route | 24% (1%, 60%) | 31% (2%, 75%) |
| \( \gamma_3 \) Risk of fever plus cough for infections by aerosol route | 56% (26%, 97%) | 72% (41%, 99%) |
| \( \theta_1 \) Proportion of household adults immune or not exposed | 90% (85%, 94%) | 65% (45%, 79%) |
| \( \theta_2 \) Proportion of household children immune or not exposed | 69% (54%, 82%) | 61% (34%, 82%) |

*The forces of infection in combination with a shared shape parameter determine the hazard associated with each competing mode of transmission. The relative contribution of each mode \( j \) is calculated as the cause-specific probabilities \( \frac{\gamma_j}{\gamma_1 + \gamma_2 + \gamma_3} \).
We compared the cause-specific probabilities of each mode of transmission as well as the associated illnesses in the control arm for influenza A and B virus infections, in Hong Kong and Bangkok respectively (Figure 3). Data for influenza A were extracted from a previous report [9]. Both influenza A and B virus infections attributed to aerosol transmission were associated with a higher risk of fever plus cough, compared with the other two modes of transmission. The point estimates of aerosol transmission were lower for influenza B compared to influenza A in both Hong Kong and Bangkok.

**Discussion**

We propose that the mode of spread associated with an influenza B virus infection affects the probability of experiencing fever plus cough for that infection. Based on that hypothesis, we estimated that approximately 37% and 26% of transmission was via the aerosol mode in households in Hong Kong and Bangkok, respectively. However, we should exercise caution in interpreting these findings because we have not been able to find literature supporting the anisotropic nature of influenza B virus infection, whereas we previously described literature supporting this

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**Figure 2.** The relative importance (cause-specific probability) of aerosol transmission in households in Hong Kong and Bangkok.

The contour lines show the proportion of secondary influenza B virus infections attributed to aerosol transmission in the control arm of each study, under varying assumptions about the efficacy of randomization to the hand hygiene and surgical mask interventions in reducing contact (x-axis) and droplet (y-axis) transmission respectively.

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**Figure 3.** The proportion of all influenza A and B virus infections attributed to each mode in the control arms of the studies in Hong Kong (blue) and Bangkok (brown), and the infections associated with fever plus cough (darker shade) or not associated with fever plus cough (lighter shade).

Data shown on influenza A were extracted from a previous study [9]. The contributions of the three modes sum to 100% within each geographic location and influenza type.

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property for influenza A virus infections [9]. Nevertheless, patterns in secondary infections and disease in the controlled trials in Hong Kong and Bangkok were consistent with this hypothesis (Figure 1). This also implicitly suggested that though hand hygiene and face masks could reduce the risk of transmission through contact or large droplets, but meanwhile increase the risk of aerosol transmission, which was associated with a greater risk of illness with fever plus cough.

Whereas we previously estimated that approximately half of within-household transmission of influenza A virus could be associated with aerosols [9], here we estimated a slightly reduced importance of aerosols for influenza B virus (Figure 3). One explanation for such a difference could be the age mix of cases of influenza A versus B, if aerosol transmission were more important among adults than children. We did not have sufficient sample size in the present study to examine whether modes of transmission might vary by age, but this would be an interesting area for further exploration.

If aerosol transmission is indeed an important mode of spread of influenza B virus, this may have important implications for control efforts. In particular measures targeting contact transmission, such as hand hygiene, and measures targeting large respiratory droplet transmission, such as surgical face masks, may not be sufficient to substantially reduce the risk of transmission. Control measures that might reduce aerosol transmission indoors include improvement in ventilation [26], modification of humidity [27], or the use of personal protective equipment that is more effective against aerosols than surgical masks. While the use of N95 respirators may not be practical in community settings and fit-testing is unlikely although required for optimal performance, other types of face masks with improved filtration compared to standard surgical masks or procedure masks may be available in the future.

There are a number of limitations to our analysis. First, our model did not include the possibility of variability in infectiousness between index cases, variability in immunity to different modes of transmission, or variability in within-household transmissibility associated with physical dimensions of the home, ventilation rates etc, and inclusion of these or other factors potentially affecting transmission dynamics could be natural extensions to our model. Because interventions were allocated randomly among households, the possibility of confounding should be minimized. Second, our model implicitly assumes that only the first infectious exposure is relevant to susceptible contacts, and once infected by that first exposure, further exposures are unimportant. Our model could be modified to allow for multiple simultaneous exposures by one or more modes, if it were understood how this might affect the course of disease. Third, while we assumed that all infections of household contacts during the 7-day follow-up were acquired within the household, it is possible that some infections were acquired outside. However in a separate study with a similar design in Hong Kong we used molecular epidemiology analyses of virus sequence data to demonstrate that many secondary influenza cases acquired infection from within the household [28], and a similar observation was reported in a household transmission study in Canada [29]. Fourth, it is possible that some secondary influenza virus infections were not confirmed due to poor quality specimens collected during home visits, or if peak influenza B viral shedding in the respiratory tract occurred between home visits at 3-day intervals. We did include serological data although this could have provided additional information on infections among household contacts. Fifth, by recruiting in outpatient clinics and using a rapid test to screen index cases, we may have introduced selection bias towards index cases with more serious illness or higher levels of virus shedding, affecting the relative importance of different modes of transmission. Finally, we did not explicitly account for imperfect adherence to the interventions, although the parameters in our model account for moderate efficacy of interventions against specific modes of transmission. Further improvements in the model might be obtained by incorporating limited data on adherence that was mainly self-reported by participants.

In conclusion, we propose that the aerosol route may be an important mode of transmission of influenza B virus in households. Further studies of non-pharmaceutical interventions in households would be improved by more careful monitoring of viral contamination on surfaces [30,31] and in the air, and inclusion of this information in transmission models.

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Author Contributions
Conceived and designed the experiments: BJC DKMI VJF HN. Analyzed the data: BJC VJF HN. Contributed to the writing of the manuscript: BJC DKMI VJF PS SJO JL TMU GMJ JSMP TC HC JMS. Interpreted data: PS SJO JL TMU GMJ JSMP TC JMS.

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