Detecting Human-to-Human Transmission of Avian Influenza A (H5N1)

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Highly pathogenic avian influenza A (HPAI) subtype H5N1 has caused family case clusters, mostly in Southeast Asia, that could be due to human-to-human transmission. Should this virus, or another zoonotic influenza virus, gain the ability of sustained human-to-human transmission, an influenza pandemic could result. We used statistical methods to test whether observed clusters of HPAI (H5N1) illnesses in families in northern Sumatra, Indonesia, and eastern Turkey were due to human-to-human transmission. Given that human-to-human transmission occurs, we estimate the infection secondary attack rates (SARs) and the local basic reproductive number, $R_0$. We find statistical evidence of human-to-human transmission ($p = 0.009$) in Sumatra but not in Turkey ($p = 0.114$). For Sumatra, the estimated household SAR was 29% (95% confidence interval [CI] 15%–51%). The estimated lower limit on the local $R_0$ was 1.14 (95% CI 0.61–2.14). Effective HPAI (H5N1) surveillance, containment response, and field evaluation are essential to monitor and contain potential pandemic strains.

Highly pathogenic avian influenza A (HPAI) subtype H5N1 is repeatedly crossing the species barrier to humans. Since December 2003, a total of 291 cases of HPAI (H5N1) have been reported in humans, resulting in 172 deaths (i.e., 59% case-fatality ratio) in 12 countries, mostly in Southeast Asia (1). Among these cases, 31 family clusters have been documented, ranging in size from 2 to 8 family members. How many of these clusters are due to a common avian source and how many are due to human-to-human transmission are important facts to determine. Should one of these HPAI (H5N1) strains gain the capacity for sustained human-to-human transmission, the resulting outbreak, if not contained, would spread worldwide through the global transportation network more rapidly than adequate supplies of vaccine matched to the new variant could be manufactured and distributed (2,3). We analyzed data from 2 of the largest of the familial clusters to ascertain if human-to-human transmission took place, and if so, how transmissible the strain was.

Methods

May 2006 Human Avian Influenza Family Cluster, Indonesia

During late April and early May 2006, a cluster of 8 cases of HPAI (H5N1) was detected and investigated by the Indonesian public health surveillance system in northern Sumatra (4–6). All case-patients were members of the same extended family. Seven of them resided within 3 adjacent houses in the village of Kubu Sembilang. The remaining patient resided with his immediate family in the village of Kabanjahe (~10 km away).

The index patient was a 37-year-old woman, thought to have been exposed to dead poultry and chicken fecal material before onset of illness. She also reportedly maintained a market stall that sold live chickens. Although her illness was not confirmed to have been caused by avian influenza (H5N1), her death on May 5, 2006, is suspected to be the result of HPAI (H5N1) infection because of her reported symptoms, illness progression, and prior contact with diseased or dead poultry.

Twenty members of her extended family are suspected to have been in contact with her, many during a family gathering on April 29, 2006 (7). At that time, she was manifesting symptoms (i.e., she had a heavy cough, was severely ill, and was prostrate). That night, 9 of these members slept in the same small room as she did (indicated by a black triangle in online Appendix Figure 1, available from www.cdc.gov/eid.)
December 2005 Human Avian Influenza Family Cluster, Eastern Turkey

From December 18, 2005, to January 15, 2006, a cluster of 8 confirmed influenza (H5N1) cases was detected in Dogubayazit District in eastern Turkey (online Appendix Figure 2, available from www.cdc.gov/EID/content/13/9/1348-appG2.htm) (10–13). These case-patients were among 21 members of 3 households located within 1.5 km of each other (14). All confirmed case-patients were hospitalized after onset of symptoms (9). Four of the confirmed case-patients died; the other 4 recovered (9). Ten of the remaining 14 household residents were hospitalized with avian influenza-like symptoms but were never confirmed to be infected with influenza (H5N1) (9). All but one of the hospitalized residents were children (6–15 years of age) (9).

Before onset of symptoms, 4 children from 1 household, 3 of whom had confirmed cases (including the index patient), were reported to have had close contact with the dead bodies of sick chickens (15). The 2 confirmed case-patients in the second household reportedly slaughtered a duck together on January 1, 2006, at the beginning of a die-off in the household’s flock (14). Two of the remaining confirmed case-patients lived in the third household and had no history of contact with sick or dying poultry. The remaining confirmed case occurred in a fourth residence located near the first household (10), but because we lacked information on the number of household members and the case-patient’s exposure history, we excluded it from these analyses. Most, if not all, of the 21 residents attended a dinner hosted by the family of the index patient on December 24, 2006, while he was symptomatic (8).

Statistical Methods

We used a previously developed statistical transmission model (16,17) to test whether human-to-human transmission occurred, and if it did, to estimate transmission parameters. In the model, persons mix with one another in households and between households. In addition, we include a common source of infection due to zoonotic exposure. Mathematical and statistical details are given in the online Technical Appendix (available from www.cdc.gov/EID/content/13/7/1348-Techapp.pdf).

Model of Probability of Transmission

We define $p_i$ as the probability that an infectious household member infects another household member in 1 day. If the distribution of the infectious period is known, we can obtain the household secondary attack rate (SAR$_h$) from $p_i$, defined as the probability that an infectious household member infects another household member over his or her infectious period. Similarly, we define the daily transmission probability ($p_e$) and the community SAR (SAR$_c$) for between household spread. Finally, we define the daily probability ($b$) that any person is infected from a zoonotic source. The contact structure used for parameter estimation is shown in the Figure. We assume that the distributions of the incubation and infectious periods are predetermined by the investigator.

We establish the likelihood function for each person and then for the whole population for statistical inference. The likelihood function for a person is equivalent to the probability of observing the realized data on that person throughout the outbreak. The likelihood function for a person labeled $i$ is built with the following steps: 1) Obtain the probability that person $i$ is infected by an infectious source labeled $j$ on day $t$, given person $i$ is not infected up to day $t-1$. If source $j$ is a person, this probability is $p_i$, for the same household, or $p_e$ for exposure in the community, multiplied by the probability of person $j$ being infectious on day $t$. The probability of person $j$ being infectious on day $t$ is derived from the symptom-onset day of person $j$ and the distribution of the infectious period. If source $j$ is zoonotic, the infection probability is $b$. The probability of escaping infection is simply 1 minus the corresponding probability of infection. 2) Take the product of the probabilities obtained in step 1 over all humans and zoonotic sources $j$ to obtain the probability of person $i$ escaping infection by any
infectious source on day $t$. 3) Take the product of the probabilities obtained in step 2 over all days before and including day $t$ to obtain the probability of person $i$ escaping infection up to day $t$. 4) If person $i$ is not infected by the end of the outbreak, the likelihood function for person $i$ is the product of the probabilities of person $i$ escaping infection up to the last day of observation. 5) If person $i$ is observed to have symptom onset on day $\tilde{t}$ and the infection time is known to be $t$, the probability of the data regarding person $i$ is the product of 3 pieces of information: a) the probability of person $i$ escaping infection up to day $t-1$, b) the probability that person $i$ is infected on day $t$, and c) the probability that the duration of the incubation period is $\tilde{t} - t$. Because we do not observe the infection time, the likelihood function for person $i$ is obtained by summing the above product, $a - c$, over all potential values of $t$.

The likelihood function for the whole population is the product of all the individual likelihood functions. In the event that human-to-human transmission occurs, SAR estimates are used to estimate the local basic reproductive number ($R_0$), which is defined as the average number of secondary cases infected by a typical index case-patient in the beginning of the outbreak (online Technical Appendix). There is potential for sustained transmission if $R_0$ is $>1$. If human-to-human transmission is determined to be occurring, then the above parameters are estimated from the symptom dates and contact information from the population under study. Data on exposed persons who do not become ill form an important component of the inference procedure.

**Statistical Test**

We set up a statistical test with the null hypothesis being that no human-to-human transmission occurs, that is, $p_1 = p_2 = 0$. The alternative hypothesis is either $p_1$ or $p_2$ is not equal to 0, or both are not equal to zero. The test statistic we use is proportional to the ratio of the maximum value of the likelihood function assuming the null hypothesis is true (null likelihood) and the maximum value of the likelihood function at the estimated parameter values (full likelihood).

Specifically, we define the likelihood ratio test statistic as $-2 \log \left( \frac{\text{null likelihood function}}{\text{full likelihood function}} \right)$. If no human-to-human transmission occurs, the 2 likelihood functions would be roughly equal, and we expect to see a likelihood ratio close to 1, and, thus, a likelihood ratio statistic close to 0. A large value of the likelihood ratio statistic is evidence of deviation from the null hypothesis. The question is how to obtain a reference set of the likelihood ratio statistic values that we would see under the null hypothesis. Given no human-to-human transmission, all the observed case-patients must have been infected by the zoonotic source. Since the exposure to the zoonotic source is assumed constant for each person on each day, the null likelihood function will not change if we reassign the infection and symptom status of the observed case-patients to a different group of people in the population. By performing such reassignment many times, we obtained a collection of datasets that were each equally likely to have been observed had there been no human-to-human transmission. The values of the likelihood ratio statistic calculated from these datasets form the null distribution for statistical testing. This method is referred to as a permutation test. The $p$ value is given by the proportion of the reference values that are equal to or larger than the observed likelihood ratio statistic value. More technical details are given in the online Technical Appendix.

The probability of infection by the zoonotic source may not be estimable together with SAR$_1$ or SAR$_2$ from an observed cluster. In such a situation, a statistical test of the occurrence of human-to-human transmission is still meaningful because the likelihood ratio test statistic is still estimable from the permuted datasets.

**Data Required**

A list of the inputs that are required for estimation and statistical testing are listed in the Table. Three categories of input parameters are required for this estimation model: outbreak-wide, individual level, and analysis parameters. The duration of the outbreak, the duration of the incubation period for the pathogen, and the minimum and maximum durations of the infectious period for the pathogen are the required outbreak-wide inputs. For each person, their residential location (neighborhood and household), their de-
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Demographic characteristics (sex and age), and whether they were a case-patient or not are required input parameters. Case-patients require additional input of their illness-onset dates, types of outcome, outcome dates, and whether or not they are the index patient in the outbreak. Hospitalization and treatment dates (considered prophylactic for nonpatients) are optional input parameters for each person. For each person who visits another residence during the outbreak period, his or her identifiers, the neighborhood and household visited, and the start and end dates of the visit are required inputs. Analysis-related inputs include the last date of community exposure to potential common sources of infection, the last date of observation, and inputs for $R_0$ estimation (mean number of residents per household and mean number of out-of-residence contacts per person per day). An expanded version of the model will require the input of other exposure information such as from schools or hospitals.

Results

For the outbreak in Indonesia, online Appendix Figure 1 shows that the incubation period had a probable range of 3–7 days and the infectious period, a probable range of 5–13 days. Thus, we let the incubation period have a uniform distribution of 3–7 days (mean 5 days) and the infectious period a uniform distribution of 5–13 days (mean 9 days). For the data shown in online Appendix Figure 1, only the household SAR (SAR$_h$) can be estimated. We determine that human-to-human spread did occur by rejecting the null hypothesis of no human-to-human transmission ($p = 0.009$). The estimated household SAR is 0.29 (95% confidence interval [CI] 0.15–0.51). Thus, a single infected person in a household infected another household member with the probability of 0.29. The average household size for rural Indonesia is ≈5 people. Because we do not have an estimate of the community SAR, we have an estimate of the lower limit of the local $R_0$, i.e., 1.14 with a 95% CI of 0.61–2.14. A sensitivity analysis on the distribution of the incubation and infectious period shows that the test and estimates for SAR$_h$ and $R_0$ are insensitive to uncertainty about these distributions within plausible ranges.

For the outbreak in Turkey, all the parameters are estimable, but we do not reject the null hypothesis of no human-to-human transmission ($p = 0.114$). Our estimate of the daily probability of infection from the common source is 0.011 (95% CI 0.005–0.025).

Discussion

We have presented statistical evidence that the strain of HPAI (H5N1) that caused the family cluster of human cases in northern Sumatra was spread from human to human and that the household SAR was 29%. This household
SAR is similar to statistical estimates for interpandemic influenza A in the United States (12.7%–30.6%) (18,19). The mean incubation period of this strain appears to have been ≈5 days, nearly twice as long as for past pandemic strains and current interpandemic strains of influenza. The CI for the estimated lower bound for the local $R_0$ covers 1. Therefore, even though we determined that human-to-human transmission probably occurred, whether the virus was capable of sustained human-to-human transmission is not clear. This virus may have required very close human contact to be transmitted. Even with no intervention, the finding that $R_0 = 1.14$ indicates that the chance that a single introduction would result in any further spread is ≈12%. In addition, the reported prophylactic use of oseltamivir may have played some role in limiting further spread. We did not find statistical evidence of human-to-human spread for the outbreak in eastern Turkey. This does not mean that no low-level human-to-human spread occurred in this outbreak, only that we lack statistical evidence of such spread. The power would be too low to detect such spread for an outbreak with 7 total cases and small SARs (17).

We did not consider the role of heterogeneity—such as age, sex, treatment status, or quarantine—in transmission. The parameters could be made to be functions of time-dependent covariates, as we have done with similar models (16,19,20). We can easily extend the model used here for covariates; however, we must have sufficient data to support such models.

Computer simulations have shown that the targeted use of influenza antiviral agents could be effective in containing a potential pandemic strain of influenza at the source (21,22), if initiated within 3 weeks of the initial case in the community, and if the $R_0$ is <1.8. This strategy, known as targeted antiviral prophylaxis, involves treating identified index patients in a mixing group and offering a single course of prophylaxis to the contacts of these index patients in predefined close contact groups, i.e., households at a minimum but also possibly neighborhood clusters, preschool groups, schools, and workplaces. In addition, the voluntary household quarantine of suspected close contacts of case-patients was recommended. Targeted antiviral prophylaxis at the household and neighborhood cluster level was carried out for the outbreak in Sumatra.

Ascertaining whether a potential pandemic strain of influenza is capable of sustained human-to-human transmission and estimating key transmission parameters are important. To estimate more than the household SAR, more detailed community data need to be collected. This would include a complete census of potentially exposed households and persons in the area where immediate transmission could occur from both potential zoonotic and human sources. Such data would enable estimation of important parameters and a more complete estimate of the $R_0$ rather than just the lower limit.

We have developed a software application, TRANSTAT, for implementing these analyses. This application provides a stand-alone environment for the entry, storage, and analysis of data from outbreaks of acute infectious diseases. A partial list of the input information is given in the Table. The statistical methods presented here can be applied to the data along with several standard epidemiologic tools. This information system would allow for real-time analysis and evaluation of control measures for an outbreak. We would encourage outbreak investigators to use this tool, taking care to input data on the exposed nonpatients as well as case-patients. The authors will provide a link to this software upon request.

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### References

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Technical Appendix

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The Likelihood

The Likelihood in a Prospective Design

We model the within household spread by the household transmission probability ($p_1$), defined as the probability that an infectious household member infects another household member in a 1 day period. The between household transmission probability ($p_2$) is the probability that an infectious person infects a person from a household different from his or her own in a 1 day period. Finally, $b$ is the probability that any person is infected from a zoonotic source due to 1 day of exposure. By prospective design we mean that a population of size $N$ are free of the disease at the beginning and followed up from day 1 to some day $T$. We assume the whole population is exposed to a constant zoonotic infective source throughout the observation period. Let $\tilde{t}_i$ be the symptom onset day for an infected person $i$. People who do not show any symptom by day $T$ will have $\tilde{t}_i = \infty$ by default. Let $c_{ij}(t)$ indicate whether there is a close contact (1) or a casual contact (0) between persons $i$ and $j$. The probability that an infective person $j$ infects a susceptible person $i$ on day $t$, is given by

$$p_{ji}(t) = p_1^{c_{ij}(t)} p_2^{1-c_{ij}(t)} f(t - \tilde{t}_j),$$

where $f(l)$ is the distribution of the infectious period. The probability that a susceptible person $i$ escapes infection from all infective sources on day $t$ is then given by

$$e_i(t) = (1-b) \prod_{j=1}^N p_{ji}(t).$$

Assume that the duration of the latent period $\delta$ has the distribution $g(l) = \Pr(\delta = l)$, $l = \delta_{\min}, \delta_{\min} + 1, \cdots, \delta_{\max}$. Define $\tilde{t} = \{\tilde{t}_j, j = 1, \cdots, N\}$, we construct the likelihood for person $i$ as
The Likelihood in Case-Ascertainment Design

Real epidemic data, such as the avian A(H5N1) influenza clusters discussed in this paper, generally come with the case-ascertained design instead of the prospective design, i.e., the whole cluster is ascertained upon the development of disease (i.e., symptom onset) of one or more index cases. To reduce the selection bias, the likelihood should be conditioning on the symptom status on the day when the index cases are ascertained. Let \( t_i^\alpha = \tilde{t}_i - \delta_{\text{max}} \) and \( t_i^\beta = \tilde{t}_i - \delta_{\text{min}} \) be the earliest and latest possible infection days of person \( i \), respectively. Let \( d_i \) denote the index case in the household of person \( i \). Following Yang et al. (1), the marginal probability of having symptom onset later than \( \tilde{t}_{d_i} \) is

\[
L_i^m(b, p_1, p_2 | \tilde{T}) = \sum_{t_{d_i} = \tilde{t}_{d_i}+1}^{\tilde{t}_i} \left\{ \left( \prod_{r=\tilde{t}_{d_i}+1}^{t_i-1} e_i(r) \right) \Pr(\tilde{t}_i > \tilde{t}_{d_i} | t) \right\} + \prod_{t=\tilde{t}_{d_i}+1}^{t_i} e_i(t).
\]

The joint conditional likelihood \( L'(b, p_1, p_2 | \tilde{T}) = \prod_i L_i^m(b, p_1, p_2 | \tilde{T}) \) will be maximized to obtain the maximum likelihood estimates. By conditioning, index cases do not contribute to the joint conditional likelihood.

The Permutation Test

The Hypotheses

Testing the existence of human-to-human transmission is equivalent to testing the hypotheses

\( H_0 : p_1 = p_2 = 0 \) vs. \( H_1 : p_1 > 0 \) or \( p_2 > 0 \),

where \( H_0 \) is the null hypothesis and \( H_1 \) is the alternative hypothesis.
The Test Statistic

When there is no person-to-person transmission, i.e., \( p_1 = p_2 = 0 \), expression (1) reduces to

\[
e_i(t) = (1 - b).
\]

Let \( L_0^c(\tilde{t}) \) denote the likelihood for the null model. The test statistic is defined as the likelihood ratio statistic:

\[
\hat{\lambda} = -2 \log \frac{\sup_b L_0^c(b \mid \tilde{t})}{\sup_b L^e(b, p_1, p_2 \mid \tilde{t})}.
\]

The Null Distribution Based on Resampling

Yang et al. (2) discussed the permutation test for the prospective design, but the same logic applies to the case-ascertained design as well. When the observed data are truly generated from \( H_0 \), i.e., there is no human-to-human transmission, we can reassign all of the observed symptom onset days (and associated infection status) to a different subset of people, and every such rearrangement has the same likelihood \( L_0^c \). By permuting symptom onset days across the population, we obtain an empirical distribution of the test statistic which can then be used to approximate the null distribution of the test statistic under \( H_0 \). Let \( (\tilde{t}_1^{[k]}, \tilde{t}_2^{[k]}, \cdots, \tilde{t}_N^{[k]}) \) be the \( k^{th} \) permuted sample of \( (\tilde{t}_1, \tilde{t}_2, \cdots, \tilde{t}_N) \), and \( \hat{\lambda}^{[k]} \) be the corresponding test statistic, \( k = 1, \cdots, M \). Then the p-value is given by \( \frac{1}{M} \sum_k I(\hat{\lambda} \geq \hat{\lambda}^{[k]}) \). This permutation test can be refined by varying symptom onset days of infected individuals in any given permuted data, as long as the null likelihood \( L_0^c \) is unchanged. The refined permutation test resamples data from a much larger sampling space and thus can attain higher statistical power. Technical details of these resampling methods can be found in Yang et al. (2). However, an important distinction from Yang et al. (2) is that the index cases should not be involved in the refinement stage for the case-ascertained design, because changing the symptom onset days of the index cases also changes the index or non-index status and makes it difficult to keep \( L_0^c \) constant.
Three Important Functions of Estimated Parameters

We summarize the degree of transmissibility of the infectious agent by the household secondary attack rate (SAR₁), where \( \text{SAR}_1 = \sum_i f(i) \left(1 - (1 - p_1)^i\right) \), and the community secondary attack rate, (SAR₂), where \( \text{SAR}_2 = \sum_i f(i) \left(1 - (1 - p_2)^i\right) \).

The basic reproductive number (\( R_0 \)) is the average number of people that a typical infected person infects in an entirely susceptible population. For a community of size \( N \) with average household size \( M \), we have the local \( R_0 = (M - 1) \times \text{SAR}_1 + (N - M) \times \text{SAR}_2 \). The maximum likelihood estimates of \( \text{SAR}_1 \) and \( \text{SAR}_2 \) are used to get an estimate of \( R_0 \). If only an estimate of \( \text{SAR}_1 \) is available, then a lower bound on \( R_0 \) can be estimated.

References
