We investigated the capacity of internal border control to limit influenza spread in an emergent pandemic in the context of Australia, a country with a low-population density and geopolitical boundaries that may facilitate restrictions. Mathematical models were used to study the time delay between epidemics in 2 population centers when travel restrictions were imposed. The models demonstrated that population size, travel rates, and places where travelers reside can strongly influence delay. The model simulations suggested that moderate delays in geographic spread may be possible with stringent restrictions and a low reproduction number, but results will be sensitive to the reproduction number and timing of restrictions. Model limitations include the absence of further importations and additional control measures. Internal border control may have a role in protecting domestic centers early in a pandemic, when importations are sparse. Our results may be useful for policymakers.

Commercial air travel has increased dramatically since the last pandemic of influenza (1). The number of international tourist arrivals recorded worldwide in 2004 was 763.2 million; 43% of these arrivals were by air (2). This increase in international travel has heightened the risk for the global spread of infectious diseases (1).

Long-distance domestic routes also carry high volumes of travelers: an estimated 40.4 million passengers traveled on Australian domestic airlines in the year ending June 30, 2005 (3), and 660 million traveled on US domestic airlines during 2005 (4). Rapid and accessible long-distance transportation facilitates the geographic spread of diseases, even those, such as influenza, that have a short incubation period (5).

If an influenza pandemic emerges, the first attempts to control its spread are likely to be made at its source, as suggested in recent modeling papers (6,7). However, if these strategies fail, individual governments will need to implement strategies to manage the pandemic when it arrives on their borders. In addition to well-publicized options for control, including antiviral prophylaxis and quarantine (6,7), travel restrictions, both external and internal, may play a role in reducing the geographic spread of the virus (8–11).

Restrictions on travel can have a sizeable economic and social impact, as seen in affected nations during the crisis with severe acute respiratory syndrome (SARS). In many countries, stringent travel restrictions will not be feasible because of high population densities and highly connected networks of transportation, infrastructure, and trade. These caveats do not apply to Australia, an island comparable in size to the United States but with a population of only 20 million. This population is concentrated in 5 large cities, along with smaller centers, primarily along its eastern and southern coastlines. These centers are widely separated; travel between them is primarily by air. During the 1918 pandemic, Australia delayed the onset of the pandemic by 1 year by imposing external border control (12).

We used mathematical models to make predictions about the effectiveness of travel restrictions and to explore the sensitivity of these predictions to disease and demographic factors. Typically, modeling studies of influenza spread are focused on predicting international or national spread between major hubs on the global air-transportation network (8,9,13), which is certainly important. In contrast, we examine the effects of travel restrictions on 2-city routes with differing characteristics. This simpler setting allows a more detailed exploration of how the delay between epidemics in 2 connected locations depends on travel restrictions, population sizes, travel rates, residence

*National Centre for Immunisation Research and Surveillance of Vaccine Preventable Diseases, Sydney, New South Wales, Australia; †The University of Sydney, Sydney, New South Wales, Australia; and ‡Australian National University, Canberra, Australian Capital Territory, Australia
of travelers, and the transmissibility of the influenza virus, with relevance to large and small centers. The analysis assumes case-patients arriving from overseas have a negligible effect, so the results apply primarily during the early stage of a pandemic. Simulating the effect of internal travel restrictions in Australia is relevant to countries with similar demographic characteristics, such as Russia, Canada, and New Zealand. The aims of our analysis were to explore the role of travel restrictions in slowing the geographic spread of an influenza pandemic and to simulate the effects of such restrictions in the context of Australia.

Methods

Two simple scenarios (Figure 1A) were used to assess the likely impact of travel restrictions on the spread of a pandemic in Australia. In the first, it was assumed that the initial cases occurred in Sydney. The growth of this epidemic and its resultant spread to Melbourne in the presence of travel restrictions were simulated. This scenario is indicative of the spread to other large centers with similar travel volumes. In the second scenario, the initial case was assumed to occur in Darwin, a smaller Australian city in close proximity to Southeast Asia, and the growth of this epidemic and spread to Sydney were simulated. The Darwin-to-Sydney scenario, with a comparatively low travel volume, represents the situation of containing the epidemic within a smaller town through the use of travel restrictions. Key parameters and assumptions are summarized in Table 1.

Data

Average daily volumes of domestic air travel between Sydney, Melbourne, and Darwin were obtained from the Australian Domestic Airline Activity report (17). Only direct flights were considered. Seasonal variations in the volume of air traffic were not taken into account. Approximately 78% of the traffic from Sydney to Melbourne and 70% of the traffic from the Northern Territory to the eastern Australian states is by air (18).

As a separate indicator of travel volumes that incorporates the average length of stay and information on the origin of travelers, we used survey estimates of nights stayed by domestic visitors to the 3 study destinations (Melbourne, Sydney, and Darwin). The data were obtained from the state government tourism websites for Victoria (20), New South Wales (NSW) (19), and the Northern Territory (21). Because details on visitor origin were only obtained at the state level, we assumed that each person in that state would make an equal contribution to visitor nights in the destination city. These values were then used to estimate the proportions of the travel volume due to each of the 2 cities on a route and to modify force of infection calculations by incorporating the average length of stay. The travel rates (weighted by length of stay) used in the simulations are provided in Table 1. Demographic data on cities and states were acquired from the Australian Bureau of Statistics population estimates for 2004 (16).

Model Structure

Simulations of influenza epidemics were computed by using a stochastic SIR model, in which the population is separated into 3 mutually exclusive classes: susceptible (S), infectious (I), and recovered (R). A stochastic model can capture random variation near the beginning of an epidemic, when the number of infectious persons is small. Homogeneous mixing is assumed, i.e., all susceptible members of the population in a city are equally likely to be infected by a given infectious person.

A schematic of the model is given in Figure 1B, and the defining equations are presented in the online Appendix, part A (available from www.cdc.gov/EID/content/13/7/1038-app.htm). The model evolves in discrete time, with the step length equal to 1 day. This time frame accords with real-life epidemics, for which incidence and other epide-
miologic data are usually recorded daily. The discrete time structure simplifies the introduction of a variable infectivity profile, incorporating a latent, noninfectious period and a changing degree of infectivity for each person during the course of his or her illness. This feature of the model is supported by virus-shedding studies (22) and enables us to contrast the effect of a highly peaked infectivity profile, similar to that used by Ferguson et al. (6), with the effect of a constant infectivity profile (7), as depicted in Figure 1C.

A key factor governing the effectiveness of our travel restrictions is the average doubling time of the attack rate during the early stages of the epidemic, when growth is exponential. The doubling time is determined by the basic reproduction number $R_0$, defined as the average number of secondary infections due to a single primary infected person in a completely susceptible population, and the form of the infectivity profile. The infectivity profile primarily influences the growth rate through the mean time (or serial interval) between cases: $\approx 2.8$ days for the peaked infectivity function and 4 days for the flat infectivity function used here. The doubling time depends linearly on the serial interval so that epidemics that use the peaked infectivity function will grow faster. This feature of the model is changing degree of infectivity for each person during the course of his or her illness. This feature of the model is supported by virus-shedding studies (22) and enables us to contrast the effect of a highly peaked infectivity profile, similar to that used by Ferguson et al. (6), with the effect of a constant infectivity profile (7), as depicted in Figure 1C.

Although influenza appears to be a highly infectious disease, with regular winter epidemics, this is largely due to its short incubation period and genetic drift, which nullifies preexisting immunity. Thus, literature estimates of the effective reproduction number for influenza are typically $<4$ (14) (whereas for measles $R_0$ is 20 [15]), although in localized outbreaks it can be considerably higher (23). We take $R_0$ to be in the range 1.5–3.5, which corresponds to attack rates of 58%–97% (including subclinical infections) in a population without prior immunity or behavioral changes in response to the pandemic.

The total period of infection, including latent period, was assumed to be 6 days (7). For each infectivity profile, the latent period was $\approx 1$ day, which is at the low end of literature estimates (other researchers have used values of 1–4 days [8,9,24]). Spread from city to city is incorporated by assuming that each person is equally likely to travel; the daily travel rates were estimated from the data sources described above. This assumption was pessimistic, since symptomatic infected persons may not travel, but it did not greatly influence the results (online Appendix, part A).

Travel restrictions were implemented as a reduction of the rate of all forms of travel. For this analysis, reductions of 80%, 90%, and 99% were compared with the base case of unrestricted travel. The values of 80% and 90% might be realistic reduction targets, whereas the value of 99% indicates what near-perfect compliance might achieve. Travel restrictions were switched on in the model at some time (measured in weeks) after the initial case occurred and remained on for the rest of the simulations.

The principal measure used in this analysis for gauging the effect of travel restrictions is $T_{20}$, the delay between the epidemic’s becoming established in city 1 and taking off in city 2. We considered the outbreak to have taken off in a city once there were 20 current infectious cases—hence, the notation $T_{20}$ for the delay between the epidemics. This choice conveniently limited comparisons to simulated epidemics that do take off. Since the model is stochastic, $T_{20}$ is random, and the results shown in the graphs are for $m_{20}$, the median value for outbreaks that take off. Ranges, when given, cover 90% of outbreak simulations.

### Table 1. Summary of parameter values, assumptions, and sources used in models of the effect of travel restrictions on pandemic influenza in Australia*

| Variable/concept                  | Value (range)/assumption                        | Source/interpretation                                         |
|-----------------------------------|------------------------------------------------|--------------------------------------------------------------|
| Reproduction no. ($R_0$)          | $1.5–3.5$                                       | Mills (14)                                                   |
| Infectivity function ($\rho$)     | Flat or peaked†                                 | Ferguson (7,8)                                               |
| Latent period                     | 1 (1–2 in sensitivity analysis) d(s)            | Ferguson (6)                                                 |
| Infectious period                 | 5 d                                            | Literature suggests 4–7 d in adults (6,7)                    |
| Mixing                            | Homogenous (within city)                       | Modeling literature (15)                                    |
| Propensity to travel              | Everyone equal                                 | Assumption                                                  |
| Populations                       | Sydney (4.2 million), Melbourne (3.6 million), Darwin (110,000) | BTRE figures (17,18), NSW, and Victoria Tourism reports (19,20) |
| Travel rate‡ Sydney ↔ Melbourne (weighted by stay length) | $(4.7 \times 10^7, 8.9 \times 10^3)$ | ABS figures (16)                                           |
| Travel rate‡ Sydney ↔ Darwin (weighted by stay length) | $(9.2 \times 10^4, 4.4 \times 10^3)$ | BTRE figures (17,18), NSW, and NT Tourism reports (19,21) |
| Travel restrictions              | 20%, 10%, or 1% of current levels               | Assumption                                                 |
| Time between 20 current cases in city 1 and city 2 ($T_{20}$) | Random variable ($T_{20}$), different for each simulation. Median value over all simulations is given by $m_{20}$. | Output variables used to measure effect of travel restrictions |

*ABS, Australian Bureau of Statistics; BTRE, Bureau of Transport and Regional Economics; NSW, New South Wales; NT, Northern Territory.
†See Figure 1, panel C, for shapes used.
‡This assumes a constant travel rate over the year with no seasonal variation in travel volumes.
The simulations were run with MATLAB version 7.04 (The MathWorks, Natick, MA, USA) with Poisson random variables simulated by the poissrnd function in version 5.02 of the Statistics Toolbox (MathWorks). Our results are based on 10,000 runs of the model.

Motivated by the results of the simulation study, we then analyzed the effects of city size and travel rates by using a deterministic approximation of the above model (details given in the online Appendix, part B). This approximation has the advantage of being much simpler to use in analyzing sensitivity to these factors, while reproducing the average behavior of the stochastic model.

**Results**

**Scenario 1 (Sydney to Melbourne)**

The median and mean numbers of days until there are 20 infectious persons in Sydney for an epidemic that began with 1 infectious person in Sydney on day 0 are presented in Table 2. Figure 2 illustrates how \( m_{20} \), the median time between the day when the number of infected persons first reached 20 in Sydney and the day when the number of infected persons first reached 20 in Melbourne, depends on \( R_0 \), the form of the infectivity profile, and the timing and severity of travel restrictions.

Each of the graphs covers 1 of the 6 combinations of the 3 values of \( R_0 \) and 2 infectivity profiles. The 4 curves shown on each graph describe the median values for each of the 4 levels of travel restrictions (none, 80%, 90%, and 99%), applied at delays from importation of the first case from 0 to 6 weeks (8 weeks for \( R_0 = 1.5 \)). The gray panes highlight the time during which the epidemic grows from 20 to 1,000 cases in Sydney.

The travel restrictions are most effective for the optimistic assumption \( R_0 = 1.5 \) and constant infectivity (Figure 2A). Figure 2B and C more closely resemble the epidemic growth rates used in recent modeling papers (6, 7). In Figure 2B \( (R_0 = 1.5, \text{peaked infectivity}), \) an increase in \( m_{20} \) from 22 to 33 days is seen for 80% restrictions, with a further increase to 52 days for 99% restrictions, if applied immediately. These improvements appear robust for delays of up to 4 weeks, but in fact a sizeable proportion of the simulations have spread to Melbourne by this point. This effect is illustrated in Figure 3A and B, in which we compare the full distribution of \( T_{20} \), in the presence of 99% travel restrictions applied at the 2- and 4-week marks, respectively. Both distributions are bimodal, but in Figure 3B, the first mode is substantial. This difference arises because a large proportion of simulated outbreaks spread to Melbourne between the 2- and 4-week marks for this combination of disease parameters, a finding that emphasizes that timing can be critical for the success of travel restrictions. Under the pessimistic assumption of \( R_0 = 3.5 \) and peaked infectivity, the impact of travel restrictions is muted, and a delay of just 2 weeks renders the restrictions ineffective.

**Scenario 2 (Darwin to Sydney)**

For an epidemic originating in Darwin, the median times until there are 20 infectious persons in Darwin are almost identical to those for scenario 1 (Table 2), although the 90% ranges are a little wider. In this scenario, \( m_{20} \) is the median time between the first day on which there are 20 infected persons in Darwin and the day on which there are 20 currently infected persons in Sydney. The effects of \( R_0 \), the infectivity function, and the delay in and severity of travel restrictions are captured in Figure 4.

These results, presented in the same format as Figure 2, show 2 key differences from those in scenario 1. The median delay, \( m_{20} \), is shorter in scenario 2, given the same combination of disease parameters, as is the time interval over which restrictions can be applied effectively. This finding appears counterintuitive because the volume of travelers on the Darwin-to-Sydney route is much smaller than that on the Sydney-to-Melbourne route.

By using the simpler model described in the online Appendix, part B, we performed a sensitivity analysis (online Appendix Figure, available from www.cdc.gov/EID/content/13/7/1038-appG.htm) on the effect of city size and travel rates on epidemic spread. This analysis implies that in scenario 2, in which there is a large difference in population size (Darwin:Sydney \( \approx 1:40 \)), infection of susceptible travelers from Sydney is the primary reason for the rapid intercity spread, despite the lower rate of travel for Sydney residents on this route. The online Appendix Figure, panel B, shows that this effect would be reduced if the rate at which Sydney residents travel to Darwin were much lower than that for Darwin residents traveling to Sydney. Such a reduction could be achieved by applying tighter restrictions on Sydney-based travelers.

| \( R_0 \) | Constant infectivity profile† | Peaked infectivity profile† |
|----------|-------------------------------|-----------------------------|
|          | Median, d | 90% range, d | Mean, d | Median, d | 90% range, d | Mean, d |
| 1.5      | 24        | 13–46        | 25.9    | 15        | 8–31        | 16.4    |
| 2.5      | 12        | 8–21         | 13.0    | 8         | 5–14        | 8.4     |
| 3.5      | 9         | 6–14         | 9.5     | 6         | 4–10        | 6.2     |

*\( R_0 \): reproduction number.
†The constant infectivity profile assumes that a person is equally infectious throughout their infectious period; the peaked infectivity profile assumes that they are most infectious early in the infectious period (see Figure 1, panel C, for the profiles used).
The ratio of city populations also influences the time interval when restrictions can be applied effectively. Travel restrictions were less effective if applied after the time at which there were 20 current cases in Darwin (Figure 4). This feature was illustrated by Figure 3C and D, which show the full distributions of $T_{20}$ for 2- and 4-week delays in restrictions, respectively. For a 2-week delay (Figure 4C), most outbreaks were delayed but a sizeable minority were not. A 4-week delay (Figure 4D) nullified any impact of the restrictions for this scenario. If, however, travel restrictions were applied immediately after the first case was detected, the increase in $T_{20}$ due to restrictions was almost identical to the increases described in scenario 1.

Now consider a situation in which a small isolated center (town A, population 1,000) attempts to remain pandemic free. Let us assume that on any given day, $N$ visitors stay in the town, and $N_t$ town members visit pandemic-affected regions. A simple stochastic model of disease spread (online Appendix, part D) can predict the probability that the outbreak can be kept out of town A in terms of $N$ and $R_0$ (online Appendix, part D). Predictions from this model agree well with simulations, as shown in the online Appendix Figure, panel C. These results indicate that travel restrictions are likely to prevent an outbreak if $N$ is reduced to $\approx 1/10$ per day.

**Sensitivity to Other Factors**

The sensitivity of the results to the duration of infection and form of the infectivity function were entirely a result of the change in the epidemic growth rate. If, for example, an additional day of latent infection were added, then the delays in spread, when the flat and peaked infectivity functions were used, were $\approx 25\%$ and $\approx 37\%$ longer, respectively, which is a considerable effect. However, epidemic

---

**Figure 2.** For an epidemic beginning in Sydney, the value of the median time delay, $m_{20}$, in the presence of travel restrictions applied at a delay of 0–6 weeks (10 and 8 weeks in [A] and [B], respectively). Assumptions are A) reproduction number ($R_0$) = 1.5, constant infectivity profile; B) $R_0 = 1.5$, peaked infectivity profile; C) $R_0 = 2.5$, constant infectivity profile; D) $R_0 = 2.5$, peaked infectivity profile; E) $R_0 = 3.5$, constant infectivity profile; F) $R_0 = 3.5$, peaked infectivity profile. The gray panes cover the periods when the epidemic grows from 20 to 1,000 infected people in Sydney; dotted, dashed, dash-dotted, and solid lines correspond to 99%, 90%, 80% and no travel restrictions, respectively.

**Figure 3.** Distributions, based on 10,000 simulations, of the time delay, $T_{20}$, given reproduction number ($R_0$) = 1.5 and the peaked infectivity function, with 99% travel restrictions imposed in scenario 1 (A) and (B) and scenario 2 (C) and (D). Scenario 1 simulates an epidemic beginning in Sydney and spreading to Melbourne. In scenario 2, the epidemic begins in Darwin and spreads to Sydney. In (A) and (C), the restrictions are imposed after 2 weeks; in (B) and (D), they are imposed after 4 weeks.
growth rates in past pandemics are typically not consistent with longer latent periods and low values of $R_0$, so these additional delays should be viewed with caution.

The sensitivity to the estimated travel volumes was relatively weak: increasing or reducing travel by a factor of 2 in each direction increases or reduces the delay by 1.5–7 days, and 4.5 or 2.5 days as compared to data in Figures 2 and 4, respectively. These results are consistent for both scenarios.

Discussion

The simulations we describe showed that although travel restrictions might delay the spread of an influenza epidemic between 2 cities by several weeks, this delay is highly sensitive to assumptions about the transmissibility of the influenza virus. A more surprising result is that the delay is also sensitive to the ratio of city sizes, differences in travel rates, and the originating city. In particular, the modeling suggests that if the epidemic begins in a smaller town, restricting visitors from entering or leaving that town is important.

Moderate delays in the pandemic could be achievable when the epidemic growth rate is low. The growth rate can be estimated from case counts during an epidemic and used in a simple formula to predict the delay due to travel restrictions (online Appendix, part D). These predictions could provide practical estimates of the benefits of longer term travel restrictions based on the first clusters of cases during an outbreak. For smaller communities with low travel rates, the probability of preventing an outbreak can also be estimated (online Appendix, part D), with good agreement with the results of our simulations (online Appendix Figure, panel C). If the estimated growth rate is high (e.g., assumptions used in Figures 2 and 4 with $R_0 = 3.5$, peaked infectivity), the additional median delay between 20 cases occurring in city 1 and 20 cases occurring in city 2 might be just 3 days, providing little benefit from longer term implementation of travel restrictions.

Our results do not account for additional importations. Thus, they are most applicable to the arrival of a pandemic in Australia, while the pandemic outside Australia remains contained or border control is effective. Our simulated delays will be overestimates if additional importations are substantial. Another concern is that stringent travel restrictions may be required for several weeks to maximize delays in spread. Inevitably, such restrictions would cause economic and social disruption, which must be balanced against any benefits from delaying the domestic spread of an epidemic.

If combined with restrictions on overseas travel, restrictions on internal travel may have a role in pandemic control, even for major centers. However, the economic impact of restrictions in major centers could be enormous, with severe consequences for service and travel industries, as seen in the SARS crisis (25), and the potential to affect trade and other sections of the economy. Some of the benefits and costs of reduced travel may also accrue without restrictions, with persons avoiding travel because of perceived risks. Our modeling suggests that travel restrictions could have a greater effect in more isolated communities that lack international ports.

Figure 4. For an epidemic beginning in Darwin, the value of the median time delay, $m_{20}$, in the presence of travel restrictions applied at a delay of 0–6 weeks (8 weeks in [A] and [B], respectively). Assumptions are (A) reproduction number ($R_0$) = 1.5, constant infectivity profile; (B) $R_0 = 1.5$, peaked infectivity profile; (C) $R_0 = 2.5$, constant infectivity profile; (D) $R_0 = 2.5$, peaked infectivity profile; (E) $R_0 = 3.5$, constant infectivity profile; (F) $R_0 = 3.5$, peaked infectivity profile. The gray panes cover the periods when the epidemic grows from 20 to 1,000 infected people in Darwin. Dotted, dashed, dash-dotted, and solid lines correspond to 99%, 90%, 80%, and no travel restrictions, respectively.
The travel restrictions we discussed have been examined in isolation, without consideration of other disease control measures. Other measures could lower the effective value of the reproduction number, or even curtail the epidemic; in these circumstances, reducing all travel by only 80% might be beneficial. Alternatively, if the $R_e$ is much higher than used here (23), internal travel restrictions would be ineffective. Limitations of our modeling approach are summarized in Table 3.

The key points in our study are that delays induced by internal border control are strongly influenced by epidemic growth rates and demographic factors such as the relative sizes of cities, travel rates, and the origin of travelers. When used without other control measures, stopping at least 99% of travel would be required to significantly increase time available for vaccine production and distribution. Although any delay in spread might be attractive for logistical purposes, the economic impact of such restrictions may be prohibitive if sustained for more than a few days. In view of these points, the situation in which they might be most applicable for extended use is in the protection of small, relatively isolated centers.

Acknowledgments

We thank colleagues at Australian National University and the University of Melbourne for helpful discussions on methods.

Partial funding from the Department of Health and Ageing and National Health and Medical Research Council grants 410224 and 358425 is gratefully acknowledged. The National Centre for Immunisation Research and Surveillance of Vaccine Preventable Diseases is supported by The Australian Government Department of Health and Ageing, the NSW Department of Health, and the Children’s Hospital at Westmead.

Dr Wood is a mathematician in the field of infectious disease modeling at the National Centre for Immunisation Research and Surveillance in Sydney, Australia. His current research interests are in influenza and other vaccine-preventable diseases, as well as underlying mathematical techniques used in modeling.

References

1. Gezairy HA. Travel epidemiology: WHO perspective. Int J Antimicrob Agents. 2003;21:86–8.
2. World Tourism Organization. Tourist arrivals by means of transport (2004). Nov 2005. [cited 2006 Feb 2]. Available from http://www.world-tourism.org/facts/eng/pdf/indicators/ITA_Transps.pdf
3. Australian Bureau of Transport and Regional Economics. Australian domestic airline activity 2004–05. Aviation statistics 102. [cited 2005 Dec 15]. Available from http://www.btre.gov.au/statistics/aviation/docs/Domestic_airlines_2004-05.pdf
4. U.S. Bureau of Transportation Statistics. 2005 domestic airline passenger traffic up 4.1 percent from 2004. 2006. [cited 2006 Mar 24]. Available from http://www.bts.gov/press_releases/2006/bts013_06/pdf/bts013_06.pdf
5. The Committee to Advise on Tropical Medicine and Travel (CATMAT) and the National Advisory Committee on Immunization (NACI). Statement on travel, influenza, and prevention. Can Commun Dis Rep. 2005;31:1–8. Erratum in: Can Commun Dis Rep. 2005;31:115.
6. Ferguson NM, Cummings DAT, Cauchemez S, Fraser C, Riley S, Meeyai A, et al. Strategies for containing an emerging influenza pandemic in Southeast Asia. Nature. 2005;437:209–14.
7. Longini IM, Nizam A, Xu S, Unghusuk K, Hanshaoworakul W, Cummings DAT, et al. Containing pandemic influenza at the source. Science. 2005;309:1083–7.
8. Ferguson NM, Cummings DAT, Fraser C, Cajka JC, Cooley PC, Burke DS. Strategies for mitigating an influenza pandemic. Nature. 2006;442:448–52.
9. Cooper BS, Pitman RJ, Edmunds WJ, Gay NJ. Delaying the international spread of pandemic influenza. PLoS Med. 2006;3:e212.
10. Hollingsworth TD, Ferguson NM, Anderson RM. Will travel restrictions control the international spread of pandemic influenza? Nat Med. 2006;12:497–9.
11. Markel H, Stern AM, Navarro JA, Michalsen JR, Monto AS, DiGiavanni C Jr. Nonpharmaceutical influenza mitigation strategies, US communities, 1918–1920 pandemic. Emerg Infect Dis. 2006;12:1961–4.
12. NSW Legislative Assembly. Report of the Director-General of Public Health NSW for the year 1919 and the report on the influenza epidemic 1919. Guillick (Australia): Sydney William Applegate; 1920.
13. Grais RF, Ellis JH, Glass GE. Assessing the impact of airline travel on the geographic spread of pandemic influenza. Eur J Epidemiol. 2003;18:1065–72. Erratum in: Eur J Epidemiol. 2004;19:395.
14. Mills CE, Robins JM, Lipsitch M. Transmissibility of 1918 pandemic influenza. Nature. 2004;432:904–6.
15. Anderson R, May R. Infectious diseases of humans. Oxford (UK): Oxford University Press; 1991.

Table 3. Limitations and effects of modeling effects of border control on pandemic influenza, Australia

| Limitations | Effects |
|-------------|---------|
| Reproduction number ($R_0$) and infectivity function for pandemic influenza are unknown. Further importations not considered. | Larger $R_0$ and a shorter average time between infections would reduce effectiveness of restrictions. Frequent importations would greatly reduce benefits of internal restrictions for cities with international airports or ports. Reductions in transmission would increase effectiveness of restrictions. |
| Other control measures (pharmaceutical and social distancing) are not considered. Heterogeneous mixing and travel patterns are not considered. | Heterogeneity could increase or reduce delays in epidemic spread. For example, high transmission among infrequent travelers (e.g., the elderly, children) would make restrictions more effective. |
| Travel rates and restrictions are based on air-travel volumes alone. | Restrictions would prevent no more than 80% of travel if non–air travel remains unrestricted, which would considerably reduce effect of restrictions. |
| Seasonal variation in travel and transmissibility are not considered. | Could lead to less or more effective restrictions if arrival of pandemic is in winter/summer. |
Internal Border Control and Spread of Pandemic Influenza

16. Australian Bureau of Statistics. Population projections, Australia 2002 to 2101. [cited 2005 Jan 12]. Available from http://www.abs.gov.au

17. Australian Bureau of Transport and Regional Economics. Australian domestic airline activity. Aviation Statistics Domestic Monthly, 2005;102. [cited 2005 Dec 15]. Available from http://www.btre.gov.au/statistics/aviation/docs/domestic_airlines_0506.pdf

18. Australian Bureau of Transport and Regional Economics. Passenger movements between Australian cities, 1970–71 to 2030–31. [cited 2006 Dec 13]. Available from http://www.btre.gov.au/docs/info-sheets/is26/is26.pdf

19. Tourism New South Wales. Travel to Sydney: year ended June 2005. 2005. [cited 2005 Jan 12]. Available from http://corporate.tourism.nsw.gov.au/corporate-live/downloads/research/sydney%20jun%202005.pdf

20. Tourism Victoria. Domestic visitation to regions of Victoria: year ending June 1999–2005. 2005. [cited 2005 Jan 12]. Available from http://www.tourism.victoria.com.au

21. Northern Territory Tourism. Darwin tourism region, results for the year ending June 2005. [cited 2005 Jan 12]. Available from: http://www.nttc.com.au/nt/system/galleries/download/NTTC_Research/June_quarterly_report_0405.pdf

22. Hayden FG, Fritz R, Lobo MC, Alvord W, Strober W, Straus SE. Local and systemic cytokine responses during experimental human influenza virus infection. Relation to symptom formation and host defense. J Clin Invest. 1998;101:643–9.

23. Fraser C, Riley S, Anderson RM, Ferguson NM. Factors that make an infectious disease outbreak controllable. Proc Natl Acad Sci U S A. 2004;101:6146–51.

24. Gani R, Hughes H, Fleming D, Griffin T, Medlock J, Leach S. Potential impact of antiviral drug use during influenza pandemic. Emerg Infect Dis. 2005;11:1355–62.

25. Smith DR, Sommers T. Assessing the economic impact of public health emergencies in international concerns: the case of SARS. Geneva: World Health Organization; 2003.

Address for correspondence: James Wood, NCIRS, Level 2, Children’s Hospital at Westmead, Westmead, New South Wales, Australia 2145; email: jamesw5@chw.edu.au

The opinions expressed by authors contributing to this journal do not necessarily reflect the opinions of the Centers for Disease Control and Prevention or the institutions with which the authors are affiliated.
Appendix

(A) Simulation Model

The model equations for a single city are given by

\[ S(t+1) = S(t) - X(t), \]
\[ i(t+1) = X(t), \]
\[ R(t+1) = i(t-D+1), \]
\[ I(t+1) = \sum_{d=0}^{D-1} i(t-d+1), \]

where \( i(t) \) denotes the number of new infected persons on day \( t \) and is determined by the Poisson random variable \( X(t) \) with mean \( \mu = S(t)p(t) \). Here,

\[ p(t) = 1 - \prod_{d=0}^{D-1} \left( 1 - \frac{S(t)p_2(t)}{N} \right)^{\rho(d)(t-d)} \]

is the probability on day \( t \) of a given susceptible person becoming infected. This equation corresponds to an illness with total infection period of \( D \) days and a varying degree of infectivity \( \rho(\tau) \) over the course of each infection. Note that \( \tau = 0, 1, \ldots, D-1, \) and \( \Sigma \rho(\tau) = 1. \)

Multiple cities are accommodated by introducing a pair of labels to indicate a person’s city of origin and destination as well as a rate of travel \( A_{kl} \) from city \( k \) to city \( l \) per day. The situations considered in this article involve only 2 cities, with just 2 rates of travel, \( A_{12} \) and \( A_{21} \). We set \( A_{12} = A_1 \) and \( A_{21} = A_2 \), for convenience. The epidemic equations for city 1 become

\[ S_1(t+1) = S_1(t) - X_1(t), \]
\[ i_1(t+1) = X_1(t), \]
\[ R_1(t+1) = i_1(t-D+1), \]
\[ I_1(t+1) = \sum_{d=0}^{D-1} i_1(t-d+1). \]

Here, \( X_1(t) \) is a Poisson random variable with mean \( \mu_1(t) = S_1(t)[A_1 p_2(t) + (1-A_1)p_1(t)] \), where \( p_1(t) \) is given by

\[ p_1(t) = 1 - \prod_{d=0}^{D-1} \left( 1 - \frac{S(t)p_2(t)}{N} \right)^{\rho(d)(1-A_1)(t-d)+A_2(t-d)}, \]
Note that the equations for city 2 and $p_2(t)$ can be obtained by replacing the label 1 with 2 and vice versa in the previous 2 equations.

The sensitivity of the results to the assumption that all infected persons travel can be tested by making an optimistic assumption that two thirds of infected persons are asymptomatic and do not travel. This leads to an additional median delay of between 2 days (reproduction number $R_0 = 3.5$, peaked infectivity) and 11 days ($R_0 = 1.5$, flat infectivity) if applied to infections acquired while a person is at home or traveling. If, however, one assumes that persons infected while traveling would return home, then the delay is less significant, 1–4 days in scenario 1 and 0–1 days in scenario 2.

**(B) Simple Model in Continuous Time**

A simpler deterministic model for the infected persons is used to analyze differences in rates of travel on a single route. If we consider only the early stages of the epidemic, well before the peak, then the number of susceptible persons is approximately equal to the total population. The epidemic equations then reduce to the pair of coupled, linear ordinary differential equations that describe the change in the number of infected persons over time:

\[
\begin{pmatrix}
\dot{I}_1 \\
\dot{I}_2
\end{pmatrix} = \left[ \begin{array}{c}
I_1 \\
I_2
\end{array} \right] = \left[ \begin{array}{c}
-(1 - A_1)^2 + \eta \phi_+ A_1^2 - \gamma / \beta \\
A_1(1 - A_2) + \phi_+ A_2 (1 - A_1) / \eta
\end{array} \right] + \left[ \begin{array}{c}
(1 - A_2) (1 - A_1) + \eta \phi_+ A_1 (1 - A_2) \\
(1 - A_2)^2 + \phi_+ A_2^2 / \eta - \gamma / \beta
\end{array} \right] \left[ \begin{array}{c}
I_1 \\
I_2
\end{array} \right]
\]

where $\beta$ is the effective contact rate, $1 / \gamma$ is the average duration of infection, $\eta$ is the population of city 1 divided by the population in city 2, $A_1$ and $A_2$ are the travel rates from city 1 to city 2 and vice versa, and $\phi_+$ and $\phi_-$ are population modifiers due to travel, with definitions

\[
\phi_+ = 1 / (1 - A_2 + \eta A_1), \ \phi_- = 1 / (1 - A_1 + A_2 / \eta).
\]

Symbolically, the solution to these equations can be written as

\[
\begin{pmatrix}
I_1(t) \\
I_2(t)
\end{pmatrix} = \exp(\beta \hat{A} t) \begin{pmatrix}
I_1(0) \\
I_2(0)
\end{pmatrix}
\]

where

\[
\hat{A} = \begin{pmatrix}
(1 - A_1)^2 + \eta \phi_+ A_1^2 - \gamma / \beta \\
A_1(1 - A_2) + \phi_+ A_2 (1 - A_1) / \eta
\end{pmatrix} + \begin{pmatrix}
(1 - A_2) (1 - A_1) + \eta \phi_+ A_1 (1 - A_2) \\
(1 - A_2)^2 + \phi_+ A_2^2 / \eta - \gamma / \beta
\end{pmatrix}.
\]

When susceptible persons are prevented from traveling, this matrix simplifies to the form

\[
\hat{A}_S = \begin{pmatrix}
1 - A_1 - \gamma / \beta \\
A_1
\end{pmatrix} \begin{pmatrix}
A_2 \\
1 - A_2 - \gamma / \beta
\end{pmatrix}.
\]
while in the absence of travel by infected persons, it can be expressed as

\[
\hat{A}_f = \begin{pmatrix}
1 - \frac{A_1 - \gamma / \beta}{A_2 \phi_2 / \eta}
\end{pmatrix} \begin{pmatrix}
A_1 \eta \phi_2 \\
1 - A_2 - \gamma / \beta
\end{pmatrix}.
\] (9)

Although the equations for the time to spread can be written in analytic form, a numerical solution is required, and hence the MATLAB (The MathWorks, Natick, MA, USA) function \textit{fsolve} is used to compute the solutions graphed in the Appendix Figure panel A.

(C) Dependence on Origin of Travelers and City Size

The dependence on these factors can be studied more generally in terms of differences in population size and travel rates by using the above deterministic version of the stochastic model used in simulations. In this context, \( T_{20} \), the time between the days on which the number of infected persons first reached 20 in city 1 and first reached 20 in city 2, is no longer random. In Figure panel A, the effect on \( T_{20} \) of varying the city 1/city 2 population ratio (\( \eta \)) is graphed (assumptions are that the city 1/city2 travel rates are equal, with \( R_0 = 1.5 \), a 6-day infection period, and the epidemic beginning in city 1). The solid curves show that the delay in epidemic spread increases noticeably as the size of city 1 increases in comparison to city 2. Figure panel B explores the effect of differences in the ratio of city 1/city 2 travel rates for 3 different city population ratios (\( \eta = 1, 1/10, \) and \( 10 \)) and shows that this factor can also strongly influence \( T_{20} \).

A better understanding of the effect of city size ratio on \( T_{20} \) is obtained by “switching off” the travel of infected persons or, alternatively, susceptible persons in the model. Switching off travel by susceptible persons implies that disease can only spread through the movement of infected persons from city 1 to city 2. Since the ratio, \( \eta \), does not alter the average number of infections per infectious person (\( R_0 \)), it should not influence the delay in intercity spread if susceptible persons do not travel. The dashed line in Figure panel A confirms this prediction. If, however, only susceptible persons can initiate journeys, then the disease can only move from city 1 to city 2 through susceptible persons from city 2 acquiring the infection in city 1 before returning to city 2. The ratio of city sizes ought to be important in this instance because the risk of acquiring infection from a single infected person is inversely proportional to the size of the population. This effect is apparent from the dotted line in Figure panel A, with \( T_{20} \) increasing linearly with \( \log(\eta) \) for all but very small values of the variable.
(D) Analytic Approximations

The effect of travel restrictions can be approximated in terms of the initial growth rate, \( r \), of the epidemic as \( T_p = 0.99 = \log(100)/r \), for 99% restrictions and similarly for other levels of travel restriction. The time taken to reach 20 cases can be used to estimate \( r \), and in fact, we can re-express \( T_p \) by using this time: \( T_p = 0.99 = T_{20} \times \log(100)/\log(20) \). In practice, estimates based on \( T_{20} \) can be somewhat inaccurate for low values of \( R_0 \) (Figure panel D).

The probability of an outbreak spreading from city 1 to city 2, as described by the simulation model above, can be approximated by using extinction probabilities for branching processes (1). If, over the course of an outbreak, an attack rate, \( AR \), occurs in the source region, then with \( N \) visitors in town A per day and \( N \) citizens of town A visiting the source region, the probability of an outbreak occurring is

\[
p_A \approx 1 - \exp(-2N(1-q)AR R_0),
\]

where \( q \) is the extinction probability for a Poisson branching process with mean \( R_0 \). As is seen in Figure panel C, where \( AR \approx (1-q) \), this approximates the simulation model very well.

Reference

1. Grimmett GR, Stirzaker DR. Probability and random processes, 3rd ed. Oxford (UK): Oxford University Press; 2001.
Appendix Figure. A) $T_{20}$ (the time between the days on which the number of infected persons first reached 20 in city 1 and first reached 20 in city 2) (now deterministic) for reproduction number $R_0 = 1.5$ and 6-day duration of infection, when the city size ratio, $\eta = N_1/N_2$, is varied. The epidemic is assumed to begin with 1 infectious person in city 1. Travel rates are equal ($A_1 = A_2$) with $A_1 = 1/1,000$; solid, dotted, and dashed lines correspond to persons being free to travel, independent of disease state, only susceptible and removed persons traveling, and only infected persons traveling, respectively. B) Persons are free to travel regardless of disease state, with $A_1 = 1/1,000$, but the reverse travel rate is varied. Solid, dotted, and dashed curves correspond to $\eta = 1$, 10, and 1/10, respectively. C) Probability of the outbreak spreading to city 2 as travel volume increases, with markers indicating results from simulations and curves from the analytic approximation. D) the analytic formula for effect of travel restrictions is compared with simulations of scenario 1 (an outbreak beginning in Sydney and spreading to Melbourne) by using the flat infectivity function. Horizontal axes are on a log-scale, and the legend in panel C also refers to panel D.