Effectiveness of travel restrictions in the rapid containment of human influenza: a systematic review
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Objective To assess the effectiveness of internal and international travel restrictions in the rapid containment of influenza.
Methods We conducted a systematic review according to the requirements of the Preferred Reporting Items for Systematic Reviews and Meta-Analyses statement. Health-care databases and grey literature were searched and screened for records published before May 2014. Data extraction and assessments of risk of bias were undertaken by two researchers independently. Results were synthesized in a narrative form.
Findings The overall risk of bias in the 23 included studies was low to moderate. Internal travel restrictions and international border restrictions delayed the spread of influenza epidemics by one week and two months, respectively. International travel restrictions delayed the spread and peak of epidemics by periods varying between a few days and four months. Travel restrictions reduced the incidence of new cases by less than 3%. Impact was reduced when restrictions were implemented more than six weeks after the notification of epidemics or when the level of transmissibility was high. Travel restrictions would have minimal impact in urban centres with dense populations and travel networks. We found no evidence that travel restrictions would contain influenza within a defined geographical area.
Conclusion Extensive travel restrictions may delay the dissemination of influenza but cannot prevent it. The evidence does not support travel restrictions as an isolated intervention for the rapid containment of influenza. Travel restrictions would make an extremely limited contribution to any policy for rapid containment of influenza at source during the first emergence of a pandemic virus.

Introduction
Travel restrictions were included in the WHO interim protocol: rapid operations to contain the initial emergence of pandemic influenza that was published in 2007 by the World Health Organization (WHO). 1 However, as they would hamper global travel and trade, such restrictions are not recommended by WHO once the global spread of pandemic influenza is established. 2,3 In 2009, some countries applied travel restrictions as one of several strategies to prevent the introduction of the influenza virus A(H1N1)pdm09 into their territories but the effectiveness of this approach has subsequently been questioned. 4 Research on influenza has focused on the evaluation of the effectiveness and impact of pharmaceutical interventions. 5 As quantitative assessment of the effectiveness of travel restrictions in pandemic situations tends to be more challenging, there are scarce data on this topic. In any meta-analysis of surveillance data from multiple studies, it is difficult to quantify and compare the effectiveness of travel restrictions because such interventions are frequently implemented with other countermeasures and without following standardized protocols. 6 However, mathematical models can be used to predict the effectiveness of each type of intervention and inform policy-makers at national and international levels. In 2009, a systematic review of studies based on such models revealed limited evidence of the effectiveness of restrictions in air travel – within and between countries – in the containment of pandemic influenza. 2 There has been no more recent systematic assessment of the effectiveness of restrictions in land, sea or air travel as isolated interventions. We therefore decided to assess the effectiveness of travel restrictions in the rapid containment of influenza strains with pandemic potential, in a systematic review that incorporated data collected during the 2009 pandemic.

Methods
Before commencement, our protocol was registered with PROSPERO – the international prospective register of scientific reviews maintained by the United Kingdom of Great Britain and Northern Ireland’s National Institute for Health Research. 4 We conducted a systematic review according to the requirements of the Preferred Reporting Items for Systematic Reviews and Meta-Analyses statement. 7 We assessed the evidence for restrictions in internal travel – travel within the same country – or international travel – travel between two or more countries – affecting the spread of influenza. We considered the air, terrestrial or maritime transportation of humans to or within countries affected by seasonal or pandemic influenza. The outcome measures of interest were epidemiological characteristics and some viral transmission parameters of influenza such as the basic reproductive number (R0). Studies eligible for inclusion were reports, reviews, meta-analyses, mathematical modelling studies and observational and experimental studies published before May 2014. Studies that only evaluated the spread of influenza in animals or animal products were excluded.

Search strategy
We searched numerous health-care databases and sources of grey literature (Box 1). Critical keywords and thesaurus heading terms were initially tailored to MEDLINE searches and then adapted for other sources as necessary. The full search

Abstracts in العربية, 中文, Français, Русский and Español at the end of each article.

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Box 1. Sources of literature included in this systematic review

**Health-care databases**
- CINAHL (Cumulative Index to Nursing and Allied Health Literature)
- Cochrane Library – Central Register of Controlled Trials
- EMBASE
- PubMed – including MEDLINE
- World Health Organization Global Index Medicus

**Evidence-based reviews**
- Bandolier
- Cochrane Library – Cochrane Database of Systematic Reviews, Database of Abstracts of Reviews of Effects, Health Technology Assessment Database, NHS Economic Evaluation Database

**Guidelines**
- United Kingdom Department of Health
- United Kingdom National Institute for Health Care and Excellence – Evidence Search
- United States Centers for Disease Control and Prevention – Guidance

**Grey literature**
- Consultation with domain experts – Martin Cetron (Centers for Disease Control and Prevention, Atlanta), John Edmunds (London School of Hygiene & Tropical Medicine, London), Peter Grove (Department of Health, London), Richard J Pitman (Oxford Outcomes, Oxford)
- OpenSIGLE system for information on grey literature in Europe
- United Kingdom National Institute for Health Care and Excellence – Evidence Search
- Web of Science

**Manual searching of relevant journals**
- Eurosurveillance
- Emerging Infectious Diseases

**Reference tracking**
- Reference lists of all studies selected for inclusion were searched to identify further relevant studies

**Citation tracking**
- Web of Science – Science Citation Index
- Google Scholar

**Internet searching**
- www.google.com
- www.dh.gov.uk
- www.hpa.org.uk – now: www.phe.gov
- www.who.int
- www.cdc.gov
- www.flu.gov

construct was included in the registered protocol. We contacted field experts and undertook reference and citation tracking to identify further relevant literature.

**Study selection**

All records identified were imported into the EndNote X6 software package (Thomson Reuters, San Francisco, United States of America). Following the removal of duplicates, all remaining records were screened for inclusion against the protocol’s eligibility criteria by two researchers. We used a threestage sifting approach to review titles, abstracts and full texts. Where disagreements arose, a third reviewer provided arbitration.

**Data extraction**

All records that met the eligibility criteria were subject to data extraction. Two reviewers independently extracted study data using a piloted form; any disagreements were resolved with a third reviewer. The full list of data items extracted is available on PROSPERO.

**Assessing risk of bias**

Risk of bias was assessed at both study and outcome level. We used an evaluation tool developed by the United States Agency for Healthcare Research and Quality for assessing such risk in reviews. Since we are not aware of a previously validated instrument to assess risk of bias in mathematical modelling studies, we developed a tool based on the principles for the construction of mathematical models recommended by the London School of Hygiene & Tropical Medicine, in consultation with an experienced modeller (see Appendix A; available at: http://www.nottingham.ac.uk/research/groups/healthprotection/documents/supporting-data/)

**Summary measures and data synthesis**

Descriptive statistics were calculated using Excel 2010 (Microsoft, Richmond, USA). We used a recognized framework to synthesize the extracted data and assessments of risk of bias in a narrative style.

**Results**

**Study selection and characteristics**

Before removal of duplicates, we identified 8836 potentially relevant records. However, only 23 studies – 19 mathematical modelling studies, one time-series analysis, two literature reviews and one systematic review – met our eligibility criteria (Fig. 1).

Of the modelling studies included, 14 used stochastic models, two used deterministic models, two used a combination of both stochastic and deterministic methods and one used a Poisson regression model. Six studies were based on meta-population models of influenza spread and one on an alternative model. The focus of the included studies was the effectiveness of internal and international travel restrictions or combined internal and international travel restrictions.

All but three of our included studies involved assessments of the impact of restrictions on air travel.
one assessed the impact of restrictions on aerial, maritime and terrestrial transportation. The characteristics of the included modelling studies and time-series analysis are presented in Appendix A.

The systematic review that we included synthesized evidence from modelling studies published between 1990 and September 2009. The literature reviews that we included evaluated evidence from mathematical modelling studies on the containment of pandemic influenza and evidence used for preparedness planning in the United Kingdom.  

Risk of bias within studies

Of the 20 studies based on mathematical modelling or time-series analysis, 17 were found to be at low risk of bias (Table 1). The other three were found to be at moderate risk of bias – because of limitations in the study design or the low quality of travel data. Methodological issues that may have led to bias included a lack of transmission variation during the progression of epidemics, seasonality, heterogeneous mixing and varying susceptibility of populations.  

The systematic review and literature reviews were at moderate risk of bias (Table 2). The systematic review was based on literature from only one health-care database and on a snow-balling strategy that could have introduced selection bias. Neither of the literature reviews included any assessment of the design and quality of the studies that were included or detailed descriptions of the eligibility criteria applied.  

Synthesis of results

Internal travel restrictions

Travel restrictions appeared to have limited effectiveness in the containment of influenza at local level (Table 3 and Table 4); Table 3 is available at: http://www.who.int/bulletin/volumes/92/12/14-135590.  

With pandemic influenza A(H1N1)pdm09 in Mongolia, the estimated delay of the pandemic peak varied between 1.0 and 1.5 weeks when 50% road and rail travel restrictions over 2–4 weeks were simulated. The corresponding impact on the attack rate was minimal – e.g. 95% travel restrictions led to a reduction of just 0.1%. A study set in the USA revealed similar findings – e.g. a delay in spread of 2–3 weeks if travel restrictions were 99% effective and implemented in conjunction with border restrictions that prevented the entry of infected travellers. Travel restrictions alone could delay spread by 1 week but only if implemented within 2 weeks of the first case. In one simulation, border controls preventing 99.9% of cases entering any given country delayed epidemic spread by up to 35 days. Another study in the USA presented analogous results – e.g. a 90% restriction on long-distance flights led to delays in the epidemic peak that ranged between a few days and a few weeks. Effectiveness of travel restrictions decreased as the transmissibility of the strain increased; travel restrictions reduced the incidence of new cases by less than 3%. According to a time-series analysis in the USA, a 50% restriction in air travel during the 2001–2002 influenza season would have delayed the peak mortality associated with novel strains of seasonal influenza by 16 days – i.e. compared with the timing of the peak in previous years.  

Internal travel restrictions in England, Scotland and Wales in the United Kingdom were predicted to have minimal impact on the magnitude of the peak and in delaying the spread of the epidemic – possibly because there are some densely populated urban areas and relatively high levels of population movement. However, in a recent review, it was estimated that a combination of internal and international travel
### Table 1. Risk of bias assessments of mathematical modelling studies or time-series analysis on the effectiveness of travel restrictions to reduce influenza transmission

| Study | Research question(s) precise and clear | Primary findings presented | Original findings | Model techniques or model structure used | Appropriate model complexity | Suitable mathematical modelling | Input data sources identified | Major model assumptions described | Relevant factors explored | Model validated | Techniques used for model fitting | Sensitivity analysis |
|-------|---------------------------------------|-----------------------------|-------------------|------------------------------------------|-----------------------------|-------------------------------|-----------------------------|---------------------------------|-----------------------------|----------------|----------------------------------|---------------------|
| Bajardi et al. (2011) | Low | Low | Low | Low | Low | Low | Low | Low | Low | Low | Moderate | Low |
| Bolton et al. (2012) | Low | Low | Low | Low | Low | Low | Low | Low | Low | Low | Moderate | Low |
| Brownstein et al. (2006) | Low | Low | Low | Low | Low | Low | Low | Low | Low | Low | Low | Moderate |
| Chong and Ying Zee (2012) | Low | Low | Low | Low | Low | Low | Low | Low | Low | Low | Moderate | Low |
| Ciofi degli Atti et al. (2008) | Low | Low | Low | Low | Low | Low | Moderate | Low | Low | Low | Moderate | Low |
| Colizza et al. (2007) | Low | Low | Low | Low | Low | Low | Low | Low | NS | Low | Low | Moderate |
| Cooper et al. (2006) | Low | Low | Low | Low | Low | Low | Low | Low | Low | Low | Low | Moderate |
| Eichner et al. (2009) | Low | Low | Low | Moderate | Low | Moderate | Low | Low | NS | N | Moderate | Low |
| Epstein et al. (2007) | Low | Low | Moderate | Low | Low | Low | Low | Low | Low | N | N | Low |
| Ferguson et al. (2006) | Low | Low | Low | Low | Low | Low | Low | Low | Low | Low | High | Low |
| Flahault et al. (2006) | Low | Low | Low | Low | Low | Moderate | Low | Moderate | Low | Low | N | Moderate |
| Germann et al. (2006) | Low | Low | Low | Low | Low | Low | Low | Low | Low | Low | High | N |
| Hsieh et al. (2007) | Low | Low | Low | Moderate | Low | Moderate | Low | Low | Low | N | N | High |
| Hollingsworth et al. (2006) | Low | Low | Moderate | Low | Low | Moderate | Low | Moderate | Low | N | N | High |
| Kernéis et al. (2008) | Low | Low | Low | Low | Low | Low | Low | Low | Low | Low | High | Low |
| Lam et al. (2011) | Low | Low | Low | Low | Low | Moderate | Low | Moderate | Low | Low | No | Low |
| Lee et al. (2012) | Low | Low | Low | Low | Low | Low | Low | Low | Low | Low | No | Low |
| Marcelino & Kaiser (2012) | Low | Low | Low | Low | Low | Moderate | Low | Moderate | Low | Low | N | Low |
| Scalia Tomba & Wallinga (2008) | Low | Low | Low | Moderate | Moderate | Moderate | Low | Moderate | Low | Low | N | High |
| Wood et al. (2007) | Low | Low | Low | Low | Low | Low | Low | Low | Low | Low | N | N |

NS: not specified.

* For each domain of interest, risk of bias was categorized as low if the authors addressed the domain adequately, moderate if the authors’ coverage of the domain was superficial or incomplete, and high if the authors reported coverage of the domain was poor.

* As this study contained mainly modelling components relevant to the outcomes, it was assessed for risk of bias as a modelling study.
Table 2. Risk of bias assessments of systematic or literature reviews on the effectiveness of travel restrictions to reduce influenza transmission

| Study | Search and evaluation criteria | Data extraction | Study quality and validity criteria | Outcomes | Intervention(s) | Incidence and exclusion criteria | Funding or sponsorship |
|-------|--------------------------------|-----------------|------------------------------------|----------|-----------------|-------------------------------|-----------------------|
| Department of Health (2011)20 | Low | Low | Moderate | Low | Low | High | Moderate | Low | Low | NS |
| Department of Health (2012)21 | Low | High | Moderate | Low | Low | High | High | Low | Low | Low |
| Lee et al. (2009)7 | Low | Low | Low | Low | Low | Low | Moderate | Low | Low | Low |

NS: not specified; UKDH: United Kingdom of Great Britain and Northern Ireland Department of Health.

For each domain of interest, risk of bias was categorized as low if the authors addressed the domain adequately, moderate if the authors’ coverage of the domain was superficial or incomplete, and high if the authors reported coverage of the domain was poor.

International travel restrictions

International travel restrictions also appeared to have limited effectiveness (Table 5 and Table 6). Low-level restrictions – i.e. restrictions of less than 70% – were the least effective in containing the spread of epidemics between countries. It was found that a 40% restriction of air travel would only delay the spread of influenza A(H1N1)pdm09 from Mexico to other countries by less than 3 days.1 In a high transmissibility scenario, a 20% or even a 50% reduction in the volume of travellers would not have any significant impact on the global spread of influenza A(H5N1).3 In a meta-population model of pandemic influenza, based on the 1968–1969 influenza A(H3N2) pandemic virus it was predicted delays in the epidemic peak of 9 and 14 days with 50% and 90% restriction of air travel, respectively.18

In Italy, relatively large delays were reported in reaching an influenza A(H5N1) peak – i.e. 7–37 days, depending on the level of influenza transmissibility and the extent of the restrictions simulated.17 Travel restrictions had no beneficial effect on attack rate if the level of strain transmissibility was moderate or high.17

In a more recent review, it was estimated that introduction of pandemic influenza into the United Kingdom could be delayed by up to 2 months if there was an almost complete – e.g. 99.9% – ban on air travel.20 However, the size of the effect was considerably reduced, to just 1–2 weeks, if the level of restriction was lowered to 90%.20 Similar observations were made in an assessment of the impact of restrictions of air, land and sea travel on the introduction of H1N1 pdm09 into Hong Kong Special Administrative Region (SAR), China.24 In this study, it was estimated that restrictions of 90% and 99% on all modes of transportation would delay the epidemic peak by up to 6 and 12 weeks, respectively, when $R_0$ was set to 1.4.45 When $R_0$ was set to 1.7, a restriction of 99% on all modes of transportation would delay the epidemic peak by up to 8 weeks and halve the cumulative attack rate. Air travel restrictions appeared to be the most effective isolated intervention, even though most infected cases would probably enter Hong Kong SAR by land travel from mainland China.45 Although one review of the evidence from mathematical modelling concluded that air travel bans would probably have a similar effect irrespective of the pandemic’s country of origin,21 another report believed that the effectiveness of such restrictions would vary according to the geographical source of the pandemic.21 If air travel bans delayed the epidemic so that it coincided with the usual influenza season, the apparent number of cases and the size of the peak in the epidemic could both increase.31 However, the opposite trends might be observed if the travel restrictions coincided with a period of low strain transmissibility.31 By restricting air travel by 95%, it should be possible to delay pandemic spread across the USA – of an infection originating in Sydney or Hong Kong SAR – by 2–3 weeks.21 However, there was no corresponding impact if the geographical origin of the pandemic was London because of London’s high flight densities and interoperability.11 The selective cancellation of a quarter of all connection flights between 300 major cities worldwide could be more effective than the closure of all the cities’ airports – reducing the number of infected travellers by an additional 19%.32 A review of air travel restrictions between Asia and the United Kingdom
Travel restrictions for the containment of influenza

| Study | Type of restrictions | Study design | Influenza strain involved | Strain transmissibility ($R_0$) | Scenario and duration of intervention | Effect estimate |
|-------|----------------------|--------------|---------------------------|---------------------------------|--------------------------------------|-----------------|
| Bolton et al. (2012) | Internal travel restrictions – i.e. blanket or reactive movement restrictions | Mathematical stochastic model | Pandemic influenza A H1N1 pdm09 | 1.6 | 95% travel restriction, 2–4 weeks | 12% reduction in ILI peak and a reduction in mean attack rate of <0.1%, even when restrictions with 95% effectiveness are implemented for 4 weeks |
| Ferguson et al. (2006) | Internal air, plus border controls, England, Scotland, and Wales in United Kingdom and USA | Mathematical stochastic model | Novel pandemic influenza strain | 1.4–2.0 | Internal travel restrictions – i.e. blanket or reactive movement restrictions – a 90–100% level of effectiveness | Reduction in attack rate of <0.1%, even when restrictions with 95% effectiveness are implemented for 4 weeks |
| Germann et al. (2006) | Internal, USA | Stochastic single-city and multi-city extended models | H5N1 pandemic influenza | 1.6, 1.9, 2.1, or 2.4 | 90% reduction in long-distance domestic travel when $R_0$ is set to 1.6, 1.9, 2.1, or 2.4, cumulative incidence per 100 inhabitants has been recorded in USA, 80 days | Decreased $R_0$ to <1, preventing spread of epidemic |
| Hsieh et al. (2007) | Internal, China | Mathematical stochastic patch model | Human seasonal influenza | NS | NS | Travel of symptomatic individuals from areas of high prevalence to areas of low prevalence eliminated |

Discussion

The results of our systematic review indicate that overall travel restrictions have only limited effectiveness in the prevention of influenza spread, particularly in those high transmissibility scenarios in which $R_0$ is at least 1.9 (Box 2). The effect size varied according to the extent and timeliness of the restrictions, the size of the epidemic, strain transmissibility, the heterogeneity of the travel patterns, the geographical source and the urban density of international travel hubs. Only extensive travel restrictions – i.e. over 90% – had any meaningful effect on reducing the magnitude of epidemics. In isolation, travel restrictions might delay the spread and peak of pandemics by a few weeks or months but we found no evidence that they would contain influenza within a defined geographical area.

Several limitations associated with our review warrant discussion. We included mathematical modelling studies that simulated very diverse scenarios with varying levels of $R_0$, geographical locations, means of transportation, strains and population characteristics. A paucity of surveillance data concerning the impact and effectiveness of nonpharmaceutical interventions meant that our observations had to be mainly based on simulations. While mathematical models are important tools that can be used to inform policy-makers, they cannot account fully for all aspects of real-life situations.
### Table 5. Simulated effects of the implementation of international travel restrictions on the spread and duration of pandemic or epidemic influenza

| Study | Type of restrictions and setting | Study design | Influenza strain involved | Strain transmissibility ($R_0$) | Scenario and duration of intervention                                                                 | Effect estimate |
|-------|----------------------------------|--------------|---------------------------|---------------------------------|-----------------------------------------------------------------------------------------------------|-----------------|
| Bajardi et al. (2011) | Air travel, global | Mathematical stochastic model | A(H1N1)pdm09 epidemic | NS | 40% restriction, < 6 weeks from epidemic notification; 90% restriction, < 6 weeks from epidemic notification; Any level of restriction, > 6 weeks from epidemic notification | ES to other countries delayed < 3 days; ES to other countries delayed < 2 weeks; No impact |
| Brownstein et al. (2006) | Internal and international air travel, USA | Time-series analysis | Seasonal influenza | 1.4, 1.7 or 2.0 | Travel restricted to and from a city with > 1000 infectious cases or worldwide when > 1000 such cases in city of origin, the 2001–2002 influenza season | Seasonal influenza season prolonged by 16 days |
| Chong and Ying Zee (2012) | Air, sea and land travel, Hong Kong Special Administrative Region, China | Mathematical stochastic model | A(H1N1)pdm09 | 1.1 | 99% air, land and sea travel; 90% air, land and sea; 99% air and land; 99% sea | EP delayed up to 1 year; ES delayed 4 and 6 weeks, respectively; ES delayed 2 and 3 months, respectively; ES delayed 1–2 and 3.5 weeks, respectively; EP delayed up to 2 weeks; EP delayed up to 1 week; EP delayed up to 1 week; No significant impact on timing of EP; EP delayed up to 8 weeks |
| Ciofi degli Atti et al. (2008) | Air travel, Italy | Mathematical global deterministic model | A(H5N1) | 1.9 | 90% air travel restriction, implemented 30 days after first case in pandemic was recorded or < 2 months after the introduction of first case in Italy; As above except 99% restriction | With $R_0$ set to 1.4, 1.7 and 2.0, EP delayed, median of 23, 10 and 6 days, respectively |
| Colizza et al. (2007) | Air travel, global | Mathematical stochastic metapopulation compartmental | A(H5N1) | 1.8 | 20% or 50% air traveller reduction at each connection | No significant impact on EP |
| Cooper et al. (2006) | Air travel, global | Mathematical stochastic metapopulation model | Epidemic and pandemic influenza | 3 | 100% susceptible, 50% air travel reduction, after first 100 symptomatic cases in each city or after 1000 cases in city of origin | EP delayed median of 7 days |

(continues...)

| Study | Type of restrictions and setting | Study design | Influenza strain involved | Strain transmissibility ($R_0$) | Scenario and duration of intervention | Effect estimate |
|-------|---------------------------------|-------------|--------------------------|-----------------|---------------------------------|----------------|
| Department of Health (2011) | Evidence-based review | Literature review | Pandemic influenza | NS | 90% air travel restriction | ES delayed 1–2 weeks |
| Department of Health (2012) | Modelling summary | Literature review | Pandemic influenza | NS | 90% air travel restriction | ES delayed 2 months |
| Eichner et al. (2009) | Air and sea travel, Pacific islands | Mathematical model[^a] | A(H1N1)pdm09, 1.5, 2.25 or 3.0 | 79% air and sea travel restriction | ES delayed 1–2 weeks |
| Epstein et al. (2007) | Air travel, global | Mathematical model[^b] | Pandemic influenza, 1.7 | Hong Kong Special Administrative Region as source of epidemic, 95% restriction implemented after 1000 infectious cases | ES delayed 13.5 days |
| Ferguson et al. (2006) | Internal air, plus border controls, England, Scotland and Wales in United Kingdom and USA | Stochastic mathematical individual-based model[^c] | Novel pandemic influenza strain, 2.0 | 90% restriction on entry of infected individuals | IOE delayed 9 days in (England, Scotland and Wales in United Kingdom) or 15 days (USA) |
| Flahault et al. (2006) | Air travel, 55 cities worldwide | Mathematical deterministic model[^d] | 1958–1969-like pandemic influenza | NS | 50% travel restriction, at the start of the pandemic or city-by-city, when there is more than one infectious case per 100,000 population | ES delayed 9 days |

[^a]: Pandemic influenza strain
[^b]: Pandemic influenza as the theoretical origin of epidemic – eliminated
[^c]: Pandemic influenza as the theoretical origin of epidemic – eliminated
[^d]: Pandemic influenza as the theoretical origin of epidemic – eliminated
The lack of available data from observational or experimental studies precluded the conduct of the meta-analysis and sensitivity analysis that formed part of the protocol that we registered. Most of the studies that we included in our review used probabilistic models that appeared to have adequate levels of complexity to simulate disease spread and the impact of interventions. In comparison, deterministic models are less complex and do not take uncertainty into account but are still useful when limited data are available and a rapid simulation is needed. Most of the studies we reviewed were limited by a lack of consideration of heterogeneous mixing, socioeconomic status and the relationship between age and immunity. Many also simulated constant strain transmissibility during epidemics – even though transmissibility can vary over time because of seasonal climactic conditions, changes in host susceptibility and the effects of interventions such as social distancing, quarantine and the use of antiviral drugs. The authors of some of the articles noted concerns that may have affected model accuracy, such as issues with the quality of air travel data – e.g. a lack of flight itineraries and the need to use crude estimates of the volume of travellers within and between countries. There was a general paucity of data on land and sea travel, although one of the studies provided comprehensive data on such travel. The tool we developed to assess the risk of bias in the mathematical modelling studies has not been validated and could have produced imprecise estimates.

The results of several studies indicate that, in reducing the global spread of influenza and the overall number of infected individuals, a combination of several different interventions is more effective than any single isolated measure. One study estimated that, when the strains involved have moderate transmissibility, a combination of antiviral prophylaxis, extensive travel restrictions and infant vaccination could reduce the cumulative attack rate by 77–87%. However, effective vaccines are not generally available at the point of emergence of a novel pandemic virus. The effectiveness of combined or single interventions can be affected by the timeliness of the implementation and this appears to be particularly relevant with strains of higher transmissibility.

Often, in the context of pandemic preparedness and response, travel re-
### Box 2. Summary of findings of the 23 studies assessed

**Internal travel restrictions: general observations**
- Have limited effectiveness
- Delay pandemic spread by about 1 week
- Delay pandemic peak by about 1.5 weeks
- Have little impact on magnitude of pandemics – e.g. they may reduce attack rates by < 2%.
- Simulated impact is particularly weak in scenarios that involve strains with high transmissibility

**Internal travel restrictions: risk of bias assessment**
- Relevant studies have low to moderate risk of bias
- Paucity of data on terrestrial travel may have led to an overestimation of the impact of travel restrictions
- Many simulations take no account of the characteristics of human populations – e.g. the mixing and variation of susceptibility across age groups – or of seasonality. Such limitations could well have affected the simulated spread of pandemic waves and impacts of interventions

**International travel restrictions: general observations**
- Have limited effectiveness – e.g. 90% air travel restriction in all affected countries may delay spread of pandemics by 3–4 weeks
- Have minimal impact on the magnitude of pandemics, typically reducing attack rates by less than 0.02%
- May prolong the seasonal influenza season
- May result in higher epidemic peak if resultant delay causes pandemic wave to coincide with seasonal influenza wave
- Simulated impact particularly weak in scenarios that involve strains with high transmissibility
- Extensive restriction of international air travel might delay introduction of a pandemic into a country by up to 2 months and delay pandemic spread by 3–4 months
- Would not prevent introduction of a pandemic into any given country
- May give time for other interventions – e.g. the production and distribution of effective vaccines and antiviral drugs
- Social and economic impacts need to be evaluated

**International travel restrictions: specific measures**
- May have benefits compared with more widespread restrictions – e.g. in one simulation, compared with the closure of all of the cities’ airports, the targeted reduction of a quarter of flight connections between 500 major cities gave a greater reduction in the number of infected travellers
- Compared with banning air travel by adults, the banning of air travel by children may be more effective at delaying the spread of a pandemic but is socially impractical

**International travel restrictions: risk of bias assessment**
- Relevant studies have low to moderate risk of bias
- A paucity of data on travel by sea and land may have led to an overestimation of the impact of air travel restrictions on the containment of influenza pandemics
- Much of the information available on air travel has a lack of detail on flight destinations and numbers of travellers and this may have led to inaccurate assumptions being made about the spread of influenza
- Again, many simulations take no account of the characteristics of human populations – e.g. the mixing and variation of susceptibility across age groups – or of seasonality. Such limitations could well have affected the simulated spread of pandemic waves and impacts of interventions
- When simulating novel pandemic strains, validation of models was an issue; mathematical models need to be validated against surveillance data to improve their value as predictive tools for policy-makers

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**Competing interests**

The University of Nottingham Health Protection and Influenza Research Group is currently in receipt of research funds from GlaxoSmithKline (GSK) and unrestricted educational grants for influenza research from F Hoffmann-La Roche and Astra Zeneca. However, this funding did not support any aspect of the present study. Prior to October 2010, JSNV-T received funding to attend influenza-related meetings and give lectures, and also consultancy fees and research funding from several manufacturers of antiviral drugs and influenza vaccines. JSNV-T was an employee of SmithKline Beecham, Roche Products and Aventis-Pasteur MSD prior to 2005 but now has no outstanding pecuniary interests by way of shareholdings, share options or accrued pension rights.
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目的
评估国内和国际出行限制对快速控制流感的有效性。

方法
我们根据系统回顾和荟萃分析首选报告项目的需
求进行了一项系统回顾。搜索医疗数据库和灰色文献
并筛选在2014年5月前发表的记录。由两位研究者独
立执行数据提取和误差风险评估。以叙事形式综合结
果。

结果
在纳入的23项研究中，整体误差风险为中低等
级。国内出行限制和国境线限制分别将流感流行传播
推迟一个星期和两个月。国际出行限制将流行病传播
和高峰期延迟几天到四个月不等。出行限制减少的新
病例发病率不到3%。流行病通知发布超过六周后或在
传播等级较高时，实施限制措施的影响效果趋于减少。

结论
广泛的出行限制可能会推迟流感的传播，但没有
阻止作用。证据不支持出行限制是一个快速控制流感
的独立干预。对于任何要在大流行性流感病毒刚刚出
现时就从源头快速控制流感的政策来说，出行限制的
作用非常有限。

Résumé
Efficacité des mesures de restriction des déplacements dans le confinement rapide de la grippe humaine: une revue systématique.

Objectif Evaluer l'efficacité des mesures de restriction des déplacements internes et internationaux dans le confinement rapide de la grippe.

Méthodes Nous avons effectué une revue systématique selon les exigences de l'énoncé des items préférables pour rendre compte des revues systématiques ou des méta-analyses (PRISMA). Nous avons effectué des recherches dans les bases de données sur la santé et la littérature grise et nous avons passé au crible les documents publiés avant mai 2014. L'extraction des données et les évaluations du risque de partialité ont été effectuées par deux chercheurs de manière indépendante. Nous avons fait la synthèse des résultats sous forme narrative.

Résultats Le risque global de partialité dans les 23 études incluses était faible à modéré. Les mesures de restrictions des déplacements internes et les mesures de restriction aux frontières internationales n'ont retardé la propagation des épidémies de grippe d'une semaine et de deux mois, respectivement. Les mesures de restriction des déplacements internationaux ont retardé la propagation et le pic de l'épidémie de périodes variant de quelques jours à quatre mois. Les mesures de restriction des déplacements ont réduit de moins de 3% l'incidence des nouveaux cas. L'impact était réduit lorsque des mesures de restriction ont été mises en œuvre plus de six semaines après la notification de l'épidémie ou lorsque le niveau de transmissibilité était élevé. L'impact des mesures de restriction des déplacements serait minime dans les centres urbains où il existe une population dense et des réseaux de transport. Nous n'avons trouvé aucune preuve que les restrictions de déplacement confineraient la grippe dans une zone géographique définie.

Conclusion Les mesures étendues de restriction des déplacements peuvent retarder la propagation de la grippe, mais ne peuvent pas l'empêcher. Les données probantes n'étant pas les restrictions de déplacement en tant qu'intervention isolée pour le confinement rapide de la grippe. Les restrictions de déplacement n'apporteraient qu'une contribution extrêmement limitée à toute politique de confinement rapide de la grippe à la source lors de la première apparition d'un virus pandémique.
Заключение

Ограничения на поездки сокращали число новых случаев менее чем на 3%. Эффект снижался, если меры по ограничению поездок принимались по истечении шести месяцев после уведомления об эпидемии или когда уровень переносимости заболевания был уже высоким. Ограничения на поездки оказывали минимальное влияние в городских центрах с высокой плотностью населения и развитенной сетью пассажирских перевозок. Доказательства того, что ограничения на поездки препятствуют распространению гриппа за пределы определенного географического региона не найдены.

Вывод

Масштабные меры по ограничению поездок могут замедлить распространение гриппа, но не могут предотвратить его. Факты, подтверждающие, что ограничения на поездки, как отдельная мера, предотвращают быстрое распространение гриппа, не найдены. Ограничения на поездки в чрезвычайно малой степени способствуют быстрой локализации гриппа в месте его возникновения при первом проявлении пандемического вируса.

Резюме

Эффективность ограничений на поездки в целях предотвращения быстрого распространения гриппа человека: систематический обзор

Цель

Оценить эффективность ограничений на внутренние и международные поездки в целях предотвращения быстрого распространения гриппа.

Методы

Был проведен систематический обзор в соответствии с рекомендациями о наиболее предпочтительных параметрах отчетности для систематических обзоров и мета-анализа. Поиск и отбор соответствующей информации был осуществлен в медицинских базах данных и неиндексированной литературе, опубликованной до мая 2014 г. Отбор данных и оценка риска систематической ошибки проводились двумя исследователями независимо друг от друга. Результаты были обобщены в форме отчета.

Результаты

Общий риск систематической ошибки в 23 включенных исследованиях был низким или умеренным. Ограничения на внутренние поездки и на пересечение международных границ задерживали распространение эпидемий гриппа на одну неделю и два месяца соответственно. Ограничения на международные поездки задерживали распространение и пик эпидемий на период от нескольких дней до четырех месяцев.

Ограничения на поездки сокращали число новых случаев менее чем на 3%. Эффект снижался, если меры по ограничению поездок принимались по истечении шести месяцев после уведомления об эпидемии или когда уровень переносимости заболевания был уже высоким. Ограничения на поездки оказывали минимальное влияние в городских центрах с высокой плотностью населения и развитенной сетью пассажирских перевозок. Доказательства того, что ограничения на поездки препятствуют распространению гриппа за пределы определенного географического региона не найдены.

Вывод

Масштабные меры по ограничению поездок могут замедлить распространение гриппа, но не могут предотвратить его. Факты, подтверждающие, что ограничения на поездки, как отдельная мера, предотвращают быстрое распространение гриппа, не найдены. Ограничения на поездки в чрезвычайно малой степени способствуют быстрой локализации гриппа в месте его возникновения при первом проявлении пандемического вируса.
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Table 3. Simulated effects of the implementation of internal travel restrictions on the spread and duration of pandemic or epidemic influenza

| Study                          | Type of restrictions and setting                                                                 | Study design                        | Influenza strain involved     | Strain transmissibility ($R_0$) | Scenario and duration interventions                                                                 | Effect estimate                                                                 |
|--------------------------------|--------------------------------------------------------------------------------------------------|-------------------------------------|--------------------------------|---------------------------------|---------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------|
| Bolton et al. (2012)           | Internal road and rail, Mongolia                                                                 | Mathematical stochastic model        | Pandemic influenza A H1N1 pdm09 | 1.6                             | 50% travel restriction, 2 weeks<br>50% travel restriction, 4 weeks                           | Pandemic peak delayed 1 week<br>Pandemic peak delayed 1.5 weeks                         |
| Brownstein et al. (2006)       | Internal and international air, USA                                                               | Time-series analysis                 | Seasonal influenza              | 1.4, 1.7 or 2.0                  | Travel restricted to and from a city with > 1000 infectious cases or worldwide when > 1000 such cases in city of origin, the 2001–2002 influenza season<br>Peak mortality due to influenza delayed 16 days | Little effect on the length of epidemic and size of peak in each local area<br>Increased spread of epidemic and desynchronization of epidemics in local areas<br>ES delayed 2–3 weeks in USA but not delayed in United Kingdom³³²⁵   |
| Department of Health (2012)    | Several scenarios                                                                                 | Literature review (mathematical models) | Pandemic influenza              | NS                              | 90% internal travel restriction between localities<br>90% internal travel restriction between localities plus total ban on international flights | Little effect on the length of epidemic and size of peak in each local area<br>Increased spread of epidemic and desynchronization of epidemics in local areas<br>ES delayed 2–3 weeks in USA but not delayed in United Kingdom³³²⁵   |
| Ferguson et al. (2006)         | Internal air, plus border controls, England, Scotland and Wales in United Kingdom and USA        | Mathematical stochastic model        | Novel pandemic influenza strain  | 1.4–20                          | Internal travel restriction – implemented when 50 cases reported in affected country – plus 99%-effective border restrictions stopping entry of infected travellers – implemented from day 30 of global pandemic<br>Internal travel restriction in USA | ES delayed 1 week in USA but not delayed in United Kingdom³³²⁵   |
| Germann et al. (2006)          | Internal, USA                                                                                     | Mathematical stochastic model        | H5N1 pandemic influenza         | 1.6, 1.9, 2.1 or 2.4             | 90% reduction in long-distance domestic travel when 10 000 symptomatic individuals have been recorded in USA, 180 days | EP delayed by a few days – when $R_0$ is relatively high – to a few weeks               |
| Lee et al. (2012)              | Restrictions on internal migration, restrictions by airplane, car, bus or ship, Republic of Korea | Mathematical stochastic single-city and multi-city extended models | Human influenza                 | 1.0, 1.2, 1.5 or 1.8             | 90% travel restriction, similar parameters all cities, constant infection force<br>90% travel restriction, similar parameters all cities, variation in infection force | Slight – unspecified – delay in EP Size of EP reduced by < 0.01%<br>Unspecified delay in EP Delayed spread of epidemic into new cities but increased risk of localized larger outbreaks |
| Study | Type of restrictions and setting | Study design | Influenza strain involved | Strain transmissibility ($R_0$) | Scenario and duration interventions | Effect estimate |
|-------|---------------------------------|-------------|--------------------------|---------------------|-----------------------------------|----------------|
| Lee et al. (2009) | Several scenarios | Systematic review (deterministic and stochastic models) | Different strains of pandemic influenza | 1.7–2.0 | Internal and international air travel restriction | ES delayed 2–3 weeks if restrictions 99% effective |
| Wood et al. (2007) | Internal, Australia | Mathematical stochastic model | Pandemic influenza | 1.5, 2.5 or 3.5 | 80% restriction of travel from Sydney to Melbourne, variable infectivity, 2 weeks after epidemic | With $R_0$ set to 1.5, ES delayed a median of 32 days |
| | | | | | As above except constant infectivity | With $R_0$ set to 1.5, 2.5 and 3.5, ES delayed a median of 30, 22 and 16 days, respectively |
| | | | | | As above except peak infectivity | With $R_0$ set to 1.5, 2.5 and 3.5, ES delayed a median of 22, 15 and 11 days, respectively |
| | | | | | 80% restriction of travel from Darwin to Sydney, constant infectivity, 2 weeks after epidemic | With $R_0$ set to 1.5, 2.5 and 3.5, ES delayed a median of 34, 17 and 13 days, respectively |
| | | | | | As above except peak infectivity | With $R_0$ set to 1.5, 2.5 and 3.5, ES delayed a median of 24, 12 and 9 days, respectively |
| | | | | | 80% travel restriction nationwide, 4 weeks after epidemic began | No impact with $R_0$ set to 1.5 |
| | | | | | 90% restriction of travel from Sydney to Melbourne, constant infectivity, 2 weeks after epidemic began | With $R_0$ set to 1.5, 2.5 and 3.5, ES delayed a median of 53, 25 and 18 days, respectively |
| | | | | | As above except peak infectivity | With $R_0$ set to 1.5, 2.5 and 3.5, ES delayed a median of 32, 17 and 13 days, respectively |
| | | | | | 90% restriction of travel from Darwin to Sydney, constant infectivity, 2 weeks after epidemic began | With $R_0$ set to 1.5, 2.5 and 3.5, ES delayed a median of 41, 20 and 15 days, respectively |
| | | | | | As above except peak infectivity | With $R_0$ set to 1.5, 2.5 and 3.5, ES delayed a median of 25, 14 and 10 days, respectively |
| | | | | | 99% restriction of travel from Sydney to Melbourne, constant infectivity, 2 weeks after epidemic began | With $R_0$ set to 1.5, 2.5 and 3.5, ES delayed a median of 75, 34 and 25 days, respectively |
| | | | | | As above except peak infectivity | With $R_0$ set to 1.5, 2.5 and 3.5, ES delayed a median of 52, 24 and 17 days, respectively |
| | | | | | 99% restriction of travel from Darwin to Sydney, constant infectivity, 2 weeks after epidemic began | With $R_0$ set to 1.5, 2.5 and 3.5, ES delayed a median of 75, 30 and 22 days, respectively |
| | | | | | As above except peak infectivity | With $R_0$ set to 1.5, 2.5 and 3.5, ES delayed a median of 46, 21 and 15 days, respectively |

EP: epidemic peak; ES: epidemic spread; NS: not specified; $R_0$: basic reproductive number.

1 A so-called SEIAR model, in which individuals who are susceptible (S), exposed (E), infectious and presented for medical care (I), infectious but not presented for medical care (A) or recovered (R) are considered.
2 A so-called SEIR model, in which individuals who are susceptible (S), exposed (E), infectious (I) or recovered (R) are considered.
3 Internal travel restrictions only effective if implemented within 2 weeks of first case in the USA. Border controls only effective if they prevent entrance of ≥99% of infective travellers and are implemented within 45 days of the start of pandemic.
4 Internal travel restrictions only effective if implemented within 2 weeks of first case in the USA.
5 With reactive movement restrictions, a 20-km exclusion zone is established around every diagnosed case – with merging of overlapping zones – and movement in and out of each exclusion zone is eliminated. With blanket movement restrictions, all journeys by an individual from that individual’s home that exceed a certain distance – often 20 or 50 km – are eliminated.
6 A so-called SIR model, in which individuals who are susceptible (S), infected (I) or recovered (R) are considered.
| Study                      | Type of restrictions and setting | Study design                                      | Influenza strain involved | Strain transmissibility ($R_0$) | Scenario and duration of intervention | Effect estimate |
|---------------------------|---------------------------------|--------------------------------------------------|---------------------------|----------------------------------|----------------------------------------|-----------------|
| Chong and Ying Zee (2012) | Air, land and sea, Hong Kong Special Administrative Region | Mathematical stochastic model$^a$ | A(H1N1)pdm2009 | 1.1, 1.4 or 1.7 | 90% air travel restriction | With $R_0$ set to 1.1, 1.4 and 1.7, CINC was 18%, 50% and 72% of NIV, respectively |
|                           |                                 |                                                  |                           |                                  | 99% air travel restriction              | With $R_0$ set to 1.1, 1.4 and 1.7, CINC was 18%, 49% and 72% of NIV, respectively |
|                           |                                 |                                                  |                           |                                  | 90% sea travel restriction             | With $R_0$ set to 1.1, 1.4 and 1.7, CINC was 15%, 55% and 73% of NIV, respectively |
|                           |                                 |                                                  |                           |                                  | 99% sea travel restriction             | With $R_0$ set to 1.1, 1.4 and 1.7, CINC was 13%, 54% and 73% of NIV, respectively |
|                           |                                 |                                                  |                           |                                  | 90% land travel restriction           | With $R_0$ set to 1.1, 1.4 and 1.7, CINC was 8%, 51% and 71% of NIV, respectively |
|                           |                                 |                                                  |                           |                                  | 99% land travel restriction           | With $R_0$ set to 1.1, 1.4 and 1.7, CINC was 5%, 46% and 71% of NIV, respectively |
|                           |                                 |                                                  |                           |                                  | 90% air and sea travel restriction    | With $R_0$ set to 1.1, 1.4 and 1.7, CINC was 18%, 48% and 70% of non-intervention value, respectively |
|                           |                                 |                                                  |                           |                                  | 99% air and sea travel restriction    | With $R_0$ set to 1.1, 1.4 and 1.7, CINC was 16%, 45% and 70% of NIV, respectively |
|                           |                                 |                                                  |                           |                                  | 90% air and land travel restriction   | With $R_0$ set to 1.1, 1.4 and 1.7, CINC was 15%, 40% and 71% of NIV, respectively |
|                           |                                 |                                                  |                           |                                  | 99% air and land travel restriction   | With $R_0$ set to 1.1, 1.4 and 1.7, CINC was 5%, 35% and 70% of NIV, respectively |
|                           |                                 |                                                  |                           |                                  | 90% land and sea travel restriction   | With $R_0$ set to 1.1, 1.4 and 1.7, CINC was 15%, 50% and 72% of NIV, respectively |
|                           |                                 |                                                  |                           |                                  | 99% land and sea travel restriction   | With $R_0$ set to 1.1, 1.4 and 1.7, CINC was 13%, 48% and 72% of NIV, respectively |
| Ciof the Atti et al. (2008) | Air travel, Italy                | Mathematical deterministic metapopulation$^b$ and individual-based model | NS                         | 1.4, 1.7 or 2.0 | 90% air travel restriction, implemented from 30 days after record of first case for the whole pandemic until 2 months after introduction of first case in Italy As above except 99% air travel restriction | With $R_0$ set to 1.4, 1.7 and 2.0, CAR was 21.2%, 30.8% and 38.7% of NIV and PDAR was 0.42%, 1.01% and 1.90% of NIV, respectively |
|                           |                                 |                                                  |                           |                                  | 99% air travel restriction, implemented from 30 days after record of first case for the whole pandemic until 2 months after introduction of first case in Italy As above except 99% air travel restriction | With $R_0$ set to 1.4, 1.7 and 2.0, CAR was 21.1%, 30.8% and 38.7% of NIV and PDAR was 0.40%, 1.03% and 1.91% of NIV, respectively |
| Study                  | Type of restrictions and setting | Study design                                      | Influenza strain involved | Strain transmissibility ($R_0$) | Scenario and duration of intervention                                                                 | Effect estimate                                                                 |
|-----------------------|---------------------------------|--------------------------------------------------|---------------------------|---------------------------------|--------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------|
| Colizza et al. (2007)  | Air travel, global              | Mathematical stochastic metapopulation model*   | A(H5N1)                   | 1.9                             | 20% or 50% air travel restrictions                                                                   | No impact on CAR                                                               |
| Epstein et al. (2007)* | Air travel, global              | Mathematical stochastic metapopulation model*   | Pandemic influenza        | 1.7                             | Hong Kong Special Administrative Region as source of epidemic, 95% restrictions implemented after 1000 infectious cases | If epidemic begins on 1 January or 1 July, it produces global means of 81,531,156 and 132,230,576 cases, respectively |
|                       |                                 |                                                  |                           |                                 | As above except Sydney, Australia, as source of epidemic                                            | If epidemic begins on 1 January or 1 July, it produces global means of 33,068,217 and 94,823,730 cases, respectively |
|                       |                                 |                                                  |                           |                                 | As above except London, United Kingdom, as source of epidemic                                       | If epidemic begins on 1 January or 1 July, it produces global means of 118,523,844 and 713,443 cases, respectively |
| Kernéis et al. (2008)* | Air travel, 52 cities worldwide | Mathematical stochastic metapopulation deterministic model* | Pandemic influenza strain (NS) | 1.8 or 49                        | Air travel restrictions of unspecified effectiveness, over various, unspecified timelines | Little effect on global burden or spatial and temporal diffusion of influenza pandemic |
| Lee et al. (2009)*    | Several scenarios               | Systematic review (deterministic and stochastic models) | Pandemic influenza (different strains) | 1.7 or 20                        | 90%, 99% or 99.9% air travel restriction                                                              | With $R_0$ set to 1.7 and 2.0 there was, respectively, no impact on overall attack rate and a 1% increase in that rate – with a 20% increase in PDAR |
| Marcelino and Kaiser (2012)* | Air travel, 500 major airports, worldwide | Mathematical stochastic metapopulation model* | A(H1N1)pdm09              | 1.7                             | Cancellation of a quarter of flight connections between 500 cities                                    | Number of circulating infected individuals reduced by an additional 19%        |

CAR: cumulative attack rate; CINC7: cumulative incidence seven months after start of epidemic; NIV: non-intervention value; NS: not specified; PDAR: peak daily attack rate; $R_0$: basic reproductive number.

* A so-called SEIR model in which individuals who are susceptible (S), exposed (E), infectious (I) or recovered (R) are considered.

* A so-called SLIR model in which individuals who are susceptible (S), latent (L), infected (I) or permanently recovered (R) are considered.

* The model took into account individuals who were nonsusceptible (NS), susceptible (S), exposed (E), infectious (I) or recovered (R).