

Modelling the epidemic spread of an H1N1 influenza outbreak in a rural university town

N. K. VAIDYA¹†, M. MORGAN²†, T. JONES³, L. MILLER², S. LAPIN² AND
E. J. SCHWARTZ^{2,3*}

¹ *Department of Mathematics and Statistics, University of Missouri – Kansas City, Kansas City, MO, USA*

² *Department of Mathematics, and* ³ *School of Biological Sciences, Washington State University, Pullman, WA, USA*

*Received 15 October 2013; Final revision 25 July 2014; Accepted 10 September 2014;
first published online 17 October 2014*

SUMMARY

Knowledge of mechanisms of infection in vulnerable populations is needed in order to prepare for future outbreaks. Here, using a unique dataset collected during a 2009 outbreak of influenza A (H1N1)pdm09 in a university town, we evaluated mechanisms of infection and identified that an epidemiological model containing partial protection of susceptibles best describes H1N1 dynamics in a rural university environment. We found that the protected group was over 14 times less susceptible to H1N1 infection than unprotected susceptibles. Our estimates show that the basic reproductive rate, R_0 , was 5·96 (95% confidence interval 5·83–6·61), and, importantly, R_0 could be decreased to below 1 and similar epidemics could be avoided by increasing the proportion of the initial protected group. Moreover, several weeks into the epidemic, this protected group generated more new infections than the unprotected susceptible group, and thus, such protected groups should be taken into account while studying influenza epidemics in similar settings.

Key words: Infectious disease epidemiology, influenza, mathematical modelling, Susceptible-Infected-Removed (SIR) model, university setting.

INTRODUCTION

Influenza infection is associated with 51 000 deaths each year in the USA, representing 2·2% of all US deaths per year [1]. The rapidly spreading pandemic influenza A(H1N1)pdm09 that emerged from Mexico in spring 2009 infected 43–89 million individuals by the end of the outbreak and was responsible for 8870–18 300 deaths, as estimated by the Centers for Disease Control and Prevention (CDC) [2]. This new virus caused death in otherwise healthy young adults

in larger numbers compared to typical influenza viruses [3]. Influenza A(H1N1)pdm09 swept through the rural university town of Pullman, WA, USA, in August 2009, and rapidly infected about 10% of the student population. In this community, which was particularly vulnerable to H1N1 infection due to more than 90% of individuals being young adults aged 18–26 years, individuals are well-informed and well-connected by the academic environment in which they interact with others in close proximity, and they are subject to minimal or no outside movement such as immigration or emigration, as well as no pharmaceutical or containment interventions. The ability to accurately understand and predict mechanisms of epidemic spread in the unique environment of a rural university town based upon collected data is of critical importance.

* Author for correspondence: Dr E. J. Schwartz, PO Box 643113, Pullman, WA 99164, USA.
(Email: ejs@wsu.edu)

† These authors contributed equally to this work.

Compartmental models [4] have been used in the past to analyse numerous influenza outbreaks including the H1N1 pandemic of 1918 and the 1968–1971 H3N2 epidemic [5], as well as to predict the potential severity of a future influenza outbreak [6] and to determine how vaccines or drug resistance may affect the spread of infection [7]. In the case of 2009 pandemic H1N1, modelling approaches were used to predict epidemic dynamics [8–11], to assess the efficacy of the CDC planned vaccination scheme for the US epidemic [12], to examine the effect of control measures [13], to assess the potential severity and transmissibility of the pandemic [14], and to determine the contagious period of this viral strain [15] (for review, see [16]).

Unlike studies of epidemic spread in urban settings [8, 9, 11], only a few studies have explored influenza dynamics in geographically contained settings, which can have quite different dynamics. Wearing *et al.* [17] used trajectory matching on incidence data to parameterize epidemic models and determine the basic reproductive rate (R_0) of an influenza outbreak in an English boarding school [18]. Mathews *et al.* [5] estimated R_0 in two populations, an H3N2 outbreak on an island in 1971 and the H1N1 pandemic in military camps in 1918. No studies, however, have examined H1N1 spread in a rural university setting, although there are dozens of such areas throughout the USA and elsewhere.

In this work we present a dataset from Washington State University (WSU) that describes the number of influenza cases reported to the university health service between August and December 2009. The small university town of Pullman, WA, USA is in a rural region surrounded by agricultural land with most students residing in town to attend the university. This dataset provides a unique opportunity to study an influenza epidemic spreading through the geographically contained community of a rural university town. Using this dataset we developed mathematical models to understand the mechanisms of infection for pandemic H1N1 [i.e. A(H1N1)pdm09] in a university student population. Specifically, we aimed to determine the role of self-protection on H1N1 dynamics, quantify the impact of the protected group and derive basic infection parameters. To our knowledge, this work is the first investigation to use compartmental modelling with data to understand H1N1 epidemic dynamics in a rural university campus setting. This may provide valuable insight into the dynamics of H1N1 infection and allow us to better prepare for the next epidemic by more accurately predicting the factors that influence H1N1 spread.

METHODS

Data collection

The epidemiological data were collected by WSU (Pullman, WA, USA) student health centre, Health and Wellness Services (HWS), from students who sought medical care from the centre. Data were collected daily from the start of the autumn semester on 22 August 2009 for 4 weeks, and thereafter on Mondays to Saturdays until the end of the epidemic on 3 December 2009. HWS is normally closed on Sundays but remained open once the epidemic was underway due to the number of ill students. Data were not available due to Sunday closure on 23 August 2009 and on Sundays from 20 September 2009 onwards, or on 26–28 November 2009 when HWS was closed for the Thanksgiving holiday. The population served by HWS includes undergraduate and graduate students. Each person who called or visited HWS with any symptom of influenza-like illness (ILI) was assumed to be positive for H1N1. Initially, all influenza that was seen and tested by HWS was confirmed to be influenza A(H1N1)pdm09. After 10 days, however, any individual with ILI was considered a case of H1N1, as per CDC recommendations [19]. The prevalence of infection was calculated assuming a recovery time of 6 days [15]; i.e. $\rho_t/N = \rho_{t-1}/N + \gamma_t/N - r_t/N$, where ρ_t is the number of individuals with infection on day t , γ_t are the new cases on day t , r_t are individuals who recover on day t , and N is the total population size (18 234).

Models

We employed a classical SIR model [20] as well as two models with protection of susceptibles (SPIR and alternate SPIR), and a quarantine model (SIQR) [21] for the purpose of modelling the 2009 H1N1 outbreak in Pullman, WA, USA. For each model we provide the expression for the basic reproductive rate, R_0 , which is defined as the average number of secondary infections that result when one infected individual is introduced into a population of susceptibles [22]. R_0 is a threshold that delineates whether an epidemic spreads ($R_0 > 1$) or dies out ($R_0 < 1$). We employed the next-generation method [23] to calculate R_0 .

Kermack–McKendrick model. In the SIR model, S represents the density of susceptible individuals, I represents the density of infected individuals, and R

represents the density of recovered individuals. The full model is given by

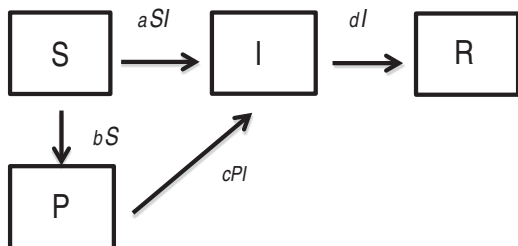


$$\begin{aligned} \frac{dS}{dt} &= -aSI, \\ \frac{dI}{dt} &= aSI - dI, \\ \frac{dR}{dt} &= dI, \end{aligned}$$

where a is the transmission rate and d is the recovery rate. The basic reproductive rate for this model is given by

$$R_0 = \frac{a}{d} S_0.$$

Model with protection. We also consider the following modification of the SIR model, where an additional compartment (P) represents susceptible individuals with self-protection from infection (hereafter, ‘protection’):



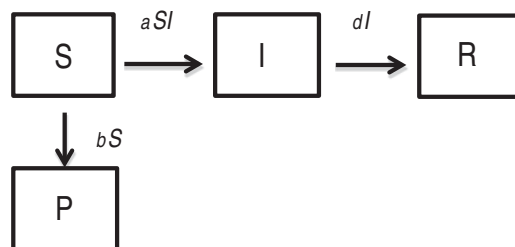
$$\begin{aligned} \frac{dS}{dt} &= -aSI - bS, \\ \frac{dP}{dt} &= bS - cPI, \\ \frac{dI}{dt} &= aSI + cPI - dI, \\ \frac{dR}{dt} &= dI, \end{aligned}$$

where a is the transmission rate of susceptible individuals without protection, b is the rate of protection, c is the rate at which protected individuals become infected, and d is the recovery rate. In this model $c > 0$ indicates that the protection from infection is partial. We found that the basic reproductive rate

for this model is given by

$$R_0 = \frac{aS_0 + cP_0}{d}.$$

Alternate model with protection that is 100% effective. In this model, 100% effectiveness of protected individuals is captured by considering absolutely no infection of individuals in P class.

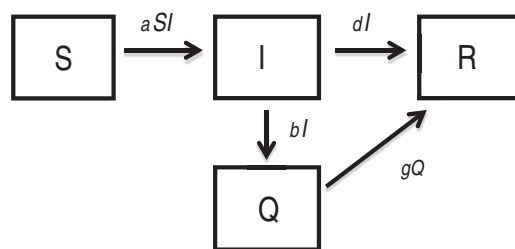


$$\begin{aligned} \frac{dS}{dt} &= -aSI - bS, \\ \frac{dP}{dt} &= bS, \\ \frac{dI}{dt} &= aSI - dI, \\ \frac{dR}{dt} &= dI. \end{aligned}$$

We found that the basic reproductive rate for this model is given by

$$R_0 = \frac{a}{d} S_0.$$

Epidemic management (quarantine) model. In this model [24], infected individuals self-quarantine at rate b , due to symptoms of infection, which removes them from mixing and infecting other susceptible individuals. Quarantined infected individuals recover at rate g .



$$\begin{aligned} \frac{dS}{dt} &= -aSI, \\ \frac{dI}{dt} &= aSI - bI - dI, \\ \frac{dQ}{dt} &= bI - gQ, \\ \frac{dR}{dt} &= dI + gQ. \end{aligned}$$

We found that the basic reproductive rate for this

model is given by

$$R_0 = \frac{a}{b+d} S_0.$$

Model assumptions and initial values

We consider this a closed population (i.e. a population with no births, deaths or continual immigration and emigration), since we considered birth and death during the semester to be insignificant due to the age group of the students, and we also considered entry and exit of students during the semester to be negligible, because students typically do not leave campus for the duration of the semester. We considered that the infection began after a single mass immigration event when students returned to campus after the summer break. Models assumed that all individuals with H1N1 contacted the university health centre, and no callers called more than once. Recovery time was 6 days [15]; thus the parameter $d = 1/6$ per day in all models. We assumed all infected individuals were symptomatic. Among the total population of size $N = 18\,234$, the initial populations in different compartments were as follows:

$$\begin{aligned} S(0) &= S_0 = 18223; & I(0) &= I_0 = 11; \\ R(0) &= P(0) = Q(0) = 0. \end{aligned}$$

Parameter estimation and model selection

We fitted several models to the data using a nonlinear least squares approach to find the model that best fits the data. We also estimated the parameter values for the potential transmission rates and protection rate. For the parameters of the best-fitting model, we calculated 95% confidence intervals using 10 000 bootstrap replicates [25].

For the purpose of model comparison, we calculated the error sum of squares (SSE), the regression sum of squares (SSR), the total sum of squares (SST) as well as R^2 , the coefficient of multiple determination, and R_{adj}^2 , the adjusted coefficient of multiple determination. The model with the greatest R_{adj}^2 , which balances the cost of using more parameters against the gain in R^2 , can be considered the model that best matches the data [26]. Furthermore, to determine if a group of predictors are jointly statistically significant, we performed the F test [27]. Note that the SIR model is the nested model of all other models considered. Using the F test, we compared the impact of the predictors of full models (SPIR, alternate SPIR, and SIQR) to the predictors of the nested SIR model in regards to best fit [26].

The test statistic used for this F test is

$$F = \frac{(SSE_j - SSE_k)/(k-j)}{SSE_k/(n-k)},$$

where SSE_j is the SSE value for the reduced model with j parameters, SSE_k is the SSE value for the full model with k parameters, and n is the number of data points. The subset of the predictors in the full model is significant if the F statistic is greater than the value of an F distribution with the appropriate degrees of freedom [26, 27].

RESULTS

Epidemiological data on the 2009 H1N1 outbreak in a rural university town

The prevalence data (the number of individuals with infection/total population size) are shown in Figure 1. The prevalence increased markedly over the first 2 weeks of the outbreak, reaching a peak of 3.57% on day 13. Cumulatively, over 2000 students were infected (i.e. more than 10% of the population). The epidemic resolved by the end of the semester.

SPIR model gives best fit to data

Kermack–McKendrick model fit. Using the recovery rate of $d = 1/6$ per day [15], the SIR model was fitted to the WSU dataset. It captures the initial rise in infection, the peak, and the resolution over the next several months, as seen in the epidemiological data. The prevalence of infected individuals is shown in Figure 2a. The R_{adj}^2 and other statistical measures of variation are shown in Table 1. We observed that the model peak fits the data better than the model tail, and therefore we tested whether a better fit would be produced by truncating the dataset on day 38, after the decline in infections following the peak. Fitting the model with the truncated dataset, however, did not show a better fit, as can be seen in Figure 2b and by the reduced R_{adj}^2 (Table 1). As the full dataset provides a better fit for this model, the full dataset was used in the subsequent studies.

SPIR model fit. Next we investigated whether we would find a better fit between the model and the data if we modified the model to include the condition that some individuals received some protection from infection during the epidemic. This protection could potentially consist of self-isolation or social distancing by susceptibles or other

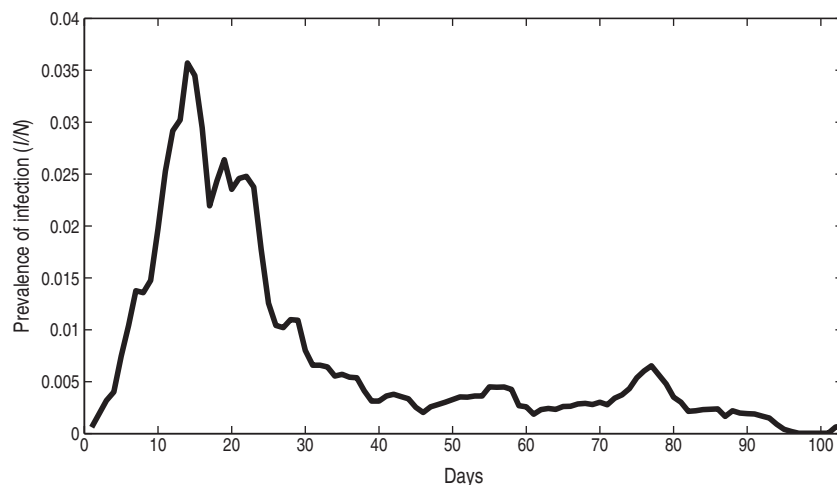


Fig. 1. University health centre data (infected fraction of total population over time).

preventative measures undertaken by susceptibles due to concern about the risk of infection. Plots of the prevalence of infection for the data and the fitted protection model (SPIR) are shown in Figure 3. The SPIR model prediction is in excellent agreement with the full dataset, and the R_{adj}^2 of the dataset with the SPIR model is greater than that with the SIR model (Table 1). This indicates that the SPIR model gives a better fit than the SIR model, signifying the importance of the protected group. Furthermore, we performed the F test and found that the better fit provided by the SPIR model is statistically significant (F test, $P < 0.001$).

Alternate SPIR model fit. The SPIR model investigated a situation where individuals in the population with some protection were still susceptible to infection. The alternate SPIR model explores a situation in which the effect of protection was absolutely perfect. Plots of the prevalence of infection for the data and model are shown in Figure 4. The F test showed that the alternate SPIR model did not produce a significantly better fit than the SIR model. The R_{adj}^2 of the dataset with the alternate SPIR model is less than that with the SPIR model (Table 1), indicating that the SPIR model gives a better fit than the alternate SPIR model. We next asked if the extra predictor in the SPIR model (compared to the alternate SPIR model), representing the transmission rate of the protected population, was a significant predictor for the goodness of fit. For this, we performed the F test with the alternate SPIR model as the nested model and the SPIR model as the full model, and found that the full model leads to a significantly better fit

of the observed data. Therefore the additional parameter in the SPIR model, the transmission rate of the protected population, contributes significantly to making the SPIR model the best-fitting model.

SIQR model fit. Next we investigated whether a better fit to the data would be found if we considered a model in which infected individuals were quarantined. Quarantine of infected individuals was thought to play a role because HWS had urged students to recover before resuming class attendance in order to slow the spread of the epidemic. Furthermore, some students reported that floors of their university dormitories were quarantined. On fitting the SIQR model to the full dataset (Fig. 5), we observed that the R_{adj}^2 of the fit with the SIQR model is less than that with the SPIR model (Table 1). This shows that the SPIR model gives a better fit to the data than the SIQR model. Further F tests confirmed that the additional predictors in the SIQR model (compared to the SIR model) do not improve the goodness of fit.

Estimation of parameters and basic reproductive rate for best-fitting model

Using the best-fitting model, SPIR, we estimated parameters a , b , and c for H1N1 epidemic dynamics in this rural university town (Table 2). Our estimates show that the transmission rate of susceptibles without protection was 5.45×10^{-5} [95% confidence interval (CI) 5.33×10^{-5} to 6.05×10^{-5}] per individual per day. During the epidemic, these susceptible individuals transfer into the protected group at the rate of 0.15 (95% CI 0.14–0.18) per day. Our results show that

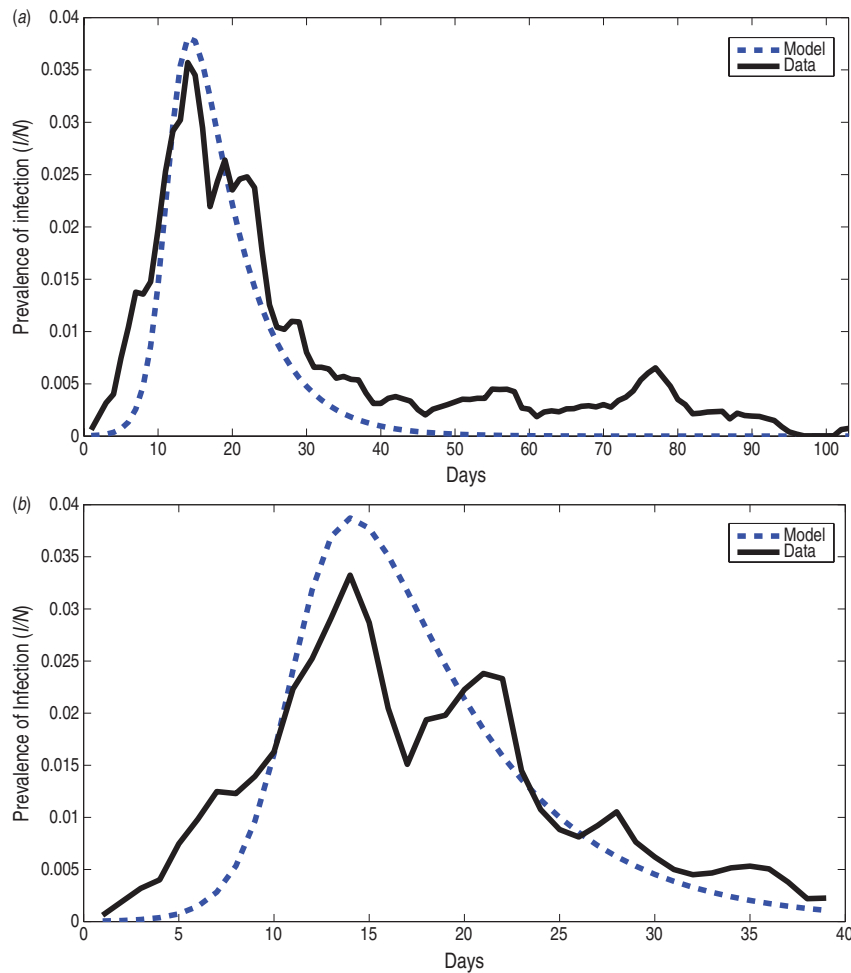


Fig. 2. (a) Prevalence of infected individuals for the Washington State University (WSU) full dataset from days 0 to 103, for data (solid line) and the SIR model (dotted line). (b) Prevalence of infected individuals for WSU truncated dataset from days 0 to 38, for data (solid line) and the SIR model (dotted line).

Table 1. *Statistics of model fitting*

Statistic*	SIR model†	SIR model‡	SPIR model†	Alternate SPIR model†	SIQR model†
SSE	0.0016	0.0013	0.0007	0.0016	0.0016
R^2	0.7886	0.5680	0.9060	0.7912	0.7886
R^2_{adj}	0.7865	0.5563	0.9032	0.7871	0.7844

* SSE, Error sum of squares; R^2 , coefficient of multiple determination; R^2_{adj} , adjusted coefficient of multiple determination.

† Modelled with full dataset.

‡ Modelled with truncated dataset.

this protection substantially reduces susceptibility to H1N1 infection by lowering the transmission rate over 14-fold, to 3.74×10^{-6} (95% CI 3.65×10^{-6} to 5.35×10^{-6}) per individual per day (Table 2). The basic reproductive rate, R_0 [22, 28, 29], obtained using our estimated parameter values from the best-fitting model (SPIR), was 5.96 (95% CI 5.83–6.61).

The protected group has a substantial impact on epidemic dynamics

We quantified the effect of the protected group. We investigated whether the epidemic could have been prevented if the initial proportion of protected individuals was increased, what proportion of the

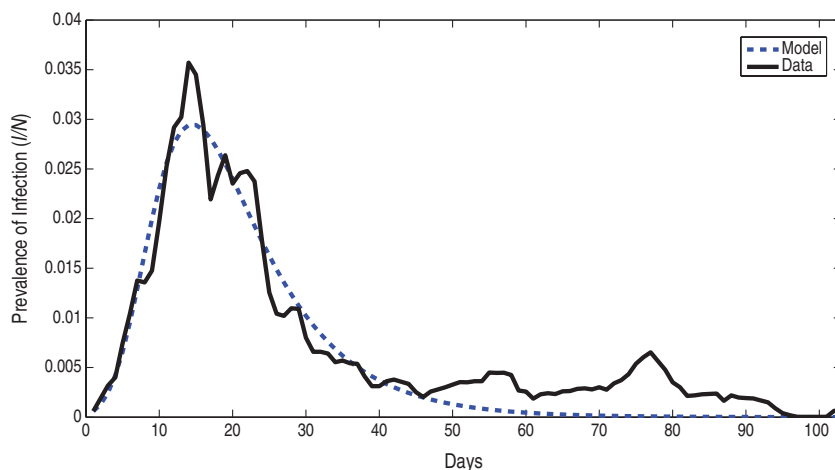


Fig. 3. Prevalence of infected individuals for the Washington State University full dataset, for data (solid line) and the SPIR model (dotted line).

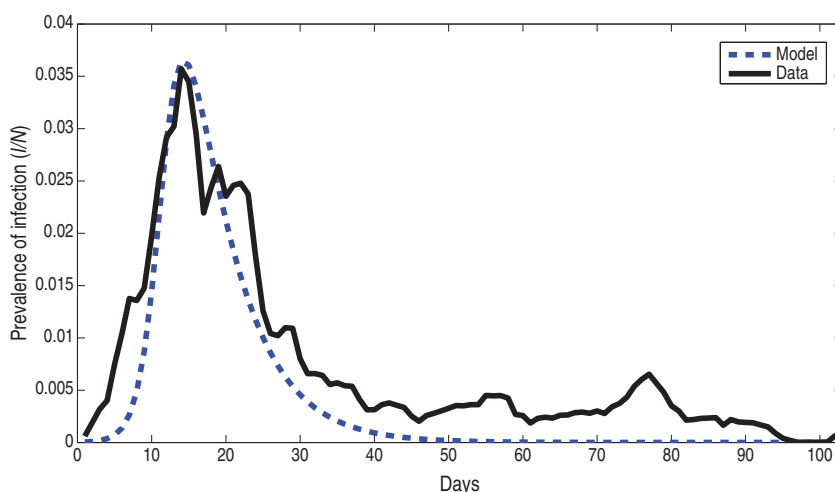


Fig. 4. Prevalence of infected individuals for the Washington State University full dataset, for data (solid line) and the alternate SPIR model (dotted line).

population was protected over the course of the epidemic, and how many new infections were generated by the protected group. We found that increasing the initial proportion of the protected group can substantially reduce R_0 (Fig. 6a). In fact, if the initial proportion of the protected population is high enough (around $\geq 90\%$), this would lower R_0 below 1. Thus a large initial protected group could prevent the epidemic. The protected group constitutes a large portion of the population, particularly after the first 3 weeks of the epidemic, reaching $>80\%$ of the population (Fig. 6b). As a result, a moderate number of new infections arise from the protected group over the course of the epidemic, although more new infections arise from the susceptible group (Fig. 6c). After the first 3 weeks of the epidemic, more new infections

arise from the protected group than from the susceptible group. Therefore, the protected group plays a substantial role in the epidemic dynamics.

DISCUSSION

In this study we investigated the transmission mechanisms and quantified the role of a protected group in the 2009 H1N1 influenza epidemic in Pullman, WA, USA. By fitting models to the epidemic data using non-linear optimization, we estimated the transmission parameters of the epidemic. Unlike many early studies of H1N1 transmission [14, 30–32] that considered either countrywide or international spread of infection, we investigated the 2009 H1N1 epidemic in a university town community subject to minimal to no outside

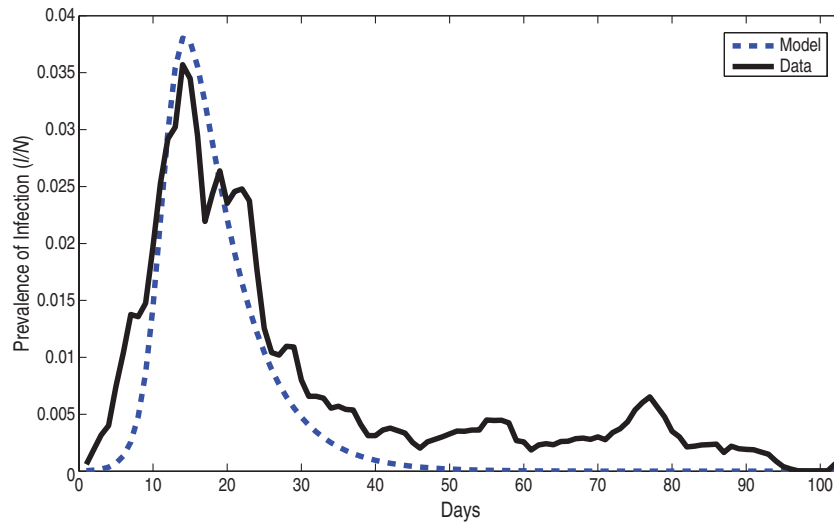


Fig. 5. Prevalence of infected individuals for the Washington State University full dataset, for data (solid line) and the SIQR model (dotted line).

Table 2. Model parameters and initial values for the best-fitting model, SPIR

Parameter	Units	Definition	Value (95% CI)*
Fixed parameter			
d	day ⁻¹	Recovery rate	1/6 [15]
Initial values			
$S(0)$	Population	Initial susceptible population	18 223
$P(0)$	Population	Initial protected population	0
$I(0)$	Population	Initial infected population	11
$R(0)$	Population	Initial recovered population	0
Estimated parameters			
a	(population*day) ⁻¹	Transmission rate (unprotected individuals)	5.45×10^{-5} (5.33×10^{-5} – 6.05×10^{-5})
b	day ⁻¹	Protection rate	0.15 (0.14–0.18)
c	(population*day) ⁻¹	Transmission rate (protected individuals)	3.74×10^{-6} (3.65×10^{-6} – 5.35×10^{-6})
R_0	(unitless)	Basic reproductive rate	5.96 (5.83–6.61)

* CI, 95% confidence intervals (generated by 10 000 bootstrap replicates).

intervention. In this unique setting, in which the population mostly consists of young adults aged 18–26 years, the spread of infection is largely governed by the interactions in the academic environment. For such environments, we identified that the SPIR model, which considers the potential effects of the protected group, was the best fitting model, with the highest R_{adj}^2 compared to the SIR model, the alternate SPIR model (with complete protection), and the SIQR (quarantine) model.

Our results suggest that a portion of the susceptible population was afforded some protection from infection during the epidemic. The protected compartment constitutes a large proportion of individuals after the first 3 weeks of the epidemic, at which point more new infections come from this population than the

susceptible population. This protection may have resulted from awareness about the epidemic and subsequent precautions undertaken by susceptible individuals to reduce their exposure, such as social distancing, increased hand washing [33] or vitamins and supplements [34]. This result is realistic, considering that the population at risk was extremely well informed. The university issued daily reports on the epidemic, urging individuals to be careful and indicating measures they could take to avoid infection. Furthermore, the population was well connected, since all individuals were involved in the educational programmes of the university, with the ability to share information through their interactions in classes and other activities. Vaccination was not thought to play a role in the protection, because the H1N1

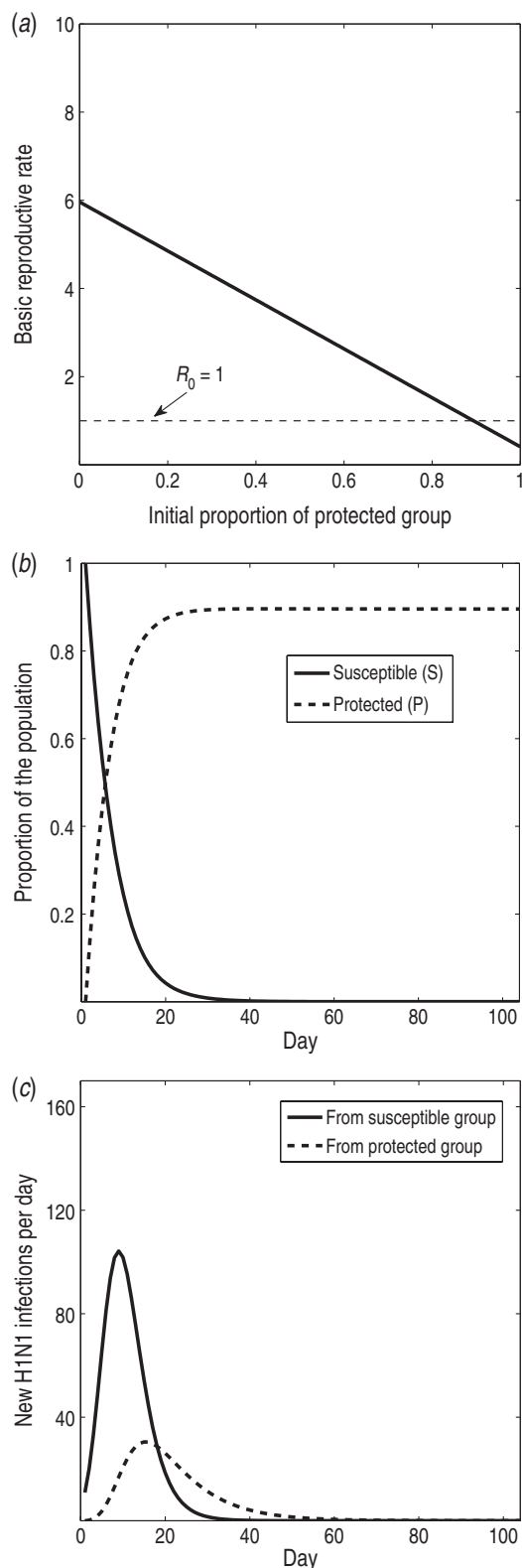


Fig. 6. (a) The basic reproductive rate, R_0 , as a function of the initial proportion of the protected group, $P(0)$ (solid line). The threshold, $R_0 = 1$, delineating whether an epidemic will spread or die out, is shown (dotted line). (b) Prevalence of susceptible individuals (solid line) and protected individuals

vaccine was not available before the outbreak; it was rolled out later in the year. Regarding quarantine, our results showed that despite the university's recommendation to stay home when ill, quarantine did not play a noticeable role.

This study makes an important contribution to the field of epidemic modelling for influenza. Theoretical modelling has been used to examine influenza spread in a geographically isolated area [6]; however, there are only a few cases where modelling and data have been combined for a closed epidemic [5, 17, 18, 35], and, to the best of our knowledge, no other studies focus on the rural university setting. The WSU H1N1 dataset is unique because it describes the epidemic from the beginning of the outbreak before the peak of infection, to its resolution, in a population on a university campus that was likely to have had little immigration or emigration. Unlike the boarding school dataset (number of data points, $n = 14$) [18] or the island dataset ($n = 50$) [5], in our study we have a much larger dataset ($n = 104$). While other studies used simulation or other modelling approaches to predict future epidemic dynamics [8–11, 14], in this work we used compartmental modelling to determine what mechanisms most likely played a role in the spread of this influenza epidemic in university town setting. Several characteristics distinguish this type of population from others, in that individuals are young adults aged between 18 and 26 years who are highly connected by the university, and they interact according to similar routines by way of adherence to the academic schedule.

Furthermore, finding reliable transmission data in closed or restricted populations to shed light on influenza dynamics is often difficult, particularly for epidemics such as the 2009 H1N1 pandemic, which affect the global population and are confounded by inconsistent data collection and the movement of individuals between different areas. Without data from a closed population, basic transmission parameters, including R_0 , can be difficult if not impossible to determine. Wearing *et al.* calculated R_0 for the boarding school epidemic to be 3.74 with the SIR model [17]. Mathews *et al.* estimated R_0 to be between 3.73 and 10.69 for the island outbreak, in which the infection was introduced to the population as a single

(dotted line) during epidemic using the SPIR model. (c) New infections arising from the susceptible group (solid line) and protected group (dotted line) during epidemic using the SPIR model.

immigration event, similar to our collected data [5]. Our SPIR model, which we found to be the best fit for the WSU H1N1 data, gave $R_0 = 5.96$ (5.83–6.61) for this rural university town environment. This value falls between those of these other estimates, and reflects that H1N1 spread rapidly through the campus community. This is reasonable; with most of the students living on or adjacent to campus, students may have spread the virus simply by social behaviour and attending classes even with influenza symptoms. Interestingly, we found that the value of R_0 is highly dependent on the initial proportion of the protected group in the university town, and with significantly higher levels of such a protected group, R_0 can be brought down to below 1 and an epidemic can be avoided.

Some limitations to this study should be noted. First, we modelled a population of students on a college campus within close proximity to one another at all times. Consequently, our results may not be generalizable to other populations. However, our results are relevant for many settings that share characteristics of our population including military units, islands, nursing homes, boarding schools, and other rural college campuses, in which the community is surrounded by a large region with low population, approximating a closed population. Rural college campuses differ from those near or in metropolitan regions, in which the immigration of infected individuals that can repeatedly seed the infection would be expected to occur more frequently and to a greater extent. Second, we used a model that assumed a closed population in which students did not leave for the duration of the epidemic. However, it may not be a strictly closed population, since a small percentage of students may commute or drop out, and non-students may travel in and out of the community. Third, we note that transmission rates vary among viruses, and hence our analyses may need to be repeated were a novel virus to appear. Furthermore, the modelling results presented here rely upon the use of deterministic modelling, which captures the dynamics of large populations but not small populations or rare events. Additionally, all students who were infected with H1N1 were assumed to have contacted HWS, and all students who contacted HWS with ILI were assumed to have had H1N1 as opposed to other illnesses. If these assumptions are inaccurate, then our results would underestimate the magnitude of the epidemic and the transmission rate, in the case that not all students with H1N1 contacted HWS, and our

results would overestimate transmission in the case that some students with ILI counted by HWS did not have H1N1. Finally, we acknowledge the limitation that the central result regarding the choice of the best-fitting model is entirely based on the R_{adj}^2 and F test of this rich dataset. Further study with more accurate data will provide a better estimate of the impact of the protected group.

The main goals of this work were to gain deeper insight into the mechanisms of epidemic spread of H1N1 and quantify the impact of the protected group in a rural college campus environment. The results identify the importance of the protected group and indicate that the protected group alone, if large enough, would be sufficient to reduce R_0 below 1. In conclusion, this work offers novel insight into influenza epidemiology and advances current knowledge of influenza epidemiology by indicating that effective epidemic control strategies in other geographically contained communities should consider the local population structure, as slowing a rapidly spreading outbreak may require different interventions than in the case of the general population.

ACKNOWLEDGEMENTS

The authors thank D. Garcia from Health and Wellness Services for providing the epidemiological data. We also thank D. Chao, E. Johnson, X. Wang, C. Cobbold, A. Dawes, and R. Tyson for useful discussions, as well as the anonymous reviewers for helpful suggestions. Finally we thank J. L. Kwon, G. Vogel, and A. Piazza for initial studies. E.J.S. acknowledges startup funding and seed grant funding from Washington State University. N.K.V. acknowledges startup funding from the University of Missouri–Kansas City and the UMRB grant from the University of Missouri Research Board.

DECLARATION OF INTEREST

None.

REFERENCES

1. **Thompson MG, et al.** Estimates of deaths associated with seasonal influenza – United States, 1976–2007. *Morbidity and Mortality Weekly Report* 2010; **59**: 1057–1062.
2. **Centers for Disease Control and Prevention.** Updated CDC estimates of 2009 H1N1 influenza cases, hospitalizations and deaths in the United States, April

- 2009–April 10, 2010 (www.cdc.gov/h1n1flu/estimates_2009_h1n1.htm). Accessed 14 May 2010.
3. **Fineberg HV.** Pandemic preparedness and response – lessons from the H1N1 influenza of 2009. *New England Journal of Medicine* 2014; **370**: 1335–1342.
 4. **Brauer F.** Compartmental models in epidemiology. *Lecture Notes in Mathematics* 2008; **1945**: 19–79.
 5. **Mathews JD, et al.** A biological model for influenza transmission: pandemic planning implications of asymptomatic infection and immunity. *PLoS ONE* 2007; **2**: e1220.
 6. **Roberts MG, et al.** A model for the spread and control of pandemic influenza in an isolated geographical region. *Journal of the Royal Society Interface* 2007; **4**: 325–330.
 7. **Alexander ME, et al.** Emergence of drug resistance: implications for antiviral control of pandemic influenza. *Proceedings of the Royal Society of London, Series B* 2007; **274**: 1675–1684.
 8. **Chao DL, et al.** FluTE, a publicly available stochastic influenza epidemic simulation model. *PLoS Computational Biology* 2010; **6**: e1000656.
 9. **Chao DL, et al.** Planning for the control of pandemic influenza A (H1N1) in Los Angeles County and the United States. *American Journal of Epidemiology* 2011; **173**: 1121–1130.
 10. **Miller L, et al.** Individual-based computational model used to explain 2009 pandemic H1N1 in rural campus community. *Journal of Biological Systems* 2013; **21**: 1340005.
 11. **Yang Y, et al.** The transmissibility and control of pandemic influenza A (H1N1) virus. *Science* 2009; **326**: 729–733.
 12. **Towers S, Feng Z.** Pandemic H1N1 influenza: predicting the course of a pandemic and assessing the efficacy of the planned vaccination programme in the United States. *Eurosurveillance* 2009; **14**: 19358.
 13. **Cruz-Pacheco G, et al.** Modelling of the influenza A (H1N1)v outbreak in Mexico City, April–May 2009, with control sanitary measures. *Eurosurveillance* 2009; **14**.
 14. **Fraser C, et al.** Pandemic potential of a strain of influenza A (H1N1): early findings. *Science* 2009; **324**: 1557–1561.
 15. **De Serres G, et al.** Contagious period for pandemic (H1N1) 2009. *Emerging Infectious Diseases* 2010; **16**: 783–788.
 16. **Boelle PY, et al.** Transmission parameters of the A/H1N1 (2009) influenza virus pandemic: a review. *Influenza and Other Respiratory Viruses* 2011; **5**: 306–316.
 17. **Wearing HJ, Rohani P, Keeling MJ.** Appropriate models for the management of infectious diseases. *PLoS Medicine* 2005; **2**: e174.
 18. **Anon.** Influenza in a boarding school. *Lancet* 1978; **1**: 587.
 19. **Centers for Disease Control and Prevention.** Interim recommendations for clinical use of influenza diagnostic tests during the 2009–10 influenza season (www.cdc.gov/h1n1flu/guidance/diagnostic_tests.htm). Accessed 29 September 2009.
 20. **Kermack WO, Mckendrick AG.** A contribution to the mathematical theory of epidemics. *Proceedings of the Royal Society of London, Series A* 1927; **115**: 700–721.
 21. **Nuno M, et al.** Mathematical models of influenza: The role of cross-immunity, quarantine and age-structure. *Lecture Notes in Mathematics* 2008; **1945**: 349–364.
 22. **Anderson RM, May RM.** *Infectious Diseases of Humans: Dynamics and Control*, 1st edn. Oxford: Oxford University Press, 1991.
 23. **Van den Driessche P, Watmough J.** Reproduction numbers and sub-threshold endemic equilibria for compartmental models of disease transmission. *Mathematical Biosciences* 2002; **180**: 29–48.
 24. **Hethcote H, Ma Z, Liao SB.** Effects of quarantine in six endemic models for infectious diseases. *Mathematical Biosciences* 2002; **180**: 141–160.
 25. **Efron B, Tibshirani R.** Bootstrap methods for standard errors, confidence intervals, and other measures of statistical accuracy. *Statistical Science* 1986; **1**: 54–75.
 26. **Devore JL.** *Probability and Statistics for Engineering and the Sciences*. Boston, MA: Brooks/Cole, 2012.
 27. **Bates DM, Watts DG.** *Nonlinear Regression Analysis and its Applications*. Hoboken, NJ: John Wiley & Sons Inc., 2007.
 28. **Heesterbeek JA.** A brief history of R_0 and a recipe for its calculation. *Acta Biotheoretica* 2002; **50**: 189–204.
 29. **Heffernan JM, Smith RJ, Wahl LM.** Perspectives on the basic reproductive ratio. *Journal of the Royal Society, Interface* 2005; **2**: 281–293.
 30. **Boelle PY, Bernillon P, Desenclos JC.** A preliminary estimation of the reproduction ratio for new influenza A(H1N1) from the outbreak in Mexico, March–April 2009. *Eurosurveillance* 2009; **14**.
 31. **Buckley D, Bulger D.** Estimation of the reproductive number for the 2009 pandemic H1N1 influenza in rural and metropolitan New South Wales. *Australian Journal of Rural Health* 2011; **19**: 59–63.
 32. **Nishiura H, Roberts MG.** Estimation of the reproduction number for 2009 pandemic influenza A (H1N1) in the presence of imported cases. *Eurosurveillance* 2010; **15**.
 33. **Cowling BJ, et al.** Facemasks and hand hygiene to prevent influenza transmission in households: a cluster randomized trial. *Annals of Internal Medicine* 2009; **151**: 437–446.
 34. **Roxas M, Jurenka J.** Colds and influenza: a review of diagnosis and conventional, botanical, and nutritional considerations. *Alternative Medicine Review* 2007; **12**: 25–48.
 35. **Murray JD.** *Mathematical Biology*, 3rd edn. New York, NY: Springer-Verlag, 2002.