SPECIAL TOPICS

Stochastic dynamic model of SARS spreading

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Abstract Based upon the simulation of the stochastic process of infection, onset and spreading of each SARS patient, a system dynamic model of SARS spreading is constructed. Data from Vietnam is taken as an example for Monte Carlo test. The preliminary results indicate that the time-dependent infection rate is the most important control factor for SARS spreading. The model can be applied to prediction of the course with fluctuations of the epidemics, if the previous history of the epidemics and the future infection rate under control measures are known.

Keywords: SARS, epidemic model, system dynamics, point process.

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It is an important topic for the research to establish the dynamic model of SARS spreading, to understand the characteristics and control factors, and to make predictions of the course of the epidemics. Most dynamic epidemic models in Chinese study are following the classic work of Anderson and May[1], by consideration of random processes in the corresponding differential equations[2-11]. Internationally, in addition to this kind of deterministic models[12-14], stochastic models[15] consist of another kind of work, either by consideration of random processes in the Markov chain and Monte Carlo simulation[19]. For epidemics of SARS, which has only a small number of tens to thousands patients in a region, and consists of a very small percentage of the entire susceptible population, a stochastic model is probably more appropriate[15,18]. So far, two epidemic models on SARS have been published[20,21], which all adopted the stochastic approach with a benefit recognized that the modeling parameters can be easily adjusted during the evolution of epidemics[22].

On one hand, epidemic equations can be applied not only to epidemics study itself, but also to other kind of social and natural scientific problems, such as biological group distribution, spreading of new technology, spreading of rumors in society, etc.[23,24]. On the other hand, scientific methods from other research areas can be made use of for epidemiology. A suggestive case comes from fluid dynamics: the fluid motion can be studied macroscopically by the partial differential equations of conservation of mass, momentum and energy; it can also be studied by tracing the motion, collision, and interaction of all the molecules, and approach the overall behavior by averaging their effects. The latter method as molecular dynamics
became tractable only after the development of advanced computers, which provide the technique possibility of calculating and tracing the motion of a large amount of molecules. In this study, we propose a stochastic model for SARS spreading by tracing the chain of infection of each patient and analyze the Vietnam case as an example. The methodology is similar to the molecular dynamics tracing every molecule to understand the macroscopic system, and the mathematic tools used is the point process similar to the stochastic model we used in the earthquake sequence analysis[25].

1 Model

In a preliminary form of SARS model, the spatial spreading and distribution can be neglected. We firstly focused on the temporal epidemic evolution within a specific area. Assuming that a SARS patient is imported, he will become infectious after the latent period. The latent period is believed to range within 1—12 d. In the model, a random parameter is assumed following the Poisson distribution. After the latent period, the patient becomes infectious, the number of people infected is determined in the computer by chance. Some infections may be traced as a result of close one-to-one contact, some may be infected in an indirect way (such as at the clinic of hospital, or flaws in the sewage system like the Amoy Garden case in Hong Kong). In any case, an averaged rate of direct or indirect infection per patient per day can be obtained, and their summation is the average infection rate for each patient. Because the distribution of infection rate is not yet well known for SARS as a new disease, we assume it to obey the Poisson distribution. The average infection rate may change at different stages of sickness for patient, and may change with social control measures taken. The length of the infectious period and prognosis of patients are all random variables. The infectious period may vary, and recover or die according to the mortality rate. The infection rate can be reduced by isolation measures. Some patients may become infectious first and then be isolated, some others may be quarantined first before they develop the symptoms, both can be produced stochastically in the model by computer. In summary, the latent period, date of infectious onset, date of isolation, and date of recovery or death, how many people newly infected each day for each patient, all these can be produced by the computer stochastically, stored in files, and updated every day of the epidemics. Therefore, with the four parameters (infection rate, latent period, infectious period, and mortality rate) known, it is possible to construct a simple dynamic model of SARS spreading. If spatial spreading of SARS is to be studied, it is necessary to add information on the spatial structure of the model, the probability of flow of people among different regions, and situation related infection rate at different regions. The flow chart for computation is shown in Fig. 1.

Among the four parameters, the latent period is not affected by mankind, while the infection rate, length of infectious period, and mortality rate are all subjected to changes produced by human measures. The morality rate does not affect the spreading of SARS in the model. Latent period, as a parameter not affected by human beings, does not vary significantly, and small changes of latent period do not make significant effects on the results. In Vietnam case, a one-day increase in the averaged latent period can produce a 2% decrease of duration of epidemics and 5% decrease in total number of patients. The duration of infectious period is reported to be about 10—14 d; change of one day can produce less than 5% changes in the results. Infection rate is the most important factor to

| Input initial values |
|----------------------|
| DO 100 I = 1, Imax (Cycle by day) |
| DO 200 J = 1, N total_sick (Scan each infected person) |
| In infectious period or not? |
| No | Yes |
| Compute number of newly infected person N will_sick |
| DO 300 K = 1, N will_sick |
| Construct infection file for each newly infected person, including information of region, infected by whom, infection date, latent period, recover or death date, isolation date, etc. All these produced by computer stochastically: |
| Update N total_sick |
| DO 400 J = 1, N total_sick |
| Daily statistics for patients |
| DO 500 J = 1, N zone |
| Daily statistics for regions |
| DO 600 I = 1, N zone |
| Final statistics |

Fig. 1. Computational flow chart of the stochastic model of SARS spreading.
affect SARS spreading. Infection rate is influenced by many factors. It is related to the course of the disease, for example, it is reported that the patient is more infectious from the 3rd to the 5th day of onset, and extremely infectious to medical workers during the operation of tracheotomia, but non-infectious in the latent period. The infection rate is also related to the population characteristics, such as population of the susceptible group, type of confluence as residential area, school and university, building set, village, etc. Each type may have different infection rates and control measures should be adjusted to fit the local conditions. There are some factors which people can control effectively to reduce the infection rate, such as early detection of SARS patients, isolation of patients and quarantine of close contacts to SARS patients, improvement of clinics to reduce cross infection, protection of medical workers, etc. All these will be embodied in one parameter: reduction of the infection rate.

Of course, in more realistic models, many other factors should be included, such as the susceptible people may be divided into different groups, and each group may have different infection rates, the recovered people may get immunity for life or just temporarily. There may be cases that the infected people do not become sick (or only have very light symptoms), and they may or may not be infectious. To consider the spatial spreading of SARS, it is necessary to construct a spatial structure of the model, to include the rate of flow or mixing of population between different regions, and the flow probability may vary between different types of regions (such as city—city, city—village), distances, means of transportation, time of seasons, holidays, or panic run away people after announcement of epidemics, etc. Although these factors are not included in the present modeling, it is not difficult to consider them in the future modeling.

2 Results

In the real world, mother nature produces a consequence randomly by her own rules: in the model, computer produces a result based on our understanding of the natural rules.

Two points should be noted: first, a good model or bad model is determined by the understanding of natural rules, we can trust the model more if we know the rules better. Second, even if the model is perfect in replication of the natural rules, it does not mean that we can exactly repeat the natural results. In the stochastic modeling, each run only produces one of the possible results randomly. The evaluation and use of the model should be based on the average results of a large amount runs, i.e. the Monte Carlo experiment. Only the average results can be compared with observation quantitatively.

Taking the example of Hanoi, Vietnam (data from WHO) as an example, Fig. 2 shows the actual daily reports of new patients evolves with time by day. The duration of the epidemics lasted 45 d, and the total number of patients infected is 62. Based on the present knowledge on SARS, the parameters chosen in our stochastic model are: the average latent period is 6.5 d, the average infectious period is 12 d. The infection rate \( r \) is defined as the number of people being infected per patient per day (unit: \( d^{-1} \)). It is found that the infection rate and its variation with time are the major factor to affect the epidemics. In this work, we made a great number of numerical tests, and by try and error, we found out the best model parameters to fit the constraints of duration of 45 d and accumulated number of 62 patients. The infection rate chosen in the final model is: before any control measures are taken, \( r_0 = 1.8 \ d^{-1} \), and among the 3rd to 5th days of sickness of the patient, the rate is 5 times higher than the base value; after control measures are taken, the average infection rate \( r \) is chosen as 0.01 \( d^{-1} \), and between the 3rd and 5th days, the rate is also 5 times higher as 0.05 \( d^{-1} \). The isolation measures are taken 7 d after the first patient is discovered. Small changes in the infection rate may produce significant changes in the results of modeling. In Vietnam case, an increase of \( r \) from 0.01 to 0.015 after control measures

![Figure 2](image-url)
were taken can result in an increase of 25% or more in the duration of epidemics and total number of patients.

Fig. 3 shows some typical results as examples of the model. The abscissa is time in day, the ordinate is the number of cases reported each day. Fig. 3(a)–(c) are some typical results similar to the actual case. Fig. 3(a) lasts 44 d of epidemics, infected 59 patients in total; Fig. 3(b) and (c) both last 44 d, and totally infected 67 patients. However, both the duration and the total number of patients may fluctuate significantly. In Fig. 3(d), the duration is 58 d and total number of patients reaches 105. In Fig. 3(e), total number of patients is only 34 within a duration of 31 d. Fig. 3(f) has a duration as long as 102 d, although the total infection is only 42 patients.

All these cases indicate some common features: i) Infected patients increase dramatically without control measures; ii) control measures show their effect not immediately, but with delays, the cases of SARS patients still increase and reach a peak a few days later because of onset of previously infected patients after the latent period; iii) effect of control measures will show up a few days later, and reported cases of SARS can reduce rapidly; iv) random fluctuations are likely to appear in the final stage, and keep the duration of epidemics longer.

These figures also show that fluctuations among different runs of the model can be significant, therefore, in order to understand the overall feature, a lot of test runs have to be carried on, and the averaged results should be used for comparison with observations. We made Monte Carlo tests, and based on 1000 times experiments, it is obtained from the model that duration of the epidemics is (49±15) d, and total number of patients is (61±22), the peak of daily reported cases appears on the (11th±3) day, and with a peak value of (8±2) cases, accumulated patients reach (28±13) at this peak day. Fig. 4 shows the frequency distribution of duration of epidemics and total cases reported.

It is a problem of special interest that if such kind of model can provide predictions on the epidemics while it is going on. Assuming that on the 13th d, it is already known that the peak of daily reported cases occurred on the 10th day of 9 cases that day and accumulated cases of 16 up to that day, considering the random fluctuations, we take parts of the results of Monte Carlotests, which satisfies the constraints that the peak appears on the 9th to 12th day, number of cases reported on the peak day is from 7 to 10 cases, and accumulated cases to the peak day ranges from 15 to 31 cases. Then we observe the subsequent evolution of these series by Monte Carlo modeling. It is found that in the circumstance that the first 12th days’ data are known, the conditional probability of subsequent development of epidemics can be obtained. It is expected that the duration of the epidemics would be (48 ± 13) d, and total number of patients would reach (58±13). Comparing corresponding predictions without knowing the 12th day data: duration of (49 ± 15) d and total number of patients of (61±22). It is found that there are only slight changes in the predicted value, but the estimation of fluctuations has been improved, especially the mean square deviation in the number of total patients reduces significantly. Frequency distribution is shown in Fig. 5, for comparison, the scale of axes is the same as in Fig. 4. The results suggest that
long as the infection rate at different stages can be estimated reasonably, it is possible to make predictions on the duration and total number of patients of the epidemics, providing not only the mean values and most probable values, but also the worst or best situations and the probability of appearance of these extremes. These can play a significant role in fighting against SARS and taking social and economical measures.

3 Discussions and conclusions

In epidemic models, a parameter called reproduction index $R$, the average number of people being infected by a patient, is considered to be critical. If it is greater than 1 at the beginning of epidemics without precaution measures, denoted as $R_0 > 1$, the epidemics will develop in deterministic models, and develop at a non-zero probability in stochastic models. If control measures are taken to reduce $R$, and make $R < 1$, the epidemics then can be controlled to decay\cite{18}. In this study, the infection rate $r$, the number of people infected per patient per day, provides detailed information of daily infection, and the summation of $r$ for all infectious days produces index $R$. In the discussion of Riley et al.\cite{20}, the infected people were divided into two categories: the normal part of infection, and the super-spreading event (SSE). A typical example of SSE was that a SARS patient from Hong Kong infected 25 staff members among the entire 26 staff members in the private French Hospital in Hanoi, Vietnam\cite{26}. Dye and Gay\cite{22} suggested that it is immature to conclude either infections can be divided into two categories as normal events and SSEs, or the so called SSE is just the long tail of a skewed distribution of infection. In this study, it is shown that a unified distribution including both normal events and the so-called SSEs can work well. The so-called super-spreaders just make the average infection rate higher. However, it is worthwhile to try more complex models in future study.

Reducing the infection rate is most important for stopping the spreading of SARS. Early control to reduce infection rate is of top priority. Numerical experiments indicate that the total case in Vietnam is only 62 because they took strict control measures to reduce the infection rate 7 d after the first case of SARS. If the measures are taken one month after the first case of SARS, at reasonable similar parameters, the total cases would be increased to about 1000; if measures are taken 45 d after the first case,
the total cases would reach nearly 3000; and if measures are taken 60 d after the first case, the total number of patients would jump to nearly 9000.

This report is just to provide a test of the suggested methodology of our SARS model, which emphasizes the use of the state of art computation ability to trace infection of each patient and construct a stochastic model for SARS spreading. This model is somehow similar to the Markov chain and Monte Carlo method in the previous studies[19], but it is more flexible to model various complexities. The preliminary test of the example of Vietnam shows the feasibility of this attempt. The model can simulate not only the different stages of SARS spreading (the accelerating stage at first, the delayed appearance of peak infection, the rapid but fluctuated decay under effective control), but also provide quantitative estimation of the duration of epidemics and total number of patients under given model parameters. During the epidemics, if the leading part has become known, it is possible to apply the model for prediction of subsequent evolution of the epidemics. In practical forecast, it is noted that an oversimplified model has no practical values, while an over complex model cannot obtain enough epidemic data to constrain the parameters. Therefore, a trade-off has to be made[19]. This work is preliminary, it needs to collect the detailed data of SARS epidemics and make use of the special capability of tracing every patient in this model, with in-depth comparison of the observation and stochastic modeling, it is possible to develop the present model to contain new findings and details on SARS research as well as the spatial spreading process.

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