Early Detection of Nosocomial Outbreaks Caused by Rare Pathogens: A Case Study Employing Score Prediction Interval

Hiroshi Nishiura a,b,*

aSchool of Public Health, University of Hong Kong, Hong Kong. 
bPRESTO, Japan Science and Technology Agency, Saitama, Japan.

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Abstract

Objectives: Nosocomial outbreaks involve only a small number of cases and limited baseline data. The present study proposes a method to detect the nosocomial outbreaks caused by rare pathogens, exploiting score prediction interval of a Poisson distribution.

Methods: The proposed method was applied to three empirical datasets of nosocomial outbreaks in Japan: outbreaks of (1) multidrug-resistant Acinetobacter baumannii \((n = 46)\) from 2009 to 2010, (2) multidrug-resistant Pseudomonas aeruginosa \((n = 18)\) from 2009 to 2010, and (3) Serratia marcescens \((n = 226)\) from 1999 to 2000.

Results: The proposed method successfully detected all three outbreaks during the first 2 months. Both the model-based and empirically derived threshold values indicated that the nosocomial outbreak of rare infectious disease may be declared upon diagnosis of index case(s), although the sensitivity and specificity were highly variable.

Conclusion: The findings support the practical notion that, upon diagnosis of index patient(s), one should immediately start the outbreak investigation of nosocomial outbreak caused by a rare pathogen. The proposed score prediction interval can permit easy computation of outbreak threshold in hospital settings among healthcare experts.

1. Introduction

Nosocomial infection refers to the infection event within medical and healthcare facilities at which medical services for some diseases or health conditions are provided. The nosocomial infection is seen not only among patients but also among patients’ relatives and healthcare workers. Because medical and healthcare facilities involve treatment of a wide spectrum of diseases and thus patients tend to be vulnerable to
infectious diseases, less virulent and commonly seen pathogens can often cause nosocomial infection, and moreover, prior antibiotic treatment tends to induce infections caused by antibiotic-resistant bacteria. The nosocomial infection can occur regardless of the size of healthcare facility, and technically it will never be eliminated. However, the nosocomial infection can sometimes influence the prognosis of patients, and so healthcare experts are expected to control an outbreak event by detecting it at the early stage. To investigate and understand the epidemiology of any nosocomial outbreaks, epidemiological surveillance would play a key role [1]. In Japan, the Ministry of Health, Labour, and Welfare has conducted a routine surveillance program of nosocomial infection [2], and each medical facility with an independent clinical laboratory section maintains the system of infectious agent surveillance report through the isolation of causative pathogens from patients’ samples. Given such a system, it is fruitful to utilize the information, in particular, by detecting any outbreak during the early stage.

To analyze the community-based surveillance data, various statistical and epidemiological studies on the early detection of community outbreaks have been conducted. Farrington and Andrews [3] comprehensively reviewed representative detection methods for the investigation of the temporal or spatiotemporal incidence data of infectious diseases. In addition to classical statistical modeling approaches, hypothesis testing for nosocomial outbreaks caused by rare pathogens, applying it to actual outbreak datasets and assessing the validity of detection.

2. Materials and Methods

2.1. Observed data and motivation

To clearly demonstrate the study motivation, the observed epidemic curves of three nosocomial outbreaks are presented in Figure 1. The outbreak data were retrieved from openly published case notification reports. Figure 1A shows the monthly incidence of multidrug resistant Acinetobacter baumannii (MDR-AB) in a tertiary hospital with approximately 1150 beds (n = 46) from 2009 to 2010. While Acinetobacter baumannii is broadly distributed in the environment, its nosocomial infection is known to easily spread from person to person, and thus it is hard to control without substantial effort [8]. Within healthcare facilities, the infection is frequently seen among patients who are intubated for respiratory support, and MDR-AB is known as a key factor to exacerbate respiratory function and elevate the risk of death [8,9]. Figure 1B shows an epidemic curve of a nosocomial outbreak caused by multidrug resistant Pseudomonas aeruginosa (MDRP) at a secondary hospital with approximately 580 beds (n = 18) from 2009 to 2010. MDRP is frequently isolated from patients staying for the long time and also those requiring surgical management or antibiotic treatment [10]. The extent of transmission is sometimes a single ward scale (e.g., through contaminated hand-washing basin), worsening clinical course of infected patients in that particular ward [11]. Figure 1C and D show the monthly counts of Serratia marcescens isolations (n = 226), counting the total samples and isolates only from blood samples at a secondary hospital with 380 beds from 1999 to 2000. Although the isolation of Serratia marcescens has not been rare in this hospital for the long time, especially when we count the total samples (Figure 1C), an abrupt increase in severe cases was observed from May to June 2000 with eight fatal outcomes. Separately counting the samples by anatomical site, isolation from blood samples showed an apparent increase in the corresponding period (Figure 1D), and indeed, many severe cases had experienced septic shock before an outbreak investigation was conducted. Serratia marcescens is commonly isolated from the respiratory and urinary tracts of hospitalized adults and is responsible for catheter-associated bacteremia, urinary tract infections and wound infections. All three diseases have greatly influenced the prognosis of patients, indicating the importance of early detection of the outbreaks and corresponding actions for swift control. If the outbreak can be detected sufficiently early, infected cases within a single ward may be managed together, with particular care for the prevention of further transmission events (e.g., admit and manage cases in the same room) and moreover, nurses
and other staffs responsible for infected cases may also be limited to particular persons. In addition, doctors can order additional laboratory testing of other suspicious cases at the early stage, and respiratory function of infected cases can be closely monitored.

In the present study, an outbreak is defined as the occurrence of defined infectious disease cases clearly in excess of the normal expectancy within a certain period of time. This study attempts to detect the outbreak based on monthly counts of cases using a statistical model. In reality, the detection does not have to rely solely on the temporal data. Usually, additional insights are gained from contact information and other risk-associated information (e.g., whether the infection is opportunistic or not), and also from examining the spatiotemporal distribution of cases within a hospital. Moreover, microbiological and clinical findings (e.g., isolation of similar genotypes from multiple patients) can help demonstrate the transmission events. Among these, the present study specifically focuses on the temporal distribution for two reasons. First, the temporal data are routinely collected even when there is no outbreak. In other words, the temporal counts of bacterial isolations or case notifications would be readily available at any time, and such a dataset should be effectively used for public health purposes. Second, for a rare pathogen, even an occurrence of a single case may be regarded as an outbreak in practical sense. Then, an outbreak defined by an occurrence of index patient(s) does not practically require rigorous statistical detection. However, issuing an alarm based on a single diagnosis (or diagnosis of the first few cases) should ideally rest on rigorous scientific grounds, and the present study offers the theoretical basis examining the relevant condition at which one can declare the outbreak caused by rare pathogens.

2.2. Outbreak and prediction interval

Here the outbreak detection is described by equations. To detect an outbreak based on the time series data, the issue of epidemiological detection is conventionally expressed as the hypothesis testing of “aberration” (i.e., if the observed data exceeds a defined threshold) under a certain type I error [3]. That is, using $\theta_1$ to define the upper bound for detecting an outbreak, the observed data $Y$ should satisfy

Figure 1. Monthly incidence of nosocomial outbreaks. (A) Multidrug resistant Acinetobacter baumannii (MDR-AB) in a tertiary hospital with approximately 1150 beds ($n = 46$) from 2009 to 2010. (B) Multidrug resistant Pseudomonas aeruginosa (MDRP) at a secondary hospital with approximately 580 beds ($n = 18$) from 2009 to 2010. (C) Total samples and (D) blood samples of Serratia marcescens ($n = 226$) at a secondary hospital with 380 beds from 1999 to 2000. The horizontal axis represents the month of diagnosis. Before the observation in the earlier year (i.e., 2009 for A and B, and 1999 for C and D), there was no report of cases since January.
\[ \Pr(Y > \theta_U | \text{no aberration}) = \alpha, \]  

(1)

where \( \alpha \) is the probability that normal observation is incorrectly detected as an outbreak and may be interpreted as the risk of false positive alarms (e.g., one may use \( \alpha = 0.025 \)). The upper bound \( \theta_U \) thus acts as the reference value for detection, and this key value should be calculated from the prediction interval, i.e., the expected range within which the population data in future lies. The interpretation of the prediction interval resembles that of the confidence interval (CI): the 95% CI of a sample indicates the “range in which the population data lie at 95% probability,” while the 95% prediction interval represents the range in which the future population data, which cannot be observed at present, lie at 95% probability based on observed data in the past [12]. Let \( x = \{x_i\} \) be the sample monthly counts of cases based on observation in month \( i \) (\( i = 1, 2, \ldots, n \)) for the length of \( n \) months, and let us consider the predicted number of cases \( y \) in \((n + 1)\)th month. Then, the interval \((L(x), U(x))\) which satisfies

\[ \Pr(L(x) \leq y \leq U(x)) = 1 - 2\alpha, \]  

(2)

is referred to as the 100\((1-2\alpha)\)% prediction interval \((L(x), U(x))\).

Among the published prediction intervals, the simplest one may be based on an assumption that the population data follow a normal distribution, and this method was actually employed by the Centers for Disease Control and Prevention and applied to various practical settings [13]. Assumting that there is no trend in the occurrence of cases, let the sample average and sample standard error be \( \bar{x} \) and \( s \), respectively. The 100\((1-2\alpha)\)% prediction interval for \((n + 1)\)th month given past observation for \( n \) months is

\[ \left( \bar{x} - z_{1-\alpha} \frac{s}{\sqrt{n}}, \bar{x} + z_{1-\alpha} \frac{s}{\sqrt{n}} \right), \]  

(3)

where \( z_{1-\alpha} \) is the 100\((1 - \alpha)\) percentile of the standard normal distribution (e.g., 1.96 for the 95% prediction interval). Although detailed derivation process of (3) is omitted here, this point is discussed in a variety of literature on the interval estimation [13,14].

There are two technical problems in applying the abovementioned prediction interval for three nosocomial outbreaks in Figure 1. First, it is strictly not appropriate to apply normal distribution to the datasets with very small counts. Nevertheless, although the prediction interval of continuous distributions tends to be studied relatively well [15,16], that of discrete distribution has not been often discussed, except for normal approximations by means of the Wald method. Second, the occurrence has been very uncommon due to causation by rare pathogens, and thus, the baseline information is extremely limited. Sometimes, the survey starts only after confirming the diagnosis of index patients.

### 2.3. Statistical model

Since the occurrence is very rare with very small number of observed cases, the present study ignores the time trend (i.e., assumes stationary process) and employs a Poisson distribution. The transmission dynamics of infectious diseases are theoretically described by the Poisson process, and the resulting number of cases with time in an endemic equilibrium is known to follow a Poisson distribution [17]. In the stationary state, there is no increase or decrease in the number of cases (i.e., the nonstationarity indicates the outbreak). Let \( X \) and \( Y \) be the random variables representing the cumulative number of cases for \( n \) months and the number of cases in \((n + 1)\)th month. It is assumed that \( X \) follows a Poisson distribution with an average \( X_n \) (where \( X_n \) is the observed sample cumulative number), and also that the predicted value in \((n + 1)\)th month similarly follows a Poisson distribution with an average \( \theta \). These satisfy the following equation:

\[ \theta = \frac{X_n}{n}. \]  

(4)

To derive the prediction interval, we consider a random variable \( W \) that represents the difference between \( Y \) in \((n + 1)\)th month and predicted value \( X/n \) for \((n + 1)\)th month (i.e., \( W = Y - X/n \)). The average of \( W \) is obtained from the averages of \( Y \) and \( X/n \), i.e.,

\[ \mu_w = \mu_y - \frac{\mu_x}{n} = \theta - \frac{n\theta}{n} = 0. \]  

(5)

The variance of \( W \) is calculated as the sum of the variances of \( Y \) and \( X/n \):

\[ \sigma_w^2 = \sigma_y^2 + \frac{\sigma_x^2}{n^2} + \theta + \frac{n\theta}{n^2} = \theta \left( 1 + \frac{1}{n} \right). \]  

(6)

Standardizing \( W \), we get

\[ z = \frac{Y - X/n}{\sqrt{\theta \left( 1 + \frac{1}{n} \right)}}. \]  

(7)

Provided that \( \theta \) and \( X_n/n \) are sufficiently large, the probability \( z \) asymptotically follows a normal distribution, and we obtain

\[ z_{1-\alpha} = \frac{Y - X_n}{\sqrt{X_n \left( 1 + \frac{1}{n} \right)}}. \]  

(8)

As we compute the Wald CI, the 100\((1 - 2\alpha)\)% prediction interval based on equation (8) is

\[ \left( \hat{Y} - z_{1-\alpha} \sqrt{\frac{X_n}{n} \left( 1 + \frac{1}{n} \right)}, \hat{Y} + z_{1-\alpha} \sqrt{\frac{X_n}{n} \left( 1 + \frac{1}{n} \right)} \right), \]  

(9)

where \( \hat{Y} \) is equal to \( X_n/n \) in the range \( X_n > 0 \). If \( X_n \) is zero (i.e., no occurrence in the past), an arbitrary small
value, e.g., $X_n = 0.5$, is conventionally adapted for the computation [18]. The approximate prediction interval (9) based on asymptotic normality is referred to as the Nelson prediction interval [19]. It should be noted that the coverage probability of the Wald CI for a normal approximation to the binomial distribution is known to be extremely small when the binomial probability is too close to 0 or 1 [20]. In the case of approximate prediction interval (9) for the Poisson distribution, the coverage probability should also be small for small number of observations, and thus, the applicability of prediction interval (9) may be limited [21]. However, the exact prediction interval is too complex for non-experts, and moreover, the exact prediction interval of discrete distribution is known not necessarily to yield better coverage probability as compared to approximate ones [20].

Hence, a score prediction interval, which is relevant to Wilson score CI that yields much better coverage probability than the Wald method in equation (9), is derived. The score prediction interval of a binomial distribution has already been proposed in a statistical study and published elsewhere [22]. To derive the score prediction interval of a Poisson distribution, let us consider a joint sampling of $X$ and $Y$, as if the predicted value of the variance of $W$ in equation (7) is $\theta_{xy} = (X+Y)/(n+1)$. Namely, we use the following quantity that asymptotically follows a normal distribution:

$$
z_{1-a} = \frac{Y - X_n}{n} \cdot \sqrt{\frac{X_n + Y}{n + 1} \left(1 + \frac{1}{n}\right)}.
$$

As mentioned above, no occurrence in the past with $X_n = 0$ is replaced by $X_n = 0.5$. The score prediction interval is derived from taking the square of both sides of equation (10) and solving it for $Y$ as a quadratic equation of $Y$ [20,23]. Thus, the $100(1 - 2\alpha)$% prediction interval is calculated as:

$$
\left(\frac{X_n + z_{1-a}^2}{2n} - z_{1-a} \sqrt{\frac{z_{1-a}^2}{4n^2} + \frac{X_n}{n^2} + \frac{X_n}{n}}\right),
$$

$$
\left(\frac{X_n + z_{1-a}^2}{2n} + z_{1-a} \sqrt{\frac{z_{1-a}^2}{4n^2} + \frac{X_n}{n^2} + \frac{X_n}{n}}\right).
$$

2.4. Application to nosocomial outbreak data

Using the prediction interval (11), the early detection was attempted for the observed three nosocomial outbreaks. For all three outbreaks, the hospital surveillance had been routinely conducted before the outbreak, and the baseline data were available from January of the corresponding earlier year of observation. As mentioned above, as long as the number of reports in the past remains zero, theoretical cumulative number $X_n = 0.5$ was used for the computation of the prediction interval. The $1^{st}$ month at which the observed number exceeded the upper 95% prediction interval was regarded as the month of successful detection.

Subsequently, the detection performance was assessed by employing the receiver operating characteristic curve, which was used for identifying an empirically defined optimal cutoff point to define an outbreak, especially by referring to the Youden index (i.e., sensitivity plus specificity minus 1) [24]. The period of outbreak was defined to be from the $1^{st}$ month to the last month with reporting of at least one case: August 2009 to August 2010 for MDR-AB, May 2009 to March 2010 for MDRP and May–June 2000 for Serratia (although a few earlier cases occurred in summer 1999, they were not clinically serious and were separated from the 2000 outbreak). Using the optimal threshold of monthly case counts, the sensitivity and specificity of outbreak detection were estimated. The 95% CIs of the sensitivity and specificity were computed using normal approximation to the binomial distribution, and similarly, the calculation of the 95% CI of the area under the curve (AUC) was made using the Wald method by means of logit transformation of the AUC.

3. Results

Figure 2 compares observed and predicted values along with the upper 95% prediction intervals for all three outbreaks. The observed number of cases initially exceeded (and thus, the outbreak was detected) in the $1^{st}$ month (August 2009) for MDR-AB and the $2^{nd}$ month for MDRP (June, 2009) and sepsis caused by Serratia marcescens (June, 2000). The calculated upper 95% prediction intervals for these months were 0.97, 1.62, and 1.64 cases for MDR-AB, MDRP, and Serratia, respectively. That is, if we round up the thresholds to the next integer, the proposed method suggests that one should use the cutoff number of 1, 2, and 2 cases to define the outbreak caused by the rare pathogens. If we round down the threshold values that are 1 or greater, all the cutoff values would suggest one case to define the nosocomial outbreak.

Empirical optimal threshold was also estimated to be one case for all three outbreaks. However, the AUC varied by outbreak and were estimated to be 100%, 78.1% (95% CI, 53.7–91.7), and 87.5% (95% CI, 26.4–99.3) for MDR-AB, MDRP, and Serratia, respectively. Since there was no month with zero report during the outbreak of MDR-AB, both the sensitivity and specificity were estimated at 100%. As for MDRP, the specificity was 100%, but the sensitivity was calculated at 56.3% (95% CI, 37.5–81.3%) due to several zero reports during the course of the outbreak. With respect to Serratia marcescens, the sensitivity was 100%, but the specificity was 60.0% (95% CI, 30.0–90.0%) due to a few isolation reports from blood samples before the outbreak.
4. Discussion

The present study proposed the score prediction interval for detecting nosocomial outbreaks caused by rare pathogens, applying the method to three actual outbreak events in Japan, caused by MDR-AB, MDRP and *Serratia marcescens*. The proposed approach is regarded as an extension of a classical method invented by Stroup et al [13], which employed a normal distribution with a static baseline, in that the nonhomogeneity (i.e., the outbreak) in the proposed approach can be identified even for rare diseases with very small number of counts for the baseline as it exploits a Poisson distribution. In all three outbreaks, the threshold to define the outbreak was computed to be one or two cases, which agreed well with empirically calculated threshold based on the receiver operating characteristic curve and Youden index. These findings support the notion that one should immediately start the outbreak investigation of any nosocomial outbreak caused by rare pathogens upon diagnosis of index patient(s).

To the best of the author’s knowledge, the present study is the first to epidemiologically support the notion that the nosocomial outbreak caused by rare pathogens should be regarded as an outbreak even when there is only one case. Based on the single report of index case, the hospital may issue an alert. Although declaring an outbreak and starting the investigation and interventions with one case can easily be justified in practice, the present study has offered a firm theoretical support for that action and demonstrated its scientific validity. However, the performances of outbreak detection (e.g., sensitivity, specificity, and AUC) were not shown to be always excellent, because of the nature of using threshold at one case, with highly variable sensitivity and specificity values. However, the imperfect performance of detection when using the defined threshold should not be regarded as a flaw of the proposed model, but rather the nature of reliance on the temporal distribution of cases. In reality, the declaration of outbreak can also account for additional information such as spatiotemporal growth of cases, the times of admission and illness onset among cases, and identification of risk factors through outbreak investigations.

An advantage of using score prediction interval is that the coverage probability is much higher than other approximate prediction intervals proposed in the past [19,21,25,26]. Moreover, the present study has shown that the analytical solution remains to be tractable, and the proposed score prediction interval permits easy computation in hospital settings among healthcare workers using spreadsheet program. Given a ward-based surveillance data for *n* months with cumulative counts $X_n$, and given that we wish to test if there is an outbreak within the ward, one can satisfy this task by comparing the observed counts in the *n* + 1th month against

$$
\theta_U = \frac{X_n}{n} + \frac{(1.96)^2}{2n} + 1.96 \sqrt{\frac{(1.96)^2}{4n^2} + \frac{X_n}{n^2} + \frac{X_n}{n}}.
$$

and, if the observed data exceeds the threshold, the hospital ward may regard it as an abnormal excess. Due to its simplicity, equation (12) has a potential to greatly help any local clinical setting (including clinical laboratory section) to issue an alarm of excess without devising any complex computer system. In fact, it is known that the isolation of MDR-AB in the laboratory section was not sufficiently informed to infection control team during the MDR-AB outbreak in Figure 1A during its early stage, and using equation (12) among clinical or laboratory experts should have at least helped recognize the abnormality.

Although there have been a number of studies aiming to detect infectious disease outbreaks employing a variety of sophisticated mathematical and statistical techniques,
not so many detection systems have been put into practice, especially at the levels of local medical and healthcare facilities. In particular, early detection approach to small scale outbreaks such as nosocomial ones has been extremely limited. In this sense, I believe that equation (12) based on the proposed method and scientific support to issue an alarm upon diagnosis of index patient(s) would greatly help in managing nosocomial outbreaks at the hospital- and ward-levels in the future.

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