April 2009

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Saeed Akhtar
Aga Khan University

Shafquat Rozi
Aga Khan University

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Recommended Citation
Akhtar, S., Rozi, S. (2009). An autoregressive integrated moving average model for short-term prediction of hepatitis C virus seropositivity among male volunteer blood donors in Karachi, Pakistan. World Journal of Gastroenterology, 15(13), 1607-1612.
Available at: https://ecommons.aku.edu/pakistan_fhs_mc_chs_chs/39
An autoregressive integrated moving average model for short-term prediction of hepatitis C virus seropositivity among male volunteer blood donors in Karachi, Pakistan

Saeed Akhtar, Shafquat Rozi

Saeed Akhtar, Department of Community Medicine and Behavioral Sciences, Faculty of Medicine, Kuwait University, PO Box 24923, Safat 13110, Kuwait
Shafquat Rozi, Department of Community Health Sciences, Medical College, Aga Khan University, Stadium Road, Karachi 74800, Pakistan

Author contributions: Akhtar S designed the study, analyzed the data, and wrote the manuscript; Rozi S participated in HCV surveillance, helped in data collection, and data management.

Supported by Department of Community Health Sciences, Faculty of Medicine, Aga Khan University, Karachi, Pakistan
Correspondence to: Saeed Akhtar, PhD, Department of Community Medicine and Behavioral Sciences, Faculty of Medicine, Kuwait University, PO Box 24923, Safat 13110, Kuwait. saeed.akhtar@hsc.edu.kw
Telephone: +965-2498-6542 Fax: +965-2533-8948
Received: September 8, 2008 Revised: December 18, 2008 Accepted: December 25, 2008
Published online: April 7, 2009

Abstract

AIM: To identify the stochastic autoregressive integrated moving average (ARIMA) model for short term forecasting of hepatitis C virus (HCV) seropositivity among volunteer blood donors in Karachi, Pakistan.

METHODS: Ninety-six months (1998-2005) data on HCV seropositive cases (1000⁻¹ × month⁻¹) among male volunteer blood donors tested at four major blood banks in Karachi, Pakistan were subjected to ARIMA modeling. Subsequently, a fitted ARIMA model was used to forecast HCV seropositive donors for 91-96 mo to contrast with observed series of the same months. To assess the forecast accuracy, the mean absolute error rate (%) between the observed and predicted HCV seroprevalence was calculated. Finally, a fitted ARIMA model was used for short-term forecasts beyond the observed series.

RESULTS: The goodness-of-fit test of the optimum ARIMA (2,1,7) model showed non-significant autocorrelations in the residuals of the model. The forecasts by ARIMA for 91-96 mo closely followed the pattern of observed series for the same months, with mean monthly absolute forecast errors (%) over 6 mo of 6.5%. The short-term forecasts beyond the observed series adequately captured the pattern in the data and showed increasing tendency of HCV seropositivity with a mean ± SD HCV seroprevalence (1000⁻¹ × month⁻¹) of 24.3 ± 1.4 over the forecast interval.

CONCLUSION: To curtail HCV spread, public health authorities need to educate communities and health care providers about HCV transmission routes based on known HCV epidemiology in Pakistan and its neighboring countries. Future research may focus on factors associated with hyperendemic levels of HCV infection.

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Key words: Hepatitis C virus; Blood donor; Ecological analysis; Autoregressive integrated moving average model; Pakistan

INTRODUCTION

Hepatitis C virus (HCV) infection poses a major public health problem in developing countries, including Pakistan. However, the results of prevalence studies have shown variable estimates in select groups including 1.8% to 3.0% in volunteer blood donors[1,2] and 16% to 20.5% in familial contacts of infected patients[3,4]. A community-based study in Hafizabad, Punjab, found a 6.5% HCV seroprevalence[5]. Using these estimates, Pakistan has been grouped into intermediate category with respect to burden of HCV infection[6]. Several routes have been implicated for nosocomial and community acquired...
HCV infection including unsafe injections, recycling of used syringes, inadequate sterilization of surgical and dental equipment, and facial shaving by barbers\cite{23}.

Public health authorities in Pakistan intermittently run educational campaigns in electronic and print media to create awareness in the general population to halt HCV spread. However, in the absence of adequate HCV surveillance, the true impact of the HCV control efforts remains uncertain. Volunteer blood donors are generally considered to be the healthier segment of any community and the proportions of HCV seropositivity among them may be considered to mirror the situation in the general population\cite{8}. We have previously reported a significant increase in HCV seroprevalence among volunteer blood donors over the past several years using data from two blood banks\cite{8}. However, there is need to expand this HCV surveillance network countrywide to obtain more reliable and representative estimates.

Recently, mathematical models have been used to project the future HCV prevalence among intravenous drug users\cite{9}, and its impact on the future development of HCV related morbidity and mortality\cite{10}. Modeling and forecasting HCV seropositivity among volunteer blood donors in Pakistan, and perhaps in other neighboring countries, might provide useful information for allocating resources, and re-shaping and planning future control activities\cite{11}. This study aimed to develop a univariate time series model for HCV seropositivity (1000\(^{-1}\) × month\(^{-1}\)) among volunteer blood donors attending four large blood banks. Specifically, the objective of this study was to identify the stochastic autoregressive integrated moving average (ARIMA) model for short term forecasting of HCV seropositivity (1000\(^{-1}\) × month\(^{-1}\)) among volunteer blood donors in Karachi, Pakistan.

**MATERIALS AND METHODS**

**Setting**

This study was conducted in Karachi—the largest cosmopolitan city and the hub of economic activity of Pakistan. It has an estimated population of 9.3 million, accounting for approximately 10% of the total population of the country. Forty three percent of the city’s population is under the age of 15 years. The population of Karachi comprises several ethnic groups defined by mother tongue, including predominantly Urdu, Sindhi, Punjabi, Pushto, and Balochi. The healthcare facilities for the population include several small and tertiary care hospitals, both in the private and public sector.

**Data**

Eight-year (1998-2005) data on monthly aggregates of number of donors attending four large blood banks (blood bank I-IV) in Karachi were available for this study. These blood banks receive blood donations only from non-remunerated volunteer blood donors. Blood bank I is part of a tertiary care hospital in the private sector and receives blood donations as replacements from friends and relatives of inpatients requiring blood transfusions. Blood banks II-IV belong to non-governmental organizations and cater for the needs of those in Karachi who need blood transfusions, including the patients with leukemia, hemophilia, thalassemia and other blood related diseases. Blood banks II-IV also receive blood donations from volunteers on an exchange basis. Prior to blood donation, each blood donor is subjected to screening for known risk factors for transfusion transmissible infections. All the blood banks follow similar criteria to receive blood donations and exclude potential donors who admit known risk factors of transfusion transmissible infections or any medical or non-medical condition associated with high risk (e.g., use of narcotic drugs, history of jaundice in the past 5 years and recent hospitalization). All four blood banks in the study use commercially available enzyme-linked immunosorbant assay kits and results are interpreted according to the manufacturer’s instructions.

As noted earlier, blood donations between January 1998 and December 2005 by men aged 18-64 years were included in this evaluation. HCV serological results of consecutive blood donations from these blood banks were available from variable starting dates depending on the completed records, to assess the proportions of HCV seropositive donors.

**Analytic approach**

We used methods developed by Box and Jenkins to build an ARIMA time series model\cite{12}. This model-building process is designed to take advantage of associations in the sequentially lagged relationships that usually exist in data collected periodically. The general form of the ARIMA model was

\[
\Delta z_t = \Phi_1 z_{t-1} + \ldots + \Phi_p z_{t-p} + \theta_1 \Delta z_{t-1} + \ldots + \theta_q \Delta z_{t-q} + \epsilon_t
\]

where:

- \(\Delta z_t\) = differenced series i.e. \(z_t - z_{t-1}\)
- \(z_t\) = set of possible observations on the time-sequenced random variable
- \(\epsilon_t\) = random shock term at time \(t\)
- \(\Phi_1, \ldots, \Phi_p\) = autoregressive parameters of order \(p\)
- \(\theta_1, \ldots, \theta_q\) = moving average parameters of order \(q\)

The series was subjected to Box-Cox transformation\cite{13}. The transformed series was then differenced at the non-seasonal level and mean corrected to induce stationarity. Sample autocorrelation and partial autocorrelation functions were used to identify the ARIMA model of the appropriate order. Estimates of the model’s parameters were obtained by the maximum likelihood method. Diagnostic checking included residual analysis and the Akaike Information Criterion was used to compare goodness-of-fit among ARIMA models. The final model was a result of several iterations of the identification, estimation, and checking process, and met the conventional criteria for the adequacy of the model\cite{14}.

**Assessment of forecast accuracy**

The last 6 observations in the data set were used for validation of the forecast accuracy of the ARIMA
RESULTS

Descriptive analysis

The crude HCV seropositivity ($1000^{-1}$) among the male volunteer blood donors during the study period was 20.3 (12,792/630,134). The mean prevalence ($1000^{-1} \times \text{month}^{-1}$) was 18.3 [95% confidence interval (CI): 16.8-19.9]. There was no statistically significant difference in HCV seropositivity ($1000^{-1}$) across various months of the years ($F = 0.201; P = 0.997$) (data not shown).

However, a substantial variation in HCV seroprevalence ($1000^{-1}$) was observed across different calendar years ($F = 47.895; P < 0.001$) (Table 1). The observed and transformed series are presented in Figure 1.

ARIMA model

The parameters' estimates for the optimum ARIMA (2,1,7) model for the series of monthly HCV seropositive donors ($1000^{-1}$) are shown in Table 2. The autocorrelation and partial autocorrelation functions of the residuals showed good-fit (Figure 2). The residual plots showed small variations around the zero mean. None of these residuals had its magnitude larger than twice the standard deviation. Residuals' autocorrelations were not significantly different from zero as a set and had constant variance, thus confirming the adequacy of the model (Ljung-Box statistic = 20.4; $P = 0.433$).

The forecasts by the ARIMA (2,1,7) model for 91-96 mo (June 2005 to December 2005) using the fitted ARIMA model was used to forecast the HCV seroprevalence ($1000^{-1} \times \text{month}^{-1}$) for 91-96 mo (June 2005 to December 2005) to contrast with the observed series of the same months. The average forecast error at prediction interval of $m$ months ($e_m$) was calculated as:

$$e_m = \sqrt{\frac{1}{m} \sum_{i=m+1}^{m+n} (\hat{y}_{i-m} - y_{i-m})^2}$$

Where $y_{i-m}$ and $\hat{y}_{i-m}$ denote the observed and forecast values for month $i + m$. Finally, the fitted ARIMA model was used for short term (January 2006 to June 2006) forecasts along with their 95% confidence limits beyond the observed series.

![Figure 1](image-url)  Hepatitis C virus seroprevalence ($1000^{-1} \times \text{month}^{-1}$) among volunteer male blood donors in Karachi, Pakistan 1998-2005. A: Observed data along with forecasts; B: Transformed series.

![Figure 2](image-url)

**Table 1** Hepatitis C virus seroprevalence ($1000^{-1} \times \text{year}^{-1}$) among male volunteer blood donors at four large blood banks in Karachi (1998-2005)

| Yr | Mean | SD  | 95% CI for mean | Minimum | Maximum |
|----|------|-----|-----------------|---------|---------|
|    |      |     | Lower limit     | Upper limit |         |
| 1998 | 13.8 | 1.5 | 12.8            | 14.7     | 12      | 16      |
| 1999 | 16.3 | 2.2 | 14.8            | 17.7     | 13      | 21      |
| 2000 | 18.5 | 2.2 | 17.1            | 19.9     | 15      | 22      |
| 2001 | 19.8 | 2.0 | 18.6            | 21.1     | 16      | 22      |
| 2002 | 19.9 | 3.0 | 18.0            | 21.8     | 15      | 26      |
| 2003 | 20.3 | 3.2 | 18.2            | 22.3     | 16      | 27      |
| 2004 | 28.3 | 2.6 | 27.6            | 30.9     | 25      | 33      |
| 2005 | 24.8 | 2.1 | 23.4            | 26.1     | 21      | 27      |
| Total | 20.3 | 5.1 | 19.3            | 21.3     | 12      | 33      |

$F = 47.9; df = 7, 88; P < 0.001$. 

The fitted ARIMA model was used to forecast the HCV seroprevalence ($1000^{-1} \times \text{month}^{-1}$) for 91-96 mo (June 2005 to December 2005) to contrast with the observed series of the same months. The average forecast error at prediction interval of $m$ months ($e_m$) was calculated as:

$$e_m = \sqrt{\frac{1}{m} \sum_{i=m+1}^{m+n} (\hat{y}_{i-m} - y_{i-m})^2}$$

Where $y_{i-m}$ and $\hat{y}_{i-m}$ denote the observed and forecast values for month $i + m$. Finally, the fitted ARIMA model was used for short term (January 2006 to June 2006) forecasts along with their 95% confidence limits beyond the observed series.
observed series of months 1–90, closely followed the pattern of observed series for the same months (Figure 1), with mean ± SD and maximum monthly absolute forecast errors (%) over 6 mo interval being 6.5% ± 3.4% and 10%, respectively. Furthermore, the short term (January 2006 to June 2006) forecasts beyond the observed series adequately captured the pattern in the data (Figure 1) and showed evidence of increasing tendency of HCV seroprevalence (1000 folds) with the mean ± SD as 24.3 ± 1.4 over the forecast interval.

**DISCUSSION**

Epidemiological surveillance of communicable diseases is one of the more traditional public health activities. Time series analysis of surveillance data on prevalence and/or incidence of various infections may be helpful in developing hypotheses to explain and anticipate the dynamics of the observed phenomena and subsequently in the establishment of a quality control system and re-allocation of resources. This method is an ecologic approach and takes advantage of the strong association in the sequentially lagged relationship that usually exists in the data collected periodically.

During the study period, the overall HCV seroprevalence (1000 folds) in volunteer blood donors was 20.3, which falls in the range of 14.9 to 38.9 known for first time blood donors in other developing countries. However, HCV seroprevalence (1000 folds) in this study was much higher than the 2.1 reported for developed countries. The low HCV seroprevalence in resource-rich countries is attributed to safe blood transfusion, whereas, in poor regions of the world, several million people acquire HCV infection each year as a result of contaminated transfusions and the re-use of infected medical devices. Therefore, public health practices adopted by the developed countries need to be strictly enforced in less developed countries to break the chain of transmission of HCV and other blood-borne pathogens.

Monitoring of HCV seropositivity among volunteer blood donors may provide clues about the effectiveness of control efforts of public health authorities and future trend of the proportion of HCV infected donors in Pakistan. In this paper, we used the ARIMA model on a time series of HCV seropositivity (1000 folds) collected monthly over a period of 96 mo on asymptomatic male volunteer blood donors from four major blood banks in Karachi. The forecasts made in a prospective manner over six months demonstrated increasing tendency of HCV seropositivity among the blood donors in this cosmopolitan city. Such a predicted increase in HCV seropositivity might result from inconsistent and naïve HCV control efforts on the part of public health officials in Pakistan. Therapeutic injections in a healthcare setting have consistently been shown as a strong risk factor for HCV infection in Pakistan, and if concerted efforts by the public health authorities are not made, might continue to contribute to the increasing load of HCV infection in this and similar settings in the region. An increasing trend among first time US blood donors of 50 to 59 years of age from 1995 to 2002 has been demonstrated. According to the authors, teenage children and young adults in 1960, and 1970s might have experimented with drug injection and were infected with HCV. These people entered into the 50 to 59 years age group during 1995 to 2002. However, in other age groups of donors in the same study and two other studies from US, and from other developed countries (France and Spain) have shown a decreasing trend of residual risk of HCV infection in blood donors. According to these investigations different factors could have played a role in this reduction, for instance, increased awareness about the factors associated with increased risk of HCV infection, voluntary deferral by potential high risk donors of 50 to 59 years of age from 1995 to 2002 has been demonstrated.

### Table 2

| Parameters | Estimate | Standard error | t-ratio |
|------------|----------|----------------|---------|
| Autoregressive parameter (θ) | 0.67 | 0.15 | 4.5 |
| Moving average parameter (θ) | -0.59 | 0.18 | 3.3 |
| θ | 0.49 | 0.15 | 3.3 |
| θ | -0.80 | 0.11 | 7.3 |
| θ | 0.74 | 0.21 | 3.5 |
| θ | -0.37 | 0.17 | 2.2 |

White noise variance = 9.74; Ljung-Box Q statistic = 20.4 (P = 0.433).

**Figure 2** Residual plots for the final ARIMA (2,1,7) model of HCV seroprevalence (1000 folds) among male volunteer blood donors in Karachi, Pakistan 1998–2005. A: Autocorrelation function; B: Partial autocorrelation function.
donors, improvement in donor recruitment, and/or an overall decrease in HCV infection level in the general population. Such factors need to be evaluated in our population in future studies.

Results from our previous study[2], and those predicted by ARIMA model for 6 mo beyond the observed data exhibited a slightly increasing tendency of HCV seropositivity among male volunteer blood donors over the forecast period. This increasing pattern of HCV seroprevalence among these asymptomatic male volunteer donors merits further investigation of factors contributing to HCV seroprevalence in this population, which is thought to be a mirror image of the situation in the general population.

Some limitations of this study need to be taken into account when interpreting the results. Our HCV seroprevalence estimates are based on ELISA, which has sensitivity of more than 95%. However, these results do not reflect possible HCV infections that do not produce detectable seropositivity during the window period of HCV infection. The exact proportion of these HCV infected, but HCV seronegative, is not known, however, it has been argued that this figure must be very small given the use of current sero-assays[20]. Our HCV seroprevalence estimates in male volunteer donor population were based on data from a limited number of blood banks; we do not know whether they reflect the national average. The blood banks that participated in this study however, account for a substantial proportion of donations made annually in Karachi. These centers are located in large metropolitan areas where the prevalence and/or incidence of HCV may be higher than the national figures. Therefore, we think we are justified in making generalizations from our data. In conclusion, in the absence of comprehensive HCV surveillance in the general population in Pakistan and perhaps in other neighboring countries, further monitoring of HCV seropositivity in blood donors and the investigation of factors associated with hyperendemic HCV infection using multivariate ARMIA models might further expand our understanding about HCV epidemiology in this region. Furthermore, effective screening of all blood donors for HCV infection at all blood banks should be seriously considered, because one single HCV infected regular blood donor could transmit the infection to several recipients.

ACKNOWLEDGMENTS
We gratefully acknowledge the support provided by the administration and staff of the participating blood banks and the Department of Community Health Sciences, Aga Khan University, Karachi, Pakistan in the conduct of this study.

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S- Editor Li LF  L- Editor Stewart GJ  E- Editor Zheng XM