Epidemiological and virological characteristics of seasonal and pandemic influenza in Lao PDR, 2008–2010

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Background Information on influenza virology and epidemiology from Lao PDR is limited and the seasonal patterns of influenza have not been previously described.

Objectives To describe epidemiological and virologic characteristics of influenza in Lao PDR to recommend public health interventions, including improvements in surveillance and response.

Patients/Methods We performed a descriptive analysis of samples taken from patients with influenza-like-illness (ILI) (fever >38°C with cough and/or sore throat) presenting at seven sentinel hospitals in three regions of Lao PDR, January 2008–December 2010. A nasopharyngeal (NP) swab or combined nasal with oropharyngeal swab was collected from patients with ILI. Samples were tested for influenza by either Luminex RVP, conventional reverse transcriptase PCR (RT-PCR) (January 2008–2009), or by real-time PCR (rRT-PCR) using US CDC reagents (February 2009 onward).

Results Of 2346 samples tested from patients with ILI, 523 (22%) were positive for influenza. The median age of those positive was 12 years (range, <1–60 year). The percentage of samples that were influenza positive was similar over the 3 years (20–23%). Each year 3–4 types/subtypes cocirculated with differing predominant type/subtype. Influenza was detected year-round with the highest proportion of positive specimens in the 3rd and 4th quarter.

Conclusions Similar to other countries in the region, we found that influenza is present year-round and has a peak activity from July to December. Dominant types or subtypes vary by year. A large proportion of patients with ILI are not influenza positive. ILI surveillance is critical for weighing disease burden, both morbidity and mortality, against the costs of advancing influenza vaccine delivery strategy.

Keywords Epidemiology, influenza, Lao PDR, virology.

Introduction Influenza is a viral infection and common seasonal illness causing epidemics worldwide. It is estimated that about 10–20% of the world’s population is affected by seasonal influenza each season, with an average of 250 000–500 000 deaths annually.1 The majority of fatal cases related to influenza in developed countries occur among people aged 65 years or older.2 In addition, in temperate climates, there is a distinct influenza seasonal pattern, typically occurring in the colder winter months.3,4 In tropical countries, however, the epidemiology is less well defined, with available data suggesting that seasonal influenza viruses circulate throughout the year with one or two peaks, particularly during the rainy season.3,5,6 Other information on influenza, including seasonality, prevalence, and estimates of disease burden, is limited in tropical countries.7,8 Such data are essential to better inform national and international prevention strategies.

Lao PDR occupies part of the Mekong Region of Southeast Asia and falls just within the northern hemisphere (18° latitude and 105° longitude), straddling both tropical and...
semi-tropical zones. More than 70% of the 5.6 million population live in rural areas. The climate is subtropical with monsoon rains from May to October. Lao PDR has limited capacity for influenza surveillance and response. However, in response to avian influenza A (H5N1) and the 2009 pandemic (influenza A (H1N1)pdm09), Lao PDR has been building capacity to respond to emerging infectious diseases, including strengthening surveillance and laboratory capacity.

Laboratory confirmation of influenza has been available at the National Centre for Laboratory and Epidemiology (NCLE) in Vientiane since 2006, and Influenza-like-Illness (ILI) virological surveillance has been ongoing since 2007. Prior to this, data on confirmed influenza infections were scarce in Lao PDR, as laboratory confirmation was only available in a few hospitals and included (i) the total number of outpatients with ILI and (ii) the number of outpatients with ILI.

Descriptive analyses of aggregate data and case data from patients with ILI were conducted in stata 10.0 (Statacorp, College Station, TX, USA). Data were compared using the chi-squared test or Wilcoxon signed-rank test, where appropriate. A two-tailed $P$-value $< 0.05$ was considered significant.

Some data from 2008 have previously been presented in an early report on the laboratory-based influenza surveillance in Lao PDR.12

**Specimen collection**

An NP swab or combined nasal swab with oropharyngeal swab was collected from all patients with ILI on one (central) or two (provincial) days a week from each site. Once samples were collected, swabs were put in Viral Transport Media, stored at 4°C, and then transported to the NCLE within 24 hours. Case data, including demographic and clinical information, were collected on a laboratory form from all patients from whom a swab was collected.

From January 2008 to January 2009, samples were tested for influenza virus by either Luminex RVP or conventional reverse transcriptase-polymerase chain reaction (RT-PCR). From February 2009 onward, samples were tested for influenza A and B viruses by real-time reverse transcriptase PCR (rRT-PCR) using US CDC reagents.

RNA from throat and nasal swab specimens from each patient was extracted using the QiAamp Viral RNA mini commercial kit (Qiagen Inc., Mettmann, Hilden, Germany) according to manufacturer’s instructions. The SuperScript III Platinum one-step quantitative RT-PCR system (Invitrogen, Carlsbad, CA, USA) was used for single-step rRT-PCR according to US CDC protocol. The primers and probes for the influenza virus (H1N1, H3N2, A (H1N1)pdm09, H5N1, and influenza B) were derived from US CDC (as a WHO Influenza Collaborating Center) protocols and shared by US CDC through the WHO National Influenza Centers’ (NICs) Network.

From 15 May 2009 onward, positive influenza A specimens were further subtyped for H1, H3, and A (H1N1)pdm09. If the specimens tested positive for influenza A/H1, H3, or A (H1N1)pdm09, then they were also inoculated into MDCK cells and isolates were identified by the haemagglutination inhibition assay (HIA) using the WHO or National Institute of Infectious Diseases (NIID), Japan influenza diagnostic kit. If the result of the specimen subtyping for H1, H3, and A (H1N1)pdm09 was negative, then subtyping was conducted for H5.

The influenza virus isolates presented in Table 2 come from specimens of a number of sources: patients who meet the ILI and severe acute respiratory infection (SARI) case definition (a person who meets the ILI case definition plus shortness of breath or difficulty breathing and who requires...
hospital admission), outbreak investigation cases, and any other ad hoc samples received for influenza testing including from enhanced surveillance during the A(H1N1) pdm09 pandemic. The isolates were identified by the haemagglutination inhibition assay (HIA) using the WHO or NIID influenza diagnostic kit. Further characterization required the use of specific ferret antiserum.

Results

On average, ILI cases account for 10% (50 390/509 313) of all patients presenting to OPD and ER of hospitals in Vientiane Capital from 2008 to 2010. The highest monthly numbers of ILI presented to surveillance sites between May and September, before and during the rainy season (Figure 1). On the basis of aggregate data, patients with ILI were younger than all OPD/ER patients (64% being under 18 years of age versus 28%) (Figure 2). Patients from whom respiratory samples were obtained and tested had similar age distribution compared with all patients presenting with ILI (Figure 2). For Vientiane Capital, only 3% (1511/50 390) of all patients with ILI were sampled and tested for influenza.

For the 3-year period, 2346 samples from patients with ILI presenting to all sentinel healthcare facilities were received in good condition and tested. The majority of samples (65%) were from hospitals in the central region, with 22% from the north and 13% from the south. The majority of samples were tested in 2010 (50%), with 36% in 2009 and 14% in 2008. The median age of patients with ILI tested was 7 years (range, <1–89 years; IQR = 1–6–21 years) and the male-to-female ratio was 1:1.1. Of these samples, results from testing of eight samples were uninterpretable and not repeated, mostly during 2008 (n = 7), and therefore excluded from further analysis.

Of the 2338 samples tested from patients with ILI, 523 (22%) samples tested positive for influenza virus. The proportion of specimens positive for influenza virus was similar in the three surveillance years (Table 1). The proportion of samples testing positive for influenza varied geographically, with 23% of samples testing positive in the central region, 16% positive from north, and 29% in the south (P < 0.05). Overall, the positivity rate was slightly higher in males (24%) than females (21%) (P < 0.05). The median age of influenza-positive patients was 12 years (range, 1–60 years; IQR = 5–23 years). Children 5–17 years of age were the highest percent positive for influenza among those with ILI (33%) followed by the 18–64 years of age group (28%). Only a small proportion (10%) of children aged <3 years were positive for influenza virus (Table 1). No adults aged 65 years or older tested positive for influenza. Influenza-infected patients were significantly older than those who tested negative for influenza viruses (median, 12 versus 5 years; P < 0.001). There were no differences in age or gender distribution of positive cases when comparing data year to year.

Influenza was detected year-round with the highest proportion of positive specimens in the 3rd and 4th quarter (Figure 3). A bi-modal seasonal pattern was seen in 2008 when there were peaks in September (39% positivity) and again in December (62%), but the sample size was small. In 2009 and 2010, the highest proportion of specimens that were influenza positive occurred in August–September (Figure 3).

Each year, influenza A and B viruses cocirculated (Table 1, Figure 4). In 2008, the dominant type was influenza B. In 2009, influenza A (H3N2) viruses predominated until A (H1N1)pdm09 emerged in July 2009, after which both subtypes cocirculated. In 2010, A (H1N1)pdm09 and influenza B were the most common types identified (Figure 4) and the temporal pattern of influenza seasonality by subtype shows a peak dominated by A (H1N1)pdm09 in weeks 37–43 (Figure 5). One specimen in 2010 was positive for both A (H1N1)pdm09 and influenza B. Nine specimens were subtype A but non-subtypable (two in 2008, seven in 2009). These samples had low viral loads that were below the threshold of detection with subtype-specific reagents. The highest proportion of influenza B specimens were reported from the north of the country (45%) and the highest proportion of A (H1N1)pdm09 from the south (56%).

![Figure 1](https://example.com/figure1.png)  
**Figure 1.** Patients meeting ILI case definition, three central hospitals in Vientiane Capital, 2008–2010. ILI, influenza-like-illness.
Virus isolation was successful for 176 of 268 influenza-positive samples (all sources) (Table 2). Hemagglutination inhibition assays identified seven isolates as “A/Brisbane/59/2007 (H1N1)-like” viruses in 2008. There were no A/H3N2 or B isolates in this year. In 2009 and 2010, with the emergence of A/H1N1pdm09, “A/California/7/2009-like” was isolated in both years. In 2009 and 2010, the majority of (H3N2)-like viruses isolates were “A/Perth/16/2009-like” and the major-
ity of B isolates were “B/Brisbane/60/2008-like.” Only one B isolate was Yamagata lineage, in 2010.

Table 1. The number samples tested and proportion positive for influenza viruses by year, geographical site and demographic characteristics; influenza-like illness (ILI) surveillance, Lao PDR, 2008–2010

| Year       | No. samples tested | No. (%) influenza positive | No. (%)* positive for influenza type/subtype |
|------------|--------------------|----------------------------|---------------------------------------------|
|            |                    |                            | A/H1 | A (H1N1) pdm09 | A/H3 | B | Other** |
| 2008       | 331                | 69 (20-9)                  | 20 (29) | 0 (0)  | 1 (1-4) | 46 (66-7) | 2 (2-9) |
| 2009       | 833                | 184 (22-1)                 | 25 (13-6) | 54 (29-4) | 93 (50-5) | 5 (2-7) | 7 (3-8)** |
| 2010       | 1174               | 270 (23-0)                 | 0 (0) | 143 (53) | 35 (12-9) | 91 (33-7) | 1 (0-4) |
| Total      | 2338               | 523                        | 45 (8-6) | 197 (37-7) | 129 (24-7) | 142 (27-2) | 10 (1-9) |
| Region     |                    |                            |      |            |       |    |        |
| Central     | 1511               | 349 (23-1)                 | 44 (12-6) | 121 (34-7) | 92 (26-4) | 82 (23-5) | 10 (2-9) |
| North       | 516                | 83 (16-1)                  | 1 (1-2) | 25 (30-1) | 20 (24-1) | 37 (44-6) | 0 (0) |
| South       | 311                | 91 (29-3)                  | 0 (0) | 51 (56-0) | 17 (18-7) | 23 (25-3) | 0 (0) |
| Gender      |                    |                            |      |            |       |    |        |
| Male        | 1210               | 291 (24-1)                 | 24 (8-3) | 105 (36-1) | 71 (24-4) | 86 (29-6) | 5 (1-7) |
| Female      | 1127               | 231 (20-5)                 | 20 (8-7) | 92 (39-8) | 58 (25-1) | 56 (24-2) | 5 (2-2) |
| Age group   |                    |                            |      |            |       |    |        |
| 0 to <3     | 782                | 78 (10-0)                  | 7 (9-0) | 37 (47-4) | 15 (19-2) | 17 (21-8) | 2 (2-6) |
| 3 to <5     | 211                | 48 (22-8)                  | 3 (6-3) | 14 (29-2) | 13 (27-1) | 18 (37-5) | 0 (0) |
| 5 to <18    | 543                | 178 (32-8)                 | 14 (7-9) | 69 (38-0) | 45 (25-3) | 48 (27-0) | 2 (1-1) |
| 18 to <65   | 726                | 203 (28-0)                 | 21 (10-3) | 69 (34-0) | 52 (25-6) | 55 (27-1) | 6 (3-0) |
| >65         | 19                 | 0 (0)                      | 0 (0) | 0 (0) | 0 (0) | 0 (0) |        |

*Denominator is total positive specimens.
**Coinfection/untypable.
***Note: a random sample of three specimens that were initially positive but untypable at National Centre for Laboratory and Epidemiology was found to be negative when retested by WHO Collaborating Centre.
the median age of patients with influenza infection was 12 years, and older children were the highest percent positive for influenza among those with ILI (positivity rate 32.8% for 5–<18 years). In neighboring Cambodia, the median age of influenza cases was 9 years old, and children aged 5–14 years were the highest percent positive for influenza (12.1%) among those with ILI. Health-seeking behavior study results from Lao PDR show reported respiratory illnesses to be significantly higher in children aged ≤15 years than in adults (4.9 versus 2.7%; P < 0.001). Other countries in the region, including Thailand, have found a large proportion of hospitalized influenza pneumonia cases are children. We expect that this may also be the case in Lao PDR. Of note, among adults over the age of 65 years, only 19 were tested and none were positive for influenza. Older patients may be less likely to be included in the surveillance because they are often referred immediately to the inpatient department for monitoring.

Influenza-like-illness cases were seen to increase from around May each year confirming that ILI appears before influenza season (July–December), suggesting the contribution of other respiratory pathogens. Our finding of year-round circulation with a peak of influenza activity during the rainy season is similar to Thailand and Northern Vietnam. Many tropical countries, including Vietnam, report a bimodal pattern. This was observed in Lao PDR in
2008 but not following the emergence of A (H1N1)pdm09, after which a single-peak around August to September was observed. Cambodia and Myanmar recently also reported the peak of influenza activity occurring during the wet season, from June to December\textsuperscript{18,19,25} but did not detect influenza year-round. Factors driving seasonality of transmission are not well defined but likely include a combination of climatic conditions, susceptibility of the population, and virus characteristics.\textsuperscript{26,27} As such, influenza activity is likely to vary year to year, and there is a need to collect and analyze multiple years of data to determine trends.

We found variability in the circulating subtypes with a different predominant strain each year. Our results clearly show the transition from A (H1N1)pdm09 into a mix of seasonally occurring viruses. Of interest, seasonal A/H1N1 has not been isolated since the emergence of A (H1N1)pdm09 in July 2009. Influenza B was not seen in quarter three and four of 2009 after emergence of A (H1N1)pdm09, but returned in 2010. Of note, in 2008 and 2009, Lao-circulating influenza strains, when compared to the most recent vaccine recommendations, suggest a closer match to the vaccine strain selection for the northern hemisphere for 2008–09 and 2009–10 respectively (Table 2).\textsuperscript{28} Vaccine strain selection was the same for the northern and southern hemisphere in 2010.\textsuperscript{28} The temporal pattern by subtype for 2010 showed a peak dominated by A (H1N1)pdm09 in the second half of 2010, which resembled the pattern of seasonality of the southern hemisphere.\textsuperscript{29}

Our findings are subject to several limitations. First, our surveillance data focuses only on patients with ILI from sentinel hospital sites, which may not be representative of the entire population of Lao PDR. Furthermore, a convenience sample on 1–2 set days of the week should be taken for testing, but this requirement is not always followed by hospitals (as shown by only 3% of ILI cases in Vientiane Capital having been sampled). As the catchment populations of the sites are unknown, extrapolation of results for estimates of disease burden cannot be made. Nevertheless, samples are collected regularly from all sites and from age-group analysis of central site data, patients from which samples are collected appear to be representative of the ages of all patients with ILI presenting to the sentinel sites. Lao PDR has historically had a limited capacity for surveillance and response, and this surveillance was only recently established and has expanded substantially since it began in 2007. As a result, over 50% of the samples were collected in 2010, data are mostly available from the three central sites (65%), and there is some missing information

\begin{table}[h]
\centering
\caption{Antigenic characterization of influenza virus isolates (all sources: including influenza-like-illness, SARI and outbreaks) in Lao PDR (n = 176)}
\begin{tabular}{|c|c|c|c|c|c|c|}
\hline
Year & Influenza virus A H1N1 & Influenza virus A H3N2 & Influenza virus A pdmH1N1-2009 & Influenza virus B  \\
& Number* & Virus & Number* & Virus & Number* & Virus  \\
\hline
2008** & 20 & “A/Brisbane/59/2007-like” & 1 & “A/Perth/16/2009-like” & 0 & “B/Califonia/7/2009-like” & 46  \\
2009*** & 47 & “A/Brisbane/59/2007-like” & 264 & “A/Perth/16/2009-like” & 295 & “A/Califonia/7/2009-like” & 14  \\
2010* & 0 & “A/Perth/16/2009-like” & 199 & “A/Califonia/7/2009-like” & 110 & “B/Califonia/60/2008-like” & 1  \\
Total & 67 & 311, 62 & 494, 34 & 170, 65 & & &  \\
\hline
\end{tabular}
\end{table}
on age and date of onset. Owing to specimen storage and transport difficulties at provincial sites, specimens are not always received within 24 hour and may arrive in suboptimal condition.

Influenza-like-illness surveillance needs to continue to be strengthened to provide more years of data in order to better understand the epidemiology and impact of influenza in Lao PDR. The relatively recent recognition of seasonal influenza patterns in Lao PDR combined with future data, including from SARI surveillance, will be important for weighing disease burden, both morbidity and mortality, against the costs of advancing influenza vaccine delivery strategy.

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