The Association Between Influenza Treatment and Hospitalization-Associated Outcomes Among Korean Children with Laboratory-Confirmed Influenza

Jacqueline K. Lim  
*International Vaccine Institute*

Tae Hee Kim  
*Asan Medical Center*

Paul E. Kilgore  
*Wayne State University, paul.kilgore@wayne.edu*

Allison E. Aiello  
*University of North Carolina at Chapel Hill*

Byung Min Choi  
*Korea University*

See next page for additional authors

Follow this and additional works at: [https://digitalcommons.wayne.edu/pharm_practice](https://digitalcommons.wayne.edu/pharm_practice)

Part of the Influenza Humans Commons, and the Pharmacy and Pharmaceutical Sciences Commons

**Recommended Citation**

Lim, Jacqueline K.; Kim, Tae Hee; Kilgore, Paul E.; Aiello, Allison E.; Choi, Byung Min; Lee, Kwang Chul; Yoo, Kee Hwan; Song, Young-Hwan; and Kim, Yun-Kyung, "The Association Between Influenza Treatment and Hospitalization-Associated Outcomes Among Korean Children with Laboratory-Confirmed Influenza" (2014). *Department of Pharmacy Practice*. 21.  
[https://digitalcommons.wayne.edu/pharm_practice/21](https://digitalcommons.wayne.edu/pharm_practice/21)

This Article is brought to you for free and open access by the Eugene Applebaum College of Pharmacy and Health Sciences at DigitalCommons@WayneState. It has been accepted for inclusion in Department of Pharmacy Practice by an authorized administrator of DigitalCommons@WayneState.
Authors
Jacqueline K. Lim, Tae Hee Kim, Paul E. Kilgore, Allison E. Aiello, Byung Min Choi, Kwang Chul Lee, Kee Hwan Yoo, Young-Hwan Song, and Yun-Kyung Kim

This article is available at DigitalCommons@WayneState: https://digitalcommons.wayne.edu/pharm_practice/21
The Association between Influenza Treatment and Hospitalization-Associated Outcomes among Korean Children with Laboratory-Confirmed Influenza

Jacqueline K. Lim, Tae Hee Kim, Paul E. Kilgore, Allison E. Aiello, Byung Min Choi, Kwang Chul Lee, Kee Hwan Yoo, Young-Hwan Song, and Yun-Kyung Kim²

¹International Vaccine Institute, Seoul; ²Department of Pediatric Pulmonology & Allergy, Asan Medical Center, Seoul, Korea; ³Department of Pharmacy Practice, Eugene Applebaum College of Pharmacy and Health Sciences, Wayne State University, Detroit, MI; ⁴Department of Epidemiology, Gillings School of Global Public Health, University of North Carolina at Chapel Hill, NC, USA; ⁵Department of Pediatrics, College of Medicine, Korea University, Seoul; ⁶Department of Pediatrics, Inje College of Medicine, Seoul, Korea

Received: 18 August 2013
Accepted: 26 February 2014

Address for Correspondence:
Yun-Kyung Kim, MD
Department of Pediatrics, College of Medicine, Korea University
Korea University Ansan Hospital, 123 Jeokgeum-ro, Danwon-gu, Ansan 425-707, Korea
Tel: +82.10-2248-2330, Fax: +82.31-403-5891
E-mail: byelhana@korea.ac.kr

This study was supported by funding from the Korean Center for Disease Control and Prevention (grant #: 2007-52-E-003), as well as from the governments of Kuwait, Sweden, and the Republic of Korea.

INTRODUCTION

Influenza viruses are common respiratory pathogens among all age groups and are major causes of excess respiratory disease-associated hospitalizations, mortality, and costs during annual epidemics and pandemics (1, 2). Children with underlying medical conditions and younger age, especially ≤ 5 yr olds, are at high risk for severe influenza-associated outcomes (3-5). Furthermore, pediatric influenza exerts a considerable socioeconomic burden in terms of direct and indirect costs, and excess health-care utilization (6-8). Children also serve as a reservoir for household transmission of influenza, causing secondary illness in the family (8, 9).

Currently available neuraminidase inhibitors are effective in reducing influenza-associated illness duration, severity, complication risks, influenza-related mortality and even antibiotic use (10-12). Often patients hospitalized with influenza-related illnesses are inappropriately treated with antibiotics as prophylactic and empiric therapy, as indicated by increased antibiotic prescriptions during the influenza season (10, 13). The emergence of drug resistance and over-prescription of antibiotics are growing public health issues, as well as increased healthcare costs without reductions in illness duration (14-16).

Republic of Korea (ROK) has a relatively high influenza vaccination coverage rate of nearly 40% in the general population (17) and this is higher than the vaccination rate among children hospitalized with acute respiratory infection in the US (18) and similar to that of healthy Ontario children in Canada (19). Nonetheless, Korea still suffers from considerable disease burden of influenza in children, where 20% of viral respiratory disease hos-
Hospitalizations are due to influenza virus infection (20). This study is the first to address treatment patterns in influenza-associated hospitalizations among Korean children. To address existing gaps in knowledge regarding childhood influenza in Korea, we evaluated the association between hospitalization duration and treatment type, using clinical and epidemiological data from children hospitalized with laboratory-confirmed influenza. We also described the economic burden by calculating medical charges incurred during hospitalization by treatment type.

MATERIALS AND METHODS

Study hospital and cohort
A retrospective review of medical records for patients admitted or treated from February 2004 through June 2007 was conducted at three academic, tertiary-care hospitals. Ansan, Anam, and Guro Hospitals, affiliated with Korea University (KU), serve the communities of Ansan city (a neighboring city of Seoul), and districts of Anam and Guro in Seoul, respectively. At these KU hospitals, nasal aspirate specimens were systematically collected from patients presenting with acute respiratory symptoms and routinely tested to identify adenovirus, parainfluenza, respiratory syncytial virus (RSV) and influenza A/B with virus. Additionally, selected patients were tested for influenza A/B and RSV by rapid antigen detection (Directigen EZ Flu A+B Test Kit, Becton-Dickinson, NJ, USA) at the clinicians’ discretion.

For this study, we included patients who: a) presented with respiratory symptoms at one of the KU hospitals, b) were ≤ 15 yr-of-age at the time of admission, and c) had laboratory-confirmation of influenza A/B. Children with mixed viral infections (with following viruses; influenza, RSV, parainfluenza, or adenovirus) and with recurrent episodes with the same virus type who returned to the hospital within two weeks from the date of the previous discharge or clinic visit were excluded from our analysis. Assembly of the study patient cohort is outlined in Fig. 1. During the 41 months, 1,039 episodes of laboratory-confirmed influenza were identified from the outpatient and inpatient departments of the three hospitals.

Virus detection
Viral culture had been performed using an enhanced cell culture method with fluorescent antibody detection. Specimens (200 μL) were inoculated onto R-Mix Ready Cells (Diagnostic Hybrids, NY, USA), and vials were centrifuged at 700 × g for 60 min at room temperature. After overnight incubation at 35°C in a CO2 incubator, cell monolayers were washed with phosphate-buffered saline, fixed with acetone, and stained with a respiratory virus fluorescent antibody pool, D3 DFA (Diagnostic Hybrids). When virus specific fluorescence was noted, virus identification was performed by using individual monoclonal antibodies (Diagnostic Hybrids). The presence of three or more cells per well with specific apple-green fluorescence was considered positive identification. Screenings were done on day 1 and 3. Additionally, selected patients were tested for influenza A/B and RSV by rapid antigen detection (Directigen EZ Flu A+B Test Kit, Becton-Dickinson, NJ, USA) at the clinicians’ discretion.

Data collection
Via medical chart reviews of 1,039 episodes of community-acquired laboratory-confirmed influenza, we extracted the following demographic and clinical variables from the KU hospital electronic medical information database: influenza virus

![Flow chart](http://jkms.org)

Fig. 1. Flow chart describing the ascertainment of the patient cohort with hospitalization identified during the study period of February 2004-June 2007.
type, dates of admission and discharge, diagnoses, signs and symptoms, body temperature, duration of fever, pre-existing medical conditions, vital signs, hematologic and radiologic test results, treatment information, and discharge status. Direct hospital medical charges were obtained from the hospital billing/registry office. Treatment-related fee was composed of charges for injection, medication, treatment material, physical therapy, surgery, anesthesia, and blood infusion. Testing fee consisted of charges for routine and specially-ordered laboratory and diagnostic tests. Hospital admission fee included admission, food, and room charges. They were summed for the total hospital charges incurred and currency was converted from Korean Won (KRW) to US dollar with the average exchange rate during study period of 1 KRW = 0.0007 US$.

The diagnosis at admission was categorized by the primary clinical manifestation. Diagnoses of pneumonia, bronchitis, croup, and asthma were grouped as lower respiratory tract illness (LRTI), while acute pharyngitis, sinusitis, and laryngitis were grouped as upper respiratory illness (URI). Urinary tract infection, sepsis, and neurologic (e.g., convulsions, febrile seizure, Guillain-Barré Syndrome), gastrointestinal (e.g., diarrhea, acute gastroenteritis), and cardiovascular conditions were grouped as ‘non-respiratory’ diseases. An axillary (tympanic membrane) temperature ≥ 37.5 (37.8)°C was defined as fever. Fever duration included the febrile days prior to admission.

Variable construction and statistical analysis
The analysis examined the relationship between the length of hospital stay (LOS) and the different types of treatment administered during hospitalization, when adjusted for potential confounding factors. Nosocomial influenza episodes with virus detection ≥ 7 days after the date of admission were excluded from the analysis sample. In accordance with current guidelines of influenza treatment, episodes treated with oseltamivir (Tamiflu®, Roche Pharmaceuticals, Basel, Switzerland) outside of the recommended two-day window after the onset of symptoms, and episodes treated with amantadine were excluded.

The mean peak body temperature for study patients measured during hospitalization, 38.6°C, was used as the cutoff value to create a dichotomous variable indicating elevated body temperature. Normal white blood cell (WBC) count was defined as WBC of 5,000-19,500/µL for children ≤ 10 months-of-age; 6,000-17,500/µL for those 11 months-2 yr-of-age; 5,500-15,500/µL for children 3-7 yr-of-age; and 4,500-13,500 (11,500)/µL for children 8-13 (14-15) yr-of-age (21). Leukopenia and leukocytosis were below the lower and above the upper values of the normal WBC range, respectively. Complications that occurred during hospitalization were categorized into secondary bacterial pneumonia, other secondary bacterial infection, encephalitis, or exacerbation of the pre-existing medical conditions. Pre-existing medical conditions were congenital diseases and acquired diseases (e.g., asthma, and cancer). Binary variables were created for the presence or absence of complications and pre-existing conditions.

Influenza hospitalizations with complete data were divided into four treatment categories: oseltamivir-only, antibiotics-only, antibiotics-plus-oseltamivir, and supportive-care without antiviral or antibiotic treatment. Categorical pair-wise comparisons were made with reference to the oseltamivir-only group using chi-square or the Fisher’s exact tests with significance at P < 0.05. Comparison of continuous variables was performed using the Student’s t-test. However, for hospital charges with skewed distributions, the Mann-Whitney test was used to make pair-wise comparisons with reference to the oseltamivir-only group. Bivariate analyses were performed to identify potential confounders and covariates of interest for the regression model, using SAS® version 9.0 (SAS Institute, Cary, NC, USA).

To adjust for variability in initial severity of illness, clinical parameters such as temperature measured at admission, presenting signs and symptoms (i.e., expectoration, diarrhea, headache, injected tympanic membrane, sore throat, vomiting, breathing difficulty, rales or wheezing), admission diagnosis, laboratory test results (i.e. WBC/neutrophil counts and chest infiltrates) were compared across treatment groups. Construction of the model with these variables allows statistical adjustment for initial variability in clinical presentation. As potential confounders, presence of underlying medical conditions and occurrence of complications during hospitalization, were included as dichotomous variables in the model to control for variations in disease presentation at admission and during clinical course that may have guided treatment-related decisions. Some variables that are closely related were assessed for multicollinearity and were limited for inclusion in the regression model. Multivariable linear regression was performed with the outcome as the log-transformed, hospitalization duration and categorical variable of influenza treatments as the main predictor. The β, % change in treatment-related charges, and 95% confidence intervals (CI) were reported.

Ethics statement
Study approvals were obtained from the institutional review boards of the KU Ansan Hospital (IRB No. ED0744 ) and the International Vaccine Institute (IRB No. 2007-008). The exemption of informed consent and assent forms was allowed by both IRBs because this study was based on a retrospective review and the patient IDs were removed from the DB provided by the hospital records office.

RESULTS

Patient characteristics
Of the 770 hospitalizations in our analysis sample, 77% were ≤
Clinical characteristics
Among the total of 770 patients, 27 were treated with oseltamivir-only, 620 were treated with antibiotics-only, 67 received antibiotics-plus-oseltamivir and 56 patients received supportive care. The mean LOS for patients in the antibiotics-only and antibiotics-plus-oseltamivir groups were significantly longer compared to the oseltamivir-only group (5.0 and 4.0 vs 3.0 days, respectively) (Fig. 2). The mean fever duration was significantly longer in the antibiotics-only group compared to the oseltamivir-only group (5.5 vs 3.8 days). Patients in the oseltamivir-only group were more likely to have received influenza rapid diagnostic tests (IRDT) than the antibiotics-only, antibiotics-plus-oseltamivir, and supportive-care groups (82% vs 3%, 55%, and 16%, respectively) (Table 2). Patients who presented with rales/wheezing were found more commonly in those three treatment groups than in the oseltamivir-only group (38%, 37%, and 41% vs 7%). Patients presenting with pharyngeal injection were significantly less common in the antibiotics-only group, compared to the oseltamivir-only group (73% vs 93%). Overall, 12% of patients (n = 96/770) suffered from complications during hospitalization, and 45% (n = 43/96) developed secondary bacterial pneumonia or other bacterial infection with few cases of neurologic problems (e.g. encephalitis) and exacerbation of pre-existing medical conditions (data not shown).

Hospitalization-associated medical charges
The mean treatment-related charges (US$171 and $111 vs $67, \(P < 0.001\) and < 0.001, respectively) and the hospital admission charges (US$287 and $240 vs $176, \(P < 0.001\) and 0.017, respectively) were significantly higher in the groups treated with antibiotics-only and antibiotics-plus-oseltamivir, compared to the group treated with oseltamivir-only (Table 3). The mean total charges for hospitalizations treated with antibiotics-only and antibiotics-plus-oseltamivir were significantly higher than charges for hospitalizations treated with oseltamivir-only (US$753 and $623 vs $508, \(P < 0.001\) and 0.059, respectively).

Predictors of increased hospital stay
Compared with the oseltamivir-only group, children treated with antibiotics-only showed 44.9% longer hospitalization compared to those who received oseltamivir-only (95% CI, 15.85-81.33; \(P = 0.001\)) (Table 4) in multivariate analysis. Treatment with antibiotics-plus-oseltamivir also showed a significantly longer LOS, compared to the oseltamivir-only group, by

Table 1. Characteristics of the analysis cohort of hospitalizations with laboratory-confirmed influenza identified during February 2004–June 2007 in Republic of Korea

| Treatment group | No. | Mean age, (yr ± SD) | Female | Influenza virus type | Admission diagnosis |
|-----------------|-----|---------------------|--------|---------------------|---------------------|
|                 |     |                     |        |                     |                     |
| Oseltamivir-only (reference) | 27  | 3.58 ± 3.91         | 12 (44.4) | 13 (48.2) | 4 (14.8) | 5 (18.5) | 12 (44.4) | 10 (37.0) |
| Antibiotics-only | 620 | 3.39 ± 2.76         | 264 (42.6) | 386 (62.3)* | 225 (36.3) | 354 (67.1)* | 184 (29.7) | 82 (13.2) |
| Antibiotics-plus-oseltamivir | 67  | 3.51 ± 2.90         | 27 (40.3) | 43 (64.2)† | 15 (22.4) | 28 (41.8)* | 29 (43.3) | 10 (14.9) |
| Supportive-care | 56  | 3.77 ± 3.36         | 17 (30.4) | 36 (64.3)‡ | 20 (35.7) | 30 (63.6)* | 15 (26.8) | 11 (19.6) |
| Total            | 770 | 3.43 ± 2.86         | 320 (41.6) | 478 (62.1) | 264 (34.3) | 417 (64.2) | 240 (31.2) | 113 (14.7) |

Values are No. (%) unless otherwise noted. *P < 0.001, for comparison between the antibiotics-only vs oseltamivir-only treatment groups; †P = 0.050, for comparison between the antibiotics-plus-oseltamivir vs oseltamivir-only treatment groups; ‡P < 0.001, for comparison between the antibiotics-only vs oseltamivir-only treatment groups; §P = 0.010, for comparison between the supportive-care vs oseltamivir-only treatment groups. LRTI, lower respiratory tract illness; URI, upper respiratory illness.
Table 2. Clinical characteristics of the 770 hospitalizations with laboratory-confirmed influenza treated with antiviral or antibiotic medication from February 2004 to June 2007

| Characteristics                              | Treatment                      |
|----------------------------------------------|-------------------------------|
|                                              | Oseltamivir-only (reference)  | Antibiotics-only             | Antibiotics plus oseltamivir | Supportive-care |
| Temperature at admission (°C) mean (SD)      | 37.65 (0.85)                  | 37.54 (0.98)                 | 37.79 (0.89)                 | 37.25 (0.80)    |
| Peak temperature during hospitalization (°C) mean (SD) | 38.50 (0.69)                  | 38.61 (0.86)                 | 38.87 (0.89)                 | 38.13 (0.99)    |
| Received influenza rapid antigen detection test | 22 (81.5)                     | 21 (3.4)*                    | 37 (55.2)*                   | 9 (16.1)*       |
| Signs                                        |                               |                               |                               |                |
| Pharyngeal injection                         | 25 (92.6)                     | 450 (72.7)*                  | 52 (77.6)                    | 43 (76.8)       |
| Pharyngeal injection                         | 2 (7.4)                       | 237 (38.2)*                  | 25 (37.3)*                   | 23 (41.1)**     |
| Pharyngeal injection                         | 3 (11.1)                      | 60 (9.7)                     | 7 (9.5)                      | 4 (5.4)         |
| Symptoms                                     |                               |                               |                               |                |
| Cough                                        | 22 (81.5)                     | 539 (86.9)                   | 59 (88.1)                    | 50 (89.3)       |
| Rhinorrhea                                   | 15 (55.6)                     | 322 (51.9)                   | 39 (58.2)                    | 38 (67.9)       |
| Expectoration                                | 14 (51.9)                     | 357 (57.6)                   | 34 (50.8)                    | 28 (50.0)       |
| Diarrhea                                     | 5 (18.5)                      | 60 (9.7)                     | 1 (1.5)*                     | 8 (14.3)        |
| Headache                                     | 6 (22.2)                      | 26 (4.2)                     | 3 (4.5)                      | 3 (5.4)         |
| Sore throat                                  | 0                             | 49 (7.9)                     | 3 (4.5)                      | 16 (28.6)*      |
| Vomiting                                     | 12 (44.4)                     | 143 (23.1)*                  | 16 (23.9)*                   | 10 (17.9)*      |
| Breathing difficulty                         | 1 (3.7)                       | 32 (5.2)                     | 2 (3.0)                      | 2 (3.6)         |
| White blood cell count                       |                               |                               |                               |                |
| Leukopenia                                   | 5 (18.5)                      | 123 (19.8)                   | 13 (19.4)                    | 18 (32.1)       |
| Normal                                       | 20 (74.1)                     | 465 (75.0)                   | 52 (77.6)                    | 36 (64.3)       |
| Leukocytosis                                 | 2 (7.4)                       | 32 (5.2)                     | 2 (3.0)                      | 2 (3.6)         |
| Neutrophil (ref. normal, 54%-62%)            |                               |                               |                               |                |
| Outside normal range                         | 26 (96.3)                     | 541 (87.3)                   | 59 (88.1)                    | 49 (87.5)       |
| Chest radiograph infiltrates                 | 5 (18.5)                      | 194 (31.3)                   | 10 (14.9)                    | 17 (30.4)       |
| Presence of complications                    | 3 (11.1)                      | 80 (12.9)                    | 4 (6.3)                      | 9 (16.1)        |
| Presence of underlying diseases              | 1 (3.7)                       | 53 (8.6)                     | 5 (7.5)                      | 7 (12.5)        |

Values are No. (%) unless otherwise noted. The 95% confidence intervals (95% CI) were reported. *P < 0.001, for comparison between the antibiotics-only vs oseltamivir-only treatment groups; †P = 0.17, for comparison between the antibiotics-plus-oseltamivir vs oseltamivir-only treatment groups; ‡P < 0.001, for comparison between the supportive-care vs oseltamivir-only treatment groups; §P = 0.022, for comparison between the antibiotics-only vs oseltamivir-only treatment groups; ‡‡P = 0.001, for comparison between the antibiotics-only vs supportive-care treatment groups; ††P = 0.004, for comparison between the antibiotics-plus-oseltamivir vs oseltamivir-only treatment groups; †‡P = 0.02, for comparison between the antibiotics-plus-oseltamivir vs oseltamivir-only treatment groups; ‡‡‡P = 0.002, for comparison between the supportive-care vs oseltamivir-only treatment groups; †††P = 0.002, for comparison between the supportive-care vs oseltamivir-only treatment groups; ††¶P = 0.001, for comparison between the antibiotics-only vs oseltamivir-only treatment groups; ††††P = 0.049, for comparison between the antibiotics-plus-oseltamivir vs oseltamivir-only treatment groups; †¶¶P = 0.010, for comparison between the supportive-care vs oseltamivir-only treatment group.

Table 3. Hospital charges (mean, SD) of patients with laboratory-confirmed influenza by treatment type, February 2004-June 2007

| Type of hospital charge          | Treatment type                      |
|----------------------------------|-------------------------------------|
|                                  | Oseltamivir-only (reference)        | Antibiotics-only            | Antibiotics plus oseltamivir | Supportive-care |
| Treatment-related                | 66.55 (37.39)                       | 170.67 (113.34)*            | 110.98 (60.28)*              | 78.41 (77.69)  |
| Hospital admission               | 175.76 (92.45)                      | 287.16 (335.04)*            | 240.23 (158.49)*             | 212.60 (112.64) |
| Testing                          | 230.35 (156.88)                     | 236.78 (193.89)             | 226.88 (138.77)              | 227.25 (167.84) |
| Total                            | 507.82 (208.36)                     | 753.47 (1247.68)*           | 623.18 (283.89)              | 556.58 (248.91) |

Mean values and standard deviation (in parenthesis) are reported in US$ with exchange rate of 1 KRW = 0.0007 US$. *P < 0.001, for comparison of the mean treatment-related fees in antibiotics-only vs the oseltamivir-only treatment groups; †P < 0.001, for comparison of the mean treatment-related fees in antibiotics-plus-oseltamivir vs the oseltamivir-only treatment groups; ‡P = 0.001, for comparison of the mean hospital admission charges in antibiotics-only vs the oseltamivir-only treatment groups; ‡‡P = 0.017, for comparison of the mean hospital admission charges in antibiotics-plus-oseltamivir vs the oseltamivir-only treatment groups; ††P < 0.001, for comparison of total hospital charges in antibiotics-only vs the oseltamivir-only treatment groups.

28.2% (95% CI, 1.96-61.11; P-value = 0.034). From the univariate analysis, patients who received the antibiotics-only therapy showed 55.8% longer hospitalization compared to those who received oseltamivir-only (95% CI, 27.99-89.66; P-value < 0.001) (data not shown). Patients who received antibiotics-plus-oseltamivir therapy also showed a significantly longer LOS by 28.5%, compared to the oseltamivir-only group (95% CI, 2.24-61.48; P-value = 0.032).

Children in the 3-5 yr-old groups showed to have 12.6% shorter LOS, compared to infants ≤ 1 yr old (95% CI, -21.9 to -2.18; P-value = 0.019). Patients with dyspnea had 18.0% longer LOS compared to those without dyspnea (95% CI, 0.28-38.86; P-value = 0.046), while those children with admission diagnosis categorized as URI showed 9.3% shorter LOS compared to those with LRTI (95% CI, -17.52 to -0.29; P-value = 0.044). Also, the presence of chest radiograph infiltrates and pre-existing medical conditions were significant indicators of lengthened hospitalization by 12.9% and 21.3%, respectively (95% CI, 3.50-23.04 and 6.27-38.32, respectively).
Our results are consistent with findings from Kaiser et al. and other treatments. IRDT in addition to culture tests than patients who received 4% received oseltamivir-only. Patients who received oseltamivir talizations, 81% of patients received antibiotic-only while only in influenza treatment options. In our sample of influenza hospi among patients treated with oseltamivir-only among different hospital admission and treatment-related charges, occurred showed that the lowest mean total hospital charges, as well as children with laboratory-confirmed influenza. Also, our data pation to hospitalization-associated outcomes among Korean This is the first study to compare influenza treatments in rela-

duction of fever in oseltamivir-treated influenza A patients (12). Our data showed a significantly longer mean LOS for patients in the antibiotics-only and antibiotics-plus-oseltamivir groups compared to the oseltamivir-only group. Less timely onset of antibiotic therapy could play a role in increased LOS, however, 99% (n = 683) of 687 antibiotic-treated patients received antibiotics within 3 days of admission in our data, indicating the timing of antibiotic therapy was unlikely the cause of observed association. In the antibi-
significantly longer mean LOS for patients in the antibiotics-only and antibiotics-plus-oseltamivir groups compared to the oseltamivir-only group. Less timely onset of antibiotic therapy could play a role in increased LOS, however, 99% (n = 683) of 687 antibiotic-treated patients received antibiotics within 3 days of admission in our data, indicating the timing of antibiotic therapy was unlikely the cause of observed association. In the antibiotic-plus-oseltamivir group, 59 patients out of 67 had received antibiotics before oseltamivir in our data. While this may be explained by the fact that these patients may have had clinical indications of bacterial pneumonia or other secondary infections, this antibiotic prescription preceding oseltamivir treatment could have led to delayed onset of oseltamivir therapy and consequently lengthening the patients’ LOS.

Several factors, indicating severe clinical course (e.g., complica-

tion) and initial differences in clinical presentation of illness,
could have potentially influenced treatment-related decisions and were taken into account in our multivariate modeling. It is possible that additional confounding variables that were not measured in this study may explain the relationship. However, the observed strength of the relationship of oseltamivir-only compared to antibiotics-only and antibiotics-plus-oseltamivir groups (i.e., 45% and 28% increases in LOS with the P value of 0.001 and 0.034, respectively) makes it unlikely that our findings are a result of residual confounding not controlled for in our models. While these findings are consistent with several reports of reduced hospitalization associated with oseltamivir treatment (10, 22), no previous studies have shown that antibiotic treatments, compared to oseltamivir therapy, is associated with longer LOS in children with laboratory-confirmed influenza.

Our finding that hospital charges were lowest among oseltamivir-only treated children also corresponds with previous studies reporting that unnecessary costs may incur among influenza-like illness (ILI) patients treated with antibiotics (13, 16, 23) while oseltamivir is the cost-effective choice of influenza treatment (24). The cost of influenza-related hospitalization in our data was shown to be lower than previously estimated in other countries (6, 7, 25). This is explained by the fact that ROK has universal population coverage by the National Health Insurance that subsidizes physician fees, hospital and prescription drugs charges via a compulsory healthcare plan (26). Despite the number of studies that have previously reported an indiscriminant and inappropriate use of antibiotics (16, 27, 28), antibiotics were demonstrated to still be a primary treatment option for children hospitalized with influenza in ROK. Observed treatment patterns may be explained, in part, by the absence of standardized treatment guidelines for seasonal influenza in ROK.

In addition to the absence of the treatment guidelines, infrequent use of the IRDT during the study period could lead to observed treatment patterns. Without test for viral pathogen, the uncertainty of diagnosis of children with influenza-like illness frequently leads clinicians to prescribe antibiotics to the children with lower respiratory symptoms and signs. Our data showed that oseltamivir-treated children were more likely to have been treated with IRDT compared to the children treated with antibiotics or supportive-care. Although all of our study subjects were ultimately proven to have laboratory-confirmed influenza, children who were treated with antibiotics or only supportive-care may have been initially suspected of having bacterial infection or non-viral respiratory illness. Such children were less likely to undergo testing by IRDT than those oseltamivir-treated, who were more likely to be confirmed with influenza on admission with IRDT. Our results as well as previous studies support the notion that IRDT are an important tool for guiding influenza treatment decision-making (29, 30). Distinguishing influenza from other respiratory diseases in children as well as identifying influenza with atypical clinical presentations is a well-recognized clinical challenge (31). As a result, increased accessibility of IRDT is a recognized priority among clinicians in ROK and elsewhere that is likely to improve patient treatment regimens and clinical outcomes, as well as potentially reduce antibiotic usage and additional laboratory testing (29, 30, 32).

In our study, we adjusted for clinical characteristics in our multivariable analysis that were previously linked to higher influenza viral load, prolonged hospitalization, secondary pneumonia, as well as other viral or bacterial complications (33-38). We found no consistent treatment-related patterns with respect to disease severity of illness when compared clinical presentation by treatment type. For example, rales/wheezing were more commonly present among patients treated with antibiotics-only, antibiotics-plus-oseltamivir, and even supportive-care, compared to those treated with oseltamivir-only. Also, chest infiltrate, which may suggest bacterial pneumonia (39), was not significantly more common in antibiotics or antibiotics-plus-oseltamivir groups compared to the oseltamivir-only group. There was no currently available and validated severity index for pediatric influenza.

In this retrospective study, our data extraction was limited to the information available on medical charts. Due to the observational nature of our study, we cannot conclude that oseltamivir shortened the LOS among hospitalized patients with influenza. Although lengthened hospitalization practically directly reflects the burden of influenza requiring hospitalization, clinical data may not entirely capture the influenza disease burden. In clinical practice at KU hospitals, the hospitalizations diagnosed with mild conditions (e.g., pharyngitis) were less likely to be treated with antibiotics, while those with more severe findings, such as rales or chest infiltrate that prompt clinicians to suspect bacterial pneumonia, were often treated with antibiotics. Thus, variability in illness severity at admission influencing clinical diagnosis was the most important confounder of this relationship and various clinical attributes were controlled for in the analysis.

Despite these limitations, a robust analysis of the treatment type was possible due to: 1) three complete and consecutive viral seasons; 2) a relatively large sample of laboratory-confirmed influenza hospitalizations; 3) the geographic and patient mix of patients from community and academic hospitals in different parts of ROK, providing a representative sample for greater generalizability of results; and 4) no oseltamivir resistance in the viral strains during our study period in ROK (anecdotal reports), minimizing the influence of oseltamivir resistance on the study outcomes. Prescription patterns could differ by clinician and facility, but the three hospitals under the umbrella of KU had highly uniform standards in patient management and treatment regimen. Moreover, all subjects were culture-confirmed, in ad-
ndition to some that had undergone IRDT, ensuring uniformity in the levels of sensitivity and specificity of influenza diagnostics. Considering the lack of data from direct comparison between antibiotic and oseltamivir treatments, our study provides a realistic illustration of influenza treatment in relation to increased hospitalization by comparing treatment options used in clinical practice in ROK. As suggested by Falagas et al., there is a need for comparative and randomized studies [40], our study findings will serve as a platform for further assessment of different influenza treatment options in clinical practice.

In ROK, the predominantly circulating strains were A/H3N2 in 2004-05, A/H1N1 in 2005-06, and A/H3N2 in 2006-07 and there were no reported oseltamivir-resistant strains of influenza during the study period of 2004-07. Notably, in the January 2011 issue of the U.S. Centers for Disease Control and Prevention’s Morbidity and Mortality Weekly Report, either oseltamivir or zanamivir was recommended for use in patients hospitalized with suspected or confirmed influenza, whether it is 2009 H1N1 virus, influenza A (H3N2) virus, or influenza B virus or when the influenza virus type or influenza A virus subtype is unknown [41]. In ROK, there is limited number of oseltamivir treatment guidelines or recommendations for seasonal influenza. Also, IRDT is not routinely used in decision-making of patient management, despite its well-known benefits [23, 32, 37]. Lately, real time RT-PCR is used more often in many facilities in ROK, which means influenza infection could be detected in earlier stage of illness with better accuracy.

Recognizing that laboratory-confirmation of influenza at an earlier point of the course of hospitalization may be particularly important in pediatric influenza treatment [29, 42], our study suggests the need for appropriate use of IRDT and RT-PCR for seasonal influenza in ROK so that more vigilant and judicious antibiotics prescription could be practiced in the clinical setting.

ACKNOWLEDGMENTS

We thank the doctors and laboratory staff of Korea University’s Ansan Hospitals for their participation in the retrospective pediatric influenza study in ROK. We thank collaborators at Myung Moon Pediatrics for their support, Dr. Betsy Foxman (School of Public Health at the University of Michigan, Ann Arbor), Dr. Bill Letson (Pediatric Dengue Vaccine Initiative), Ms. Sunheang Shin, and Ms. Deborah Hong, as well as the statisticians and administrative staff at the International Vaccine Institute for their helpful comments during the analysis and preparation of this manuscript.

DISCLOSURE

The authors have no relevant financial relationships or potential conflicts of interest to disclose regarding the material discussed in this manuscript.

ORCID

Jacqueline K. Lim http://orcid.org/0000-0001-6744-7825
Byung Min Choi http://orcid.org/0000-0003-9831-2353
Kwang Chul Lee http://orcid.org/0000-0003-3552-8721
Kee Hwan Yoo http://orcid.org/0000-0001-6490-4293
Young-Hwan Song http://orcid.org/0000-0001-6355-9440
Yun-Kyung Kim http://orcid.org/0000-0003-4396-8671

REFERENCES

1. Molinari NA, Ortega-Sanchez IR, Messonnier ML, Thompson WW, Worley PM, Weintraub E, Bridges CB. The annual impact of seasonal influenza in the US: measuring disease burden and costs. Vaccine 2007; 25: 5086-96.
2. Newall AT, Scuffham PA. Influenza-related disease: the cost to the Australian healthcare system. Vaccine 2008; 26: 6818-23.
3. Neuzil KM, Wright PF, Mitchel EF Jr, Griffin MR. The burden of influenza illness in children with asthma and other chronic medical conditions. J Pediatr 2000; 137: 856-64.
4. Neuzil KM, Zhu Y, Griffin MR, Edwards KM, Thompson JM, Tolleson SJ, Wright PF. Burden of interpandemic influenza in children younger than 5 years: a 25-year prospective study. J Infect Dis 2002; 185: 147-52.
5. Bhat N, Wright JG, Broder KR, Murray EL, Greenberg ME, Glover MI, Likos AM, Posey DL, Klimov A, Lindstrom SE, et al. Influenza-associated deaths among children in the United States, 2003-2004. N Engl J Med 2005; 353: 2559-67.
6. Ampofo K, Gesteland PH, Bender J, Mills M, Daly J, Samore M, Byington C, Pavia AT, Srivastava R. Epidemiology, complications, and cost of hospitalization in children with laboratory-confirmed influenza infection. Pediatrics 2006; 118: 2409-17.
7. Keren R, Zaoutis TE, Sadekemire S, Luan XQ, Coffin SE. Direct medical cost of influenza-related hospitalizations in children. Pediatrics 2006; 118: e1321-7.
8. Neuzil KM, Hohlbein C, Zhu Y. Illness among schoolchildren during influenza season: effect on school absenteeism, parental absenteeism from work, and secondary illness in families. Arch Pediatr Adolesc Med 2002; 156: 986-91.
9. Heikkinen T, Silvennoinen H, Peltola V, Ziegler T, Vainionpaa R, Vuo rinen T, Kainulainen L, Puhakkka T, Jarri T, Toikka P, et al. Burden of influenza in children in the community. J Infect Dis 2004; 190: 1561-66.
10. Kaiser L, Wat C, Mills T, Mahoney P, Ward P, Hayden F. Impact of oseltamivir treatment on influenza-related lower respiratory tract complications and hospitalizations. Arch Intern Med 2003; 163: 1667-72.
11. Nicholson KG, Aoki FY, Osterhaus AD, Trottier S, Carewicz O, Mercier CH, Rode A, Kinnersley N, Ward P. Efficacy and safety of oseltamivir in treatment of acute influenza: a randomised controlled trial: Neuraminidase Inhibitor Flu Treatment Investigator Group. Lancet 2000; 355: 1845-50.
12. Sato M, Saito R, Sato I, Tanabe N, Shobugawa Y, Sasaki A, Li D, Suzuki Y, Sato M, Sakai T, et al. Effectiveness of oseltamivir treatment among chil-
dren with influenza A or B virus infections during four successive winters in Niigata City, Japan. Tohoku J Exp Med 2008; 214: 113-20.

13. Neuzil KM, Mellen BG, Wright PF, Mitchel EF Jr, Griffin MR. The effect of influenza on hospitalizations, outpatient visits, and courses of antibiotics in children. N Engl J Med 2000; 342: 225-31.

14. Carrat F, Schwarzer M, Houssset B, Valleron AJ. Antibiotic treatment for influenza does not affect resolution of illness, secondary visits or lost workdays. Eur J Epidemiol 2004; 19: 703-5.

15. Chidiac C, Maulin L. Using antibiotics in case of influenza. Med Mal Infect 2006; 36: 181-9.

16. Gao B, Li R, Liu YM, Cao ZX, Geng XQ, Lau LT, Lu J, Wu L, Cai SF, Bai RT, et al. The impact of antibiotic treatment in patients with influenza-like illness. Zhonghua Jie He He Hu Xi Za Zhi 2008; 31: 483-7.

17. Kee SY, Lee JS, Cheong HJ, Chou BC, Song JY, Choi WS, Jo YM, Seo YB, Kim WI. Influenza vaccine coverage rates and perceptions on vaccination in South Korea. J Infect 2007; 55: 273-81.

18. Poehling KA, Edwards KM, Weinberg GA, Szilagyi P, Staat MA, Iwane KM, et al. Influenza burden in febrile infants: a Korean case. Health Policy 2005; 74: 133-45.

19. Cao B, Li R, Liu YM, Cao ZX, Geng XQ, Lau LT, Lu J, Wu L, Cai SF, Bai RT, et al. The underrecognized burden of influenza in young children. N Engl J Med 2006; 355: 31-40.

20. Poehling KA, Edwards KM, Weinberg GA, Szilagyi P, Staat MA, Iwane KM, et al. Influenza vaccine coverage rates and perceptions on vaccination in South Korea. J Infect 2007; 55: 273-81.

21. Pee SY, Choi Y, Lee GC, Lee JW, Kilgore PE. Burden of viral respiratory disease hospitalizations among children in a community of Seoul, Republic of Korea, 1995 - 2005. Scand J Infect Dis 2008; 40: 946-53.

22. Gums JG, Pelletier EM, Blumentals WA. Oseltamivir and influenza-related complications, hospitalization and healthcare expenditure in healthy adults and children. Expert Opin Pharmacother 2008; 9: 151-61.

23. Ciesla G, Leader S, Stoddard J. Antibiotic prescribing rates in the US ambulatory care setting for patients diagnosed with influenza, 1997-2001. Respir Med 2004; 98: 1093-101.

24. Reisinger K, Greene G, Aultman R, Sander B, Gyldmark M. Effect of influenza treatment with oseltamivir on health outcome and costs in otherwise healthy children. Clin Drug Invest 2004; 24: 395-407.

25. Hassan F, Lewis TC, Davis MM, Gebremariam A, Dombkowski K. Impact of influenza rapid diagnostic tests (IRDT) on the diagnosis of influenza and on the management of influenza in children in ambulatory pediatric setting. Arch Pediatr 2009; 16: 288-93.

26. Reisinger K, Greene G, Aultman R, Sander B, Gyldmark M. Impact of influenza rapid diagnostic tests (IRDT) on the diagnosis of influenza and on the management of influenza in children in ambulatory pediatric setting. Arch Pediatr 2009; 16: 288-93.

27. Peltola V, Ziegler T, Ruuskanen O. Influenza A and B virus infections in children. Clin Infect Dis 2003; 36: 299-305.

28. Low D. Reducing antibiotic use in influenza: challenges and rewards. Clin Microbiol Infect 2008; 14: 298-306.

29. Noyola DE, Demmuler GJ. Effect of rapid diagnosis on management of influenza A infections. Pediatr Infect Dis J 2000; 19: 303-7.

30. De La Rocque F, Lécuyer C, Wollner C, d’Athys P, Pecking M, Thollot F, Cohen R. Impact of influenza rapid diagnostic tests (IRDT) on the diagnosis of influenza and on the management of influenza in children in ambulatory pediatric setting. Arch Pediatr 2009; 16: 288-93.

31. Poehling KA, Edwards KM, Weinberg GA, Szilagyi P, Staat MA, Iwane KM, et al. Influenza burden in febrile infants: a Korean case. Health Policy 2005; 74: 133-45.

32. Bonner AB, Monroe KW, Talley LI, Klasner AE, Kimberlin DW. Impact of the rapid diagnosis of influenza on physician decision-making and patient management in the pediatric emergency department: results of a randomized, prospective, controlled trial. Pediatrics 2003; 112: 363-7.

33. Eriksson M, Bennett R, Nilsson A. Wheezing following lower respiratory tract infections with respiratory syncytial virus and influenza A in infancy. Pediatr Allergy Immunol 2000; 11: 193-7.

34. Heymann PW, Carper HT, Murphy DD, Platts-Mills TA, Patrie J, McLaughlin AP, Erwin EA, Shaker MS, Hellem M, Peerzada J, et al. Viral infections in relation to age, atopy, and season of admission among children hospitalized for wheezing. J Allergy Clin Immunol 2004; 114: 239-47.

35. Ciesla G, Leader S, Stoddard J. Antibiotic prescribing rates in the US ambulatory care setting for patients diagnosed with influenza, 1997-2001. Respir Med 2004; 98: 1093-101.

36. Poehling KA, Edwards KM, Weinberg GA, Szilagyi P, Staat MA, Iwane KM, et al. Influenza vaccine coverage rates and perceptions on vaccination in South Korea. J Infect 2007; 55: 273-81.

37. Daley AJ, Nallusamy R, Isaacs D. Comparison of influenza A and influenza B virus infection in hospitalized children. J Paediatr Child Health 2000; 36: 332-5.

38. Ruiz Laiglesia F, Zubizarreta García J, Agud Aparicio J, Villasante Claudio F, Ayensa Dean C, Sánchez N阮lares M, Torrubia Pérez C. Febrile syndrome in hospitalized patients. Am Med Interna 1992; 9: 367-71.

39. Dawood FS, Fiore A, Kamimoto L, Nowell M, Reingold A, Gershman K, Meek J, Halder J, Arnold KE, Ryan F, et al. Influenza-associated pneumonia in children hospitalized with laboratory-confirmed influenza, 2003-2008. Pediatr Infect Dis J 2010; 29: 585-90.

40. Falagas ME, Vouloumanouk EK, Baskouta E, Rafailidis PI, Polyzos K, Rello J. Treatment options for 2009 H1N1 influenza: evaluation of the published evidence. Int J Antimicrob Agents 2010; 35: 421-30.

41. Fiore AE, Fry A, Shay D, Guinovart L, Bree EE, Uyeki TM, Centers for Disease Control and Prevention (CDC). Antiviral agents for the treatment and chemoprophylaxis of influenza: recommendations of the Advisory Committee on Immunization Practices (ACIP). MMWR Recomm Rep 2011; 60: 1-24.

42. Jennings LC, Skopnik H, Burckhardt I, Hribar I, Del Piero L, Deichmann KA. Effect of rapid influenza testing on the clinical management of pediatric influenza. Influenza Other Respir Viruses 2009; 3: 91-8.