Role of Human Metapneumovirus, Influenza A Virus and Respiratory Syncytial Virus in Causing WHO-Defined Severe Pneumonia in Children in a Developing Country

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Abstract

Objective: The role of respiratory viruses in causing severe, life threatening pneumonia in children in developing countries is not well established. Our study aims to determine the role of human metapneumovirus (HMPV), influenza A virus and respiratory syncytial virus (RSV) in children, aged 6 weeks to 2 years, hospitalized with WHO defined severe pneumonia (tachypnea plus any general danger sign or chest in-drawing) at a public sector hospital in Karachi, Pakistan.

Methods: This study was conducted from November 2010 to September 2011 at Abbasi Shaheed Hospital, a large public tertiary care hospital in Karachi, Pakistan. Children admitted with WHO-defined severe pneumonia were enrolled and throat swabs were obtained to detect respiratory viruses using real time RT-PCR. Chest x-rays of all subjects were obtained and independently interpreted by two radiologists to diagnose radiologic pneumonia.

Results: 169 children were enrolled. HMPV was detected in 24 (14.2%), influenza A virus in 9 (5.3%) and RSV in 30 (17.8%) children admitted with severe pneumonia. Of 9 patients with influenza A, 8 tested positive for H1N1. Viral etiology was found in 18% of radiologically confirmed pneumonia. HMPV infections peaked in February and April, influenza A was prevalent in January, June and November and RSV infections were most prevalent from June to September.

Conclusion: HMPV, influenza A and RSV are common causes of WHO-defined severe pneumonia in hospitalized children in Karachi. Knowledge regarding the viral etiology of pediatric pneumonia and individual viral seasonality can help in the recommendation and implementation of appropriate management strategies.

Introduction

Pneumonia is the second leading cause of childhood mortality worldwide, with estimated 1.25 million deaths per year [1]. In Pakistan, the estimated community based incidence of pneumonia in children aged less than five years is 1–4 episodes per 100 children per year, which accounts for 13% of all under five deaths every year [2]. While the role of respiratory viruses including HMPV, influenza A and RSV is well established in causing self-limiting upper respiratory tract infections or mild pneumonia, their contribution in causing severe and radiologically proven pneumonia in developing countries is less clear [3]. This knowledge is critical to guide the development of preventive and therapeutic interventions against these viruses, and to make a case for the use of existing interventions like the influenza vaccine [4].

A challenge in finding the viral etiology of severe pneumonia in developing countries is the general unavailability of appropriate diagnostic facilities [5]. Even if diagnostic tests are available, they are restricted to high resource, private tertiary care hospitals, whereas most of the deaths due to pneumonia occur in the community or in public sector facilities [2]. To determine the etiology of severe, potentially fatal pneumonia in developing countries, studies need to be conducted in health care facilities which cater to the lower income population, which is at the highest risk for adverse health outcomes. We therefore conducted a longitudinal surveillance study at a busy public sector hospital in Karachi to determine the role of respiratory viruses in WHO-defined severe pneumonia, including radiologically proven pneumonia, in children 6 weeks to 2 years of age.

Methods

This was an 11 month long study conducted between November 2010 and September 2011 at Abbasi Shaheed Hospital in Karachi, Pakistan. Abbassi Shaheed is a public sector hospital that serves the residents of the northern part of Karachi with an estimated population of nearly 1 million. Karachi is located on the coast of Arabian sea and has a relatively mild climate. The latitude and longitude of Karachi are 24.8508° N, 67.0181° E. The city has two main seasons; summer and winter, and receives the monsoon...
rains from July to September. The humidity levels usually remain high from March to November, while very low in winter. This study covers both seasonal periods.

In this study, WHO defined criteria for severe pneumonia, i.e. tachypnea (respiratory rate >60/min in children <2 months, 50/min in children 2–12 months and >40/min in children >12 months) and chest indrawing or any other danger sign [6], were used to identify children 6 weeks to 2 years old, hospitalized at the Abbasi Shaheed Hospital. Children were excluded if they were previously enrolled in the study or if their parents refused to participate. This study received ethical approval from Ethics Review Committee of Aga Khan University, Karachi Pakistan. After written informed consent from the parents or guardians, a baseline questionnaire was completed and pharyngeal swabs were obtained by the study physician. These were transported to the Infectious Diseases Research lab (IDRL) at the Aga Khan University (AKU) for viral testing using real time RT-PCR. The proportion of severe pneumonia cases associated with HMPV, influenza A and RSV was determined. In addition, chest x-rays of all enrolled subjects were obtained and emailed to two radiologists at AKU for interpretation using WHO standard definition for radiologically confirmed pneumonia [7]. These two radiologists were not given any clinical information about the child, and were blinded to each other’s interpretations. A third radiologist was used in case of discordant interpretation of the first two radiologists [7]. Descriptive analysis was done to calculate median age of children, and proportion of respiratory virus etiology. Associations between clinical manifestations and etiological agents were analyzed using chi square test for categorical data and t-test for continuous data, considering a P-value <0.05 as significant value. Statistical analysis was calculated using Stata software (Version 11).

### PCR Analysis

A throat swab was obtained using a commercial flocked swab (Diagnostics Hybrid Inc.) in Universal Transport Medium (Diagnostics Hybrid Inc.) and was transported to the IDRL. RNA was extracted from frozen aliquots (stored at −80°C) using QIAamp Viral RNA Mini Kit (Qiagen). Later, real time RT-PCR assays were performed using primers and probes for HMPV [CDC Reference Number: I-036-06], RSV [8], influenza A and A(H1N1)pdm09 [9]. All samples were first tested for influenza A, and those testing positive for influenza A were subsequently tested for A(H1N1)pdm09. Influenza B was not tested. Positive and negative controls were used in each run, and influenza A positive samples were re-confirmed and serotyped at a reference laboratory at North American Medical Research Unit 3 in Cairo, Egypt or in the reference lab of National Institute of Health, Pakistan.

### Results

From November 2010 to September 2011, 387 children between the age of 6 weeks and 2 years with signs suggestive of severe pneumonia were screened. 247 (64%) of these were found to

| Table 1. A comparison of baseline characteristics of HMPV, Influenza A and RSV infected children. |
|-----------------|-----------------|-----------------|
| **Baseline characteristics**       | **HMPV* (n = 24)** | **Influenza A (n = 9)** | **RSV** (n = 30) |
| Age in months (median, IQR)       | 6(4–10)          | 10(6–11)         | 5(3–7)         |
| Male                           | 13 (54.2%)       | 3 (33.3%)        | 20 (69%)       |
| Smokers in the house            | 3 (12.5%)        | 5 (55.6%)        | 5 (16.7%)      |
| Number of household members (median, IQR) | 7.5(7–11) | 7 (6–8) | 6(5–11) |
| Ever Vaccinated                 | 21 (87.5%)       | 6(66.7%)         | 19(63.3%)      |

Abbreviations:
*HMPV: Human metapneumovirus.
**RSV: Respiratory syncitial virus.

| Table 2. A comparison of the clinical features of HMPV, Influenza A and RSV infected children. |
|-----------------|-----------------|-----------------|
| **Clinical features** | **HMPV** | **Influenza A** | **RSV** |
| **Positive** | **Negative** | **Positive** | **Negative** | **Positive** | **Negative** |
| **(n = 24)** | **(n = 145)** | **(n = 9)** | **(n = 160)** | **(n = 30)** | **(n = 139)** |
| Wheezing/noisy breathing | 8(33.3%) | 93(64.1%) | 0.003 | 3(33.3%) | 98(61.2%) | 0.082 | 18(64.3%) | 83(59.7%) | 0.682 |
| Chest indrawing | 16(66.7%) | 60(41.4%) | 0.021 | 6(66.7%) | 70(43.8%) | 0.179 | 12(40%) | 64(46%) | 0.546 |
| Unable to drink | 15(62.5%) | 49(33.8%) | 0.007 | 4(44.4%) | 60(37.5%) | 0.676 | 10(33.3%) | 54(38.9%) | 0.572 |
| Difficult to arouse/abnormally sleepy | 5(20.8%) | 45(31%) | 0.310 | 3(33.3%) | 47(29.4%) | 0.800 | 6(20%) | 44(31.6%) | 0.205 |
| Radiologically confirmed pneumonia | 2(8.3%) | 49(33.7%) | 0.01 | 1(11.1%) | 50(31.2%) | 0.190 | 6(20.7%) | 45(32.3%) | 0.197 |

Abbreviations:
*HMPV: Human metapneumovirus.
**RSV: Respiratory syncitial virus.

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be eligible for the study, out of which 169 (68%) children with
WHO-defined severe pneumonia were enrolled. The demograph-
ic characteristics of children who tested positive for viruses are
summarized in table 1. The median age of the study subjects was 6
months (IQR 3–9 months). At least one of the three viruses was
detected in 61 (36%) of the enrolled subjects. HMPV was detected
in 24 (14.2%), influenza A virus in 9 (5.3%) and RSV in 30
(17.8%) subjects. Out of the 9 patients with influenza A, 8 tested
positive for H1N1. The clinical features and risk factors among
children with each viral etiology is summarized in table 2. Inability
to drink was a prominent feature in HMPV (62.5%) and influenza
A (44.4%) infected patients, whereas wheezing (64.3%) was most
noticeable in RSV infected individuals. Wheezing was found less
frequently in patients with influenza A (33.3%) and HMPV
(33.3%). Antibiotics were used prior to admission in 69.6% of
positive patients. Majority of children infected with influenza A were found
to have smokers in the house (55.6%, p = 0.042).

Pneumonia was radiologically confirmed in 51 (30.1%) of the
study subjects. Of these, 9 (18%) were positive for at least one of
the three viruses tested. HMPV was detected in 2 (4%) influenza
A in 1 (2%) and RSV in 6 (12%) of the radiologically confirmed
pneumonia cases. Radiologically confirmed pneumonia was found
in 2/24 (8.3%) HMPV, 1/9 (11%) influenza A and 6/30 (20%)
RSV infected children.

The monthly distribution of each of the three viruses and the
corresponding temperature and precipitation data are shown in
figure 1. The highest number of rainy days was observed in the
months of July, August and September. The lowest temperatures
were seen from December to January and the highest from May to
July. Most infections with RSV were reported from June to
September 2011. The peak incidence of RSV was in July (n = 6,
50%) and September (n = 5, 50%). HMPV showed one major
peak in February (n = 10, 63%) and one minor peak in April
(n = 2, 25%), and was absent from May to November. Influenza A
showed three minor peaks, in January (n = 6, 17%), June (n = 1,
6%) and November (n = 2, 13%) and was not detected during the
rest of the year.

Discussion

This study is the first to describe the viral etiology of WHO
defined severe pneumonia in hospitalized children in Pakistan
using real time RT-PCR assays. We found that 36% of patients
with severe pneumonia were positive for HMPV, influenza A or
RSV. This finding is similar to other hospital-based etiological
data collected in Bangladesh, India, Gambia and central Australia
[10,11]. On average, viruses are generally isolated in approxi-
ately 30–50% of the cases of severe pneumonia hospitalized in
developing countries [12].

RSV was the predominant virus in our study. It was isolated
from 30 (17%) children, reinforcing the results of several studies in
other countries which have demonstrated a similar magnitude of
prevalence of this virus in childhood pneumonia [13,14]. In a
study conducted in 2006 in rural Bangladesh on children aged 0–
24 months, RSV was found in 21 out of 38 (55%) cases admitted
with pneumonia [15]. The Bangladesh study however, included
cases of non-severe pneumonia, which may account for the
difference. A study conducted in rural Kenya identified 16.5%
cases of RSV associated severe pneumonia, which is consistent
with our findings [14].

We found that 20% of RSV positive, severe pneumonia cases in
our study had radiologically confirmed pneumonia. This is lower
than what was found in a similar study in Bangladesh (66.7%)
[11,14]. Our strict radiological interpretation was perhaps a factor
in the lower proportion of radiologically confirmed pneumonia
seen in our study. Nonetheless, RSV positive cases had the highest
proportion of radiologically confirmed pneumonia (20%) as
compared to other viral causes in our study. i.e HMPV (0%) and
influenza A(11%). While the possibility of secondary bacterial
infection cannot be ruled out, our study shows that viruses are
often associated with radiologically confirmed pneumonia.

RSV epidemics generally occur during the winter and spring
months, though there is considerable variation [4]. A study
conducted in a refugee population in Thailand showed a highly
seasonal pattern of RSV infections, occurring predominantly in
October each year, just after the peak rainfall [16]. We found RSV
infections year round in our study with peaks from July to
September, perhaps due to the heavy rainfall observed during this
period. As shown in Fig. 1, the highest precipitation occurred in
the months of July, August and September, which corresponds
with the RSV peak incidence. This is an important association and
makes this season an important target for intervention. RSV
vaccination or prophylaxis, once available, may offer considerable
public health benefit for vulnerable population in this season.
HMPV was found in 14% of children hospitalized with severe pneumonia in our study. This is a higher burden compared to what was recently seen in a prospective 5 year surveillance study in the United States, where only 6% of the hospitalized children tested positive for HMPV [17]. A similar study conducted in Israel found HMPV in 13% of the children hospitalized with pneumonia [18]. Though HMPV seasonality is not well-defined, some studies point towards it being more prominent in the winter season. Majority of cases in the prospective surveillance conducted in the United States were found during the months of January through April [17]. A study in Italy showed that a high incidence of HMPV infection (25.3%) was observed during the 2005–2006 winter-spring season, whereas a much lower rate of infection (4.7%) was found during the next season [19]. As observed in figure 1, the lowest temperatures in Karachi were seen from the months of December to February, which parallel a relatively higher incidence of HMPV infections detected in our study.

The incidence of influenza-virus-associated severe pneumonia requiring hospitalization is important because influenza is an infection that can be prevented with vaccines [20]. Infection with influenza virus predisposes children to infections with other common organisms associated with severe illness, such as pneumococcal and staphylococcal pneumonia [21]. During the year our study was conducted, influenza A was found at a lower rate as compared to RSV and HMPV. The incidence of influenza A in our study (5%) was similar to many other studies conducted in various parts of the world. In a study conducted in El Salvador from 2008–2010, 608 cases of severe pneumonia were tested for respiratory viruses and 37 (6%) were positive for influenza virus [22]. A slight peak in the incidence of influenza in January and February in our study coincides with a lower temperature observed during these months, as seen in figure 1.

Our study has several limitations. This surveillance was conducted for 11 months and the seasonal pattern of respiratory viruses may vary from year to year. Multiple year surveillance is needed to establish seasonality of different respiratory viruses reliably. However, the seasonal pattern seen in our study is consistent with data from other regional countries. While testing for influenza, we only tested for influenza A and not for influenza B. We did not try to elicit bacterial etiology of severe pneumonia in our patients, though all our study subjects received antibiotics. Hence estimates of viral-bacterial co-infections cannot be made. Nonetheless, our study shows that either alone or as part of viral-bacterial co-infections, respiratory viruses play an important role in clinically severe pneumonia in children in Pakistan.

In summary, we found that a considerable proportion of WHO-defined severe pneumonia and radiologically proven pneumonia hospitalizations in children in Pakistan are associated with respiratory viruses. Respiratory viruses should be the focus of additional efforts to decrease pneumonia-associated morbidity and mortality in developing countries and defining the burden of these viruses using reliable and sensitive diagnostic tests is an important first step in this regard. Many children with typical respiratory viral symptoms are not brought to the hospital until a secondary bacterial infection develops. A preventive vaccine given at the correct time could prevent many secondary bacterial infections. While our study shows important trends, subsequent year surveillance is also needed to assess if this seasonal pattern is consistent in this region.

Author Contributions

Conceived and designed the experiments: AA AR SM AZ. Performed the experiments: AA AR FA. Analyzed the data: AA AR MZB FA. Contributed reagents/materials/analysis tools: AA AR MZB. Wrote the paper: AA AR MZB SM AZ.

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