Surveillance of respiratory viruses in the outpatient setting in rural coastal Kenya: baseline epidemiological observations

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Abstract
Background: Endemic and seasonally recurring respiratory viruses are a major cause of disease and death globally. The burden is particularly severe in developing countries. Improved understanding of the source of infection, pathways of spread and persistence in communities would be of benefit in devising intervention strategies.

Methods: We report epidemiological data obtained through surveillance of respiratory viruses at nine outpatient health facilities within the Kilifi Health and Demographic Surveillance System, Kilifi County, coastal Kenya, between January and December 2016. Nasopharyngeal swabs were collected from individuals of all ages presenting with acute respiratory infection (ARI) symptoms (up to 15 swabs per week per facility) and screened for 15 respiratory viruses using real-time PCR. Paediatric inpatient surveillance at Kilifi County Hospital for respiratory viruses provided comparative data.

Results: Over the year, 5,647 participants were sampled, of which 3,029 (53.7%) were aged <5 years. At least one target respiratory virus was detected in 2,380 (42.2%) of the samples; the most common being rhinovirus 18.6% (1,050), influenza virus 6.9% (390), coronavirus 6.8% (387), parainfluenza virus 6.6% (371), respiratory syncytial virus (RSV) 3.9% (219) and adenovirus 2.7% (155). Virus detections were higher among <5-year-olds compared to older children and adults (50.3% vs 32.7%, respectively; \(\chi^2(1) = 177.3, P=0.0001\)). Frequency of viruses did not differ significantly by facility (\(\chi^2(8) = 13.38, P=0.072\)). However, prevalence was significantly higher among inpatients than outpatients in <5-year-olds for RSV (22.1% vs 6.0%; \(\chi^2(1) = 159.4, P=0.0001\)), and adenovirus (12.4% vs 4.4%; \(\chi^2(1) = 56.6, P=0.0001\)).

Conclusions: Respiratory virus infections are common amongst ARI outpatients in this coastal Kenya setting, particularly in young
children. Rhinovirus predominance warrants further studies on the health and socio-economic implications. RSV and adenovirus were more commonly associated with severe disease. Further analysis will explore epidemiological transmission patterns with the addition of virus sequence data.

**Keywords**
Outpatient; Respiratory viruses; Surveillance; Real-time PCR; Nasopharyngeal samples; Coastal Kenya

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Introduction
Acute respiratory infection (ARI) is a major cause of morbidity and mortality globally, principally affecting young children and the elderly, and with majority of the burden occurring in low-resource settings\textsuperscript{1-2}. Viruses are recognized as major cause of both mild ARI and of severe acute lower respiratory tract infection (LRTI)\textsuperscript{3}. Respiratory syncytial virus (RSV), rhinovirus and influenza A are often identified as the most common viruses associated with ARI\textsuperscript{3-5}, but a wide range of viruses are to be found in ARI presentations to the hospital and outpatient settings\textsuperscript{6}. The changing landscape of ARI due to the widespread use of conjugated bacterial vaccines could lead to an increased prominence of viral causes of these illnesses\textsuperscript{6-7}.

Given this context, greater emphasis on the control of virus-associated ARI is likely. Vaccination as an intervention for the control of viral respiratory infections faces considerable hurdles. These include continuous or rapid pathogen evolution (e.g. influenza)\textsuperscript{8}, high serotype diversity (e.g. rhinovirus)\textsuperscript{9} or target populations not appropriate for current vaccines (e.g. RSV)\textsuperscript{1}. Consequently, innovation in vaccination strategies for these pathogens (e.g. targeting of schools and households), mass use of antivirals or non-pharmacological methods such as social distancing (e.g. school closures), are options to consider. Designing strategies to control respiratory viruses would be assisted by detailed knowledge of the patterns of spread, i.e. of distinct pathways of transmission in communities, at various social organizational levels from the household, school, local community, nationally and beyond country boundaries.

The present work forms part of a larger project under the title SPReD (Studies of the Pathways of transmission of Respiratory virus Disease), which aims to map patterns of spread of a range of respiratory viruses using epidemiological and nucleotide sequence data across different settings in Kenya. Here we present the baseline results of 1 year of respiratory virus epidemiological surveillance at health facilities within a well-defined coastal Kenyan population and a comparison with contemporaneous data on inpatient admissions to Kilifi County Hospital (KCH).

Methods

Study site
This study was conducted on the coast of Kenya, within the Kilifi Health and Demographic Surveillance System (KHDSS)\textsuperscript{12-14}. The KHDSS area was defined and mapped for demographic surveillance, clinical and epidemiological research by the KEMRI-Wellcome Trust Research Programme (KWTRP) in the year 2000. It is located in Kilifi County along the coastal fringe and covers an area of 891 km\textsuperscript{2}, 50 km north and south, and 30 km west, of the KCH. The KHDSS monitors a population of around 296,000 residents (2016 census) through household enumeration visits conducted every 4 months. The major economic activity of most residents is subsistence farming of maize, cassava, cashew nuts and coconuts, as well as goats and dairy cattle\textsuperscript{15}.

The KHDSS area has 21 public health facilities (including the KCH) receiving out-patients, which operate under the Kenya Ministry of Health (MoH). In total, nine of these facilities were selected for this study: Matsangoni, Ngerenya, Mtondia, Sokoke, Mavueni, Jaribuni, Chasimba, Pingilikani and Junju (Figure 1). The facilities were purposively selected to provide a broad representation across the geographical region, covering major road networks into the location and variation in population density. All specimen processing and testing was carried out at KEMRI-Wellcome Trust Research Programme laboratories in Kilifi.

Patient recruitment and specimen collection

Participant recruitment and specimen collection was integrated within the routine patient care at the nine selected outpatient facilities led by a resident clinician or nurse. Each facility had one or two sampling days per week, usually scheduled from Monday to Friday, between 9.00 am and 1.00 pm. On each sampling day, a study fieldworker stationed at the health facility, assisted by the resident clinician or nurse, would describe the study to the attending patients. Any person present with listed signs and symptoms of ARI would be asked to see the fieldworker for further screening and obtainment of consent as they await review by the nurse or clinician. Patients of any age presenting with one or more ARI symptoms of cough, sneezing, nasal congestion, difficulty breathing, or increased respiratory rate for age (as defined by the World Health Organisation\textsuperscript{16}) were eligible. New-borns aged less than 7 days and patients with ARI for more than 30 days were excluded. Written individual informed consent was sought from adult patients and parents/guardians of patients below 18 years. The study commenced in December 2015 with piloting before enhanced surveillance of 15 samples per site per week from January to December 2016.

The sample size of 15 samples per site per week was determined based on outpatient data on the number of respiratory infection cases seen per month per selected health facility and our previous experience with inpatient surveillance for respiratory viruses at KCH. To optimize the study power to detect a diversity of the respiratory viruses, we estimated to collect approximately 7000 NPS specimen within a 1-year study period. Out of the 7000 NPS, we expected 1050 (15\%), 700 (10\%), 490 (7\%) and 940 (7\%) specimens to be positive for rhinoviruses, RSV, coronaviruses, and influenza viruses, respectively.

The selection of participants each week was on a ‘first-come first-served’ basis as they presented to the health facility on the set sampling days. A standardised questionnaire (Supplementary File 1) was used to collect biodata, presenting symptoms as well as the treatment provided which was entered directly into a study database using a computer tablet. A nasopharyngeal swab (NPS) was collected from each participant by inserting a sterile nylon-flocked plastic-shafted swab (503CS01, Copan Diagnostics, Flocked Swab Technologies, Italy) into one nostril to a distance where the tip located in the deep nasopharynx and was twisted 3 times before it was gently removed (in total taking about 10 seconds)\textsuperscript{17}. The NPS sample was stored in universal virus transport media (Copen Diagnostics, USA) and kept at approximately 8°C in an ice-packed cool box for return to KWTRP within 4 h of collection.
Ethical considerations
All individuals, parents and guardians gave written informed consent for themselves or their children to participate in this study. The study was approved by the KEMRI-Scientific and Ethical Review Unit (SERU# 3103) and the University of Warwick Biomedical and Scientific Research Ethics Committee (BSREC# REGO-2015-6102).

Laboratory procedures
NPS collections received at KWTRP virology laboratory were stored in 2-ml vials at −80°C until use. Using previously described methods, RNA was extracted from the respiratory specimens by Qiacube HT using an RNeasy extraction kit (Qiagen, Germany) and screened for RSV (A and B), rhinovirus (HRV), human coronaviruses (OC43, NL63, E229), influenza viruses (FLU-A, B, and C), parainfluenza viruses (PIV 1-4), adenovirus (ADV) and human metapneumovirus (HMPV), using a multiplex real-time PCR assay system. Samples with cycle threshold (Ct) of <35.0 were defined as positive for the target virus. Residual NPS samples were stored at −80°C.

Statistical analysis
Statistical analysis was conducted using STATA version 13.1 (College Station, Texas). Summary statistics were produced for the data to give the proportions of virus positives by age and by location. Comparative data was obtained from the paediatric ward of KCH for patients aged <5 years admitted with acute LRTI from a contemporaneous respiratory virus surveillance. Chi-squared and Fisher’s exact tests were used to test associations of virus occurrence with age, calendar month, facility, setting (outpatient or inpatient) and other demographic characteristics. Frequency distribution graphs were generated for all virus targets. Graphs for temporal patterns for each virus were generated.

Figure 1. A map of the Kilifi Health and Demographic Surveillance System (HDSS) area, coastal Kenya, expanded from a map of Kenya, showing population density (person per Km²) and the health facilities where the study was conducted in 2016. The red circles show the nine participating health facilities while the green markers show the other public health facilities within the KHDSS area.
Baseline characteristics of participants and sites
A total of 5647 participants were recruited from January to December 2016. The median age was 4 years (interquartile range (IQR): 1–15 years), 53.6% (n=3029) were children under 5 years of age, and 42.3% (2389) were male. The frequency of distribution of symptoms among participants by site, age and virus target are shown in Table 1. Although the proportions of each symptom by health facility, age category and virus target, were often statistically significantly different (Table 1), the rank

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Fever, %</th>
<th>Chest indrawing, %</th>
<th>Crackles, %</th>
<th>Wheeze, %</th>
<th>Cough, %</th>
<th>Nasal discharge, %</th>
<th>Nasal flare, %</th>
<th>Difficulty breathing, %</th>
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χ² P value 0.0001 0.835 0.0001 0.001 0.0001 0.0001 0.0001 0.0001 0.0001 0.0001

Age Category

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<th>Fever, %</th>
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<th>Crackles, %</th>
<th>Wheeze, %</th>
<th>Cough, %</th>
<th>Nasal discharge, %</th>
<th>Nasal flare, %</th>
<th>Difficulty breathing, %</th>
<th>Total participants, n</th>
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<td>1.72</td>
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<td>91.97</td>
<td>78.78</td>
<td>4.21</td>
<td>12.62</td>
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<td>6–11 mths</td>
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χ² P value 0.0001 0.005 0.127 0.0001 0.019 0.0001 0.0001 0.0001

Virus Target

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<th>Crackles, %</th>
<th>Wheeze, %</th>
<th>Cough, %</th>
<th>Nasal discharge, %</th>
<th>Nasal flare, %</th>
<th>Difficulty breathing, %</th>
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<td>1</td>
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χ² P value 0.0001 0.02 0.0001 0.019 0.024 0.026 0.007

Sex

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χ² P value 0.0001 0.051 0.002 0.841 0.105 0.0001 0.159 0.129

Table 1. Distribution of respiratory symptoms among study participants from a study of 9 outpatient health facilities in Kilifi County coastal Kenya over the year 2016.
order of magnitude of each sign or symptom was approximately the same. A majority of participants (85.3%, n=4819), presented with symptoms of cough and nasal discharge. A history of fever was identified in n=3541 participants (62.7%), but declined with age from 10–19 years onwards. Symptoms indicative of lower respiratory tract involvement, crackles, chest indrawing and fast breathing for age were uncommon (1.2%, n=65). The number of participants in each facility for this study differed significantly by the month in which recruitment occurred ($\chi^2(11) = 78.26$, P=0.001). There were interruptions during the 5th to 13th of December 2016 due to a health workers’ industrial dispute and 1 week during the Christmas break that led to lower recruitment for that month. Only 56 participants were recruited in December from all the nine facilities out of the 585 expected per month.

**Virus detection**

The prevalence of NPS collections positive for one or more respiratory virus target was 42.1% (n=2380). The median age of the virus-positive patients was 2 years (IQR, 1–9). The proportion of virus-positive individuals differed significantly by age, sex and education status, but not by sampling site (Table 2). The prevalence of respiratory virus detections was higher among young children (<5 years) than older children and adults (≥5 years).

### Table 2. Characteristics of respiratory virus positive and negative participants from a study of 9 outpatient health facilities in Kilifi County coastal Kenya over the year 2016.

<table>
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<tr>
<th>Characteristic</th>
<th>NPS virus positive(n)</th>
<th>NPS virus negative(n)</th>
<th>Total(n)</th>
<th>P value</th>
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<tr>
<td></td>
<td>(n=2380)</td>
<td>(n=3267)</td>
<td>(n=5647)</td>
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<td>Age in years</td>
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<td>Median (IQR*)</td>
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<td>5 (1-20)</td>
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IQR, Interquartile range; Chi2 test P value
(50.3% vs 32.7%, respectively; \( \chi^2 (1) = 177.3, P=0.0001 \)). Infants aged below 6 months with ARI constituted 9.3% (n=523) of the participants, and 58% (302) of these had one or more respiratory virus targets detected. Of all virus-positive individuals, 70% (1665/2380) were not in any form of education, with the vast majority of these pre-school, i.e. 83.2% (1385/1665). During study piloting (month of December 2015), respiratory samples from 153 participants were collected; 24.2% (n=37) were children under 5 years of age and 37.9% (n=58) were positive for one or more virus.

The most common respiratory viruses detected in this study were rhinovirus 18.6% (n=1050), influenza virus 6.9% (n=390), coronavirus 6.8% (n=387), parainfluenza virus 6.6% (n=371), RSV 3.9% (n=219) and adenovirus 2.7% (n=155). The frequency of viruses did not differ significantly by facility of recruitment (\( \chi^2(8) = 13.38, P=0.072 \) (Table 2). The distribution of proportions of viruses detected from each health facility were similar, with rhinovirus being the most common virus detected from all the sites (Figure 2). Furthermore, for each virus group, the proportion of all positive individuals were distributed roughly evenly across all health facilities, with the exception of HMPV (Figure 3). Note for comparison that during the pilot phase of December 2015 during which 56 samples were collected, influenza virus type A; 37.9% (n=22), RSV; 20.6% (n=12) and rhinovirus; 17.2% (n=10) were the most commonly detected viruses.

The age distribution of detections showed variation between viruses (Figure 4). For example, RSV B was most frequent among participants 0 to 23 months, adenovirus in children 6 to 35 months, and influenza B in those aged 5 to 19 years.

Multiple viruses were detected in 4.4% of the participants (n=246), with 2, 3 and 4 co-infecting viruses detected in n=221, n=19 and n=6 of the participants, respectively. Rhinovirus occurred in 153 participants with multiple viruses; coronaviruses in 131 participants; adenovirus in 55 participants; parainfluenza virus type 3 in 36 participants and RSV B in 28 participants. Rhinoviruses and adenoviruses co-infected with all the other 14 target viruses while RSV group B was found not to co-infect with all, except HMPV.

Only 1.4% of the participants (n=33) with a virus-positive NPS sample were referred to the KCH for specialized management. Treatment given to participants with ARI symptoms who tested positive for any of the virus targets were: antibiotics 81.4% (n=1947), antihistamines (chlorphenamine maleate) 60.4% (n=1437) and paracetamol 86% (n=2045). The most common antibiotic drugs prescribed were amoxicillin (39.7%, n=945) and trimethoprim/sulfamethoxazole (35%, n=834). Amoxicillin/ clavulanic acid (0.3%, n=8) and ciprofloxacin (0.7%, n=16) were less commonly prescribed.

Seasonality of the detected viruses

We observed a seasonal pattern in occurrence for some of the viruses (Figure 5). Rhinovirus and adenovirus appeared to occur throughout the year. Influenza virus occurrence during the year differed between the types. Influenza type B, the most commonly detected influenza type, occurred predominantly between March and August. Influenza type C occurred mostly between September and December, while influenza type A was detected most commonly in January, February, November and December. RSV group B was predominant over RSV group A; cases for both groups arose predominantly in the first 4 months of the year. Coronavirus OC43 occurred mostly during the months of June to August and E229 at the end of the year; NL63 occurred least in the third quarter of the year. Seasonal patterns for parainfluenza viruses and HMPV were difficult to discern.

Comparison of inpatient vs outpatient virus detections

A comparison of the distribution of viruses between the outpatient and the inpatient setting among children under 5 years of age is shown in Figure 6. A total of 49.0% NPS samples (n=282) from 575 inpatients with LRTI aged under 5 years were positive for one of the target respiratory viruses. RSV was the most common virus detected among hospitalized cases of ARI, with significantly higher prevalence 22.1% (n=127) than for outpatients 6.9% (182) (\( \chi^2(1)=159.4, P=0.0001 \)). Adenovirus was also of high prevalence and significantly more common 12.4% (n=71) than in outpatients 4.4 % (134) (\( \chi^2(1)=56.6, P=0.0001 \)). Rhinovirus was at high prevalence in both settings, but more so in the outpatients. Coronavirus OC43 and parainfluenza virus type 4 were more prevalent in outpatient than inpatients. All remaining viruses, though proportionally less common, showed higher prevalence in the outpatient setting. The distribution of Ct values for each virus target was similar for both inpatient and outpatient samples.

Discussion

In this study on the coast of Kenya, we describe the frequency of occurrence, and the spatial and temporal distribution of 15 viruses associated with respiratory illness in the outpatient setting. We also compare the distribution of viruses associated with respiratory infection among children under 5 years of age in the outpatient and inpatient settings.

To provide baseline data for investigating the transmission patterns and pathways of spread for common respiratory viruses, we used molecular techniques to detect viruses circulating throughout a well-defined population of nearly 300,000 inhabitants in an area of ~900 km² in rural coastal Kenya. We observed a prevalence of 42% virus infection among the selected patients presenting with symptoms of ARI in outpatient departments of nine health facilities. These findings indicate that viruses are prominently associated with respiratory tract infections of sufficient severity for individuals to seek medical attention within this community.

Consistent with observations elsewhere\(^{23,24}\), most of the ARI outpatients in this study were young children under 5 years; 50.3% of the participants in this age group had a virus positive NPS sample. This demonstrates that the under 5’s are a highly vulnerable age group for medically important ARI. The results are comparable to those from the outpatient health facilities in refugee camps and at a national referral hospital, in Kenya, where
Figure 2. Percentage of nasopharyngeal swab samples positive for each of 15 respiratory virus targets for the nine health facilities (all facilities together and each individually) from ARI surveillance in the Kilifi Health and Demographic Surveillance System, Kenya, January to December 2016. RSVA, RSV group A; RSBV, RSV group B; HRV, human rhinovirus; PIV1-PIV4, parainfluenza virus types 1–4; ADV, adenovirus; OC43, human coronavirus OC43; NL63, human coronavirus NL63; E229, human coronavirus E229; FLUA-FLUC, influenza viruses types A-C; HMPV, human metapneumovirus.
The most frequently detected virus in this study was rhinovirus. This is in accordance with previous community-based studies, for example the Tecumseh project in Michigan. Serological and molecular epidemiological studies show rhinoviruses to exist as many types (currently over 160), with little cross-type protection (amongst those classified immunologically), which may explain the high prevalence and absence of seasonality in respiratory illness caused by rhinovirus infection in the community. Persistence of rhinovirus might be ascribed to the frequent introduction of new virus strains into the community unconstrained by prior circulation of other types.

In this study, we did not see any major difference in distribution of viruses across the health facilities. This could have been attributed to the relatively small size of the demographic surveillance system area (891 km²). This allows for the possibility of population mixing and frequent interactions, especially during social events, leading to the rapid spread of viruses across the KHDSS area. However, definitive understanding of the temporal and spatial patterns of spread requires the addition of sequence data to infer relatedness of circulating viruses.

Influenza, RSV and coronavirus exhibit a clear seasonality pattern in occurrence, whereas rhinovirus and adenovirus are detected throughout the year. Of note is that during the first quarter of the year, other than rhinovirus, RSV is predominant amongst the detected viruses. Currently little is known about the mechanisms underlying virus dominance, interaction, co-existence and competition. Studies are warranted to investigate occurrence and interactions of multiple respiratory viruses in the nasopharynx of the individual over time (i.e. across the seasons), and to explore the possible effect of eliminating a virus such as
Figure 4. Age-distribution of detections of 15 virus targets in nasopharyngeal swabs (NPS) identified from ARI surveillance at nine health facilities in the Kilifi Health and Demographic Surveillance System, Kenya, January to December 2016. The primary Y axis shows frequency while secondary Y axis shows proportion in each age group. RSVA, RSV group A; RSVB, RSV group B; HRV, human rhinovirus; PIV1–PIV4, parainfluenza virus types 1–4; ADV, adenovirus; OC43, human coronavirus OC43; NL63, human coronavirus NL63; E229, human coronavirus E229; FLUA-FLUC, influenza viruses types A–C; HMPV, human metapneumovirus.
Figure 5. The distribution by month of the proportion of virus-positive nasopharyngeal swab samples over the period January to December 2016 for each of 15 virus targets, obtained through ARI surveillance at nine health facilities in the Kilifi Health and Demographic Surveillance System, Kenya. Secondary Y axis records number of samples collected from recruits per month. RSVA, RSV group A; RSVB, RSV group B; HRV, human rhinovirus; PIV1-PIV4, parainfluenza virus types 1-4; ADV, adenovirus; OC43, human coronavirus OC43; NL63, human coronavirus NL63; E229, human coronavirus E229; FLUA-FLUC, influenza viruses types A-C; HMPV, human metapneumovirus.
Figure 6. Virus surveillance comparison between inpatient and outpatient facilities for children under 5 years in the Kilifi Health and Demographic Surveillance System (KHDSS), Kenya, 2016. Panel A compares the proportion of nasopharyngeal swab samples positive for each of the 15 virus targets in samples collected from severe pneumonia admissions to Kilifi County Hospital (grey bars) and from outpatients presenting to nine health facilities (black bars). Violin plots show the distribution (median, IQR) for detectable rPCR cycle threshold (Ct) values (i.e. Ct<=40) from respiratory samples for KHDSS outpatients (B) and for KCH inpatients (C). Threshold used for determining positive and negative samples shown by dashed line (Ct=35.0). RSVA, RSV group A; RSVB, RSV group B; HRV, human rhinovirus; PIV1-PIV4, parainfluenza virus types 1-4; ADV, adenovirus; OC43, human coronavirus OC43; NL63, human coronavirus NL63; E229, human coronavirus E229; FLUA-FLUC, influenza viruses types A-C; HMPV, human metapneumovirus.
RSV through vaccination. Such information will be useful to guide policy on priority respiratory viruses to focus for intervention.

We also find the wide use of antibiotics to treat majority of patients presenting with symptoms of ARI most likely caused by viruses. This raises concern over antimicrobial stewardship, with increased risk of antimicrobial resistance to first-line antibiotic agents and unnecessary use of expensive second-line antibiotics in treating mild acute respiratory disease.

In contrast to the outpatient setting, where rhinovirus is the most common virus associated with ARI among children under 5 years; in the inpatient setting, RSV and adenovirus are the leading cause of severe respiratory illness. From the long-term in-patient surveillance at KCH the observation for RSV is not unusual, but 2016 had an unusually high occurrence of adenovirus cases (data not shown for other years). The pattern in distribution of virus load (equated to Ct values) suggests that the cut off of 35.0 for the MPX real time PCR diagnostic method is generally suitable for all viruses, both in outpatient and inpatients (i.e. irrespective of disease severity and possible viral load), excepting for HMPV, PIV-1 and CoV 22E in outpatients and PIV-2 in inpatients and outpatients, where sensitivity may be an issue and a contributor to low prevalence in this study.

The major limitation of this study is that data are for 1 year only and caution should be applied in inferring seasonal patterns. This is exacerbated by the low numbers of participants recruited in December due to countrywide industrial action by nurses. In addition, there is competition for viruses due to the design of sampling we used, of selected 15 samples per facility per week. An epidemic for one target virus might influence the proportions of other viruses observed but this might not necessarily imply seasonality. The detection of multiple viruses in one individual makes it difficult to determine the viral pathogen responsible for the respiratory illness at the time of recruitment. Only a sub-sample of all ARI presentations were recruited and the underlying denominator not recorded, which prevents an estimation of community incidence of presentations that would be useful for comparative purposes.

Conclusions
In a sample of 5647 participants, about 40% of the ARI outpatient visits to the Kenyan Coast were associated with respiratory virus infection. Virus ARI is predominant among children <5 years, and relatively uncommon amongst school-going children. Rhinoviruses, influenza viruses, parainfluenza viruses, coronaviruses and RSV are most commonly associated with ARI over one year of community surveillance, whereas RSV and adenoviruses are the predominant respiratory virus among hospitalized patients with ARI. Virus occurrence (temporal and age-related) is similar across all facilities within the KHDSS area. Studies of the socio-economic implications of this burden are warranted, especially for rhinovirus infections that predominated. Further analysis of virus sequence data will delineate patterns of spread of viruses causing ARI illness in this setting.

Data availability
The replication data and analysis scripts for this manuscript are available from the Harvard Dataverse: https://doi.org/10.7910/DVN/ZX7NS4#. As the dataset contains potentially identifying information on participants, it is stored under restricted access. Details on eligibility for access and a request form are available from http://kemri-wellcome.org/about-us/#ChildVerticalTab_15 for consideration by our Data Governance Committee (dgc@kemri-wellcome.org).

Competing interests
No competing interests were disclosed.

Grant information
This work was supported by the Wellcome Trust (102975, 203077).

The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Acknowledgements
We thank all the study participants for their contribution of study samples and data. We also thank the Dispensary /Health Centre management committees for allowing us to conduct the study within their health facilities. We are grateful to the field study team: Grace Mwange, Timothy Mwaringa, Emily Nyale, Mercy Mzungu, Lucy Kazungu, Clinton Mwatata and Robert Garero who dedicated their time to make sure recruitment of participants is successful. We acknowledge the laboratory staff of KEMRI Wellcome Trust Research Programme, Virus Epidemiology and Control Research Group for the screening of all samples and uploading the data for analysis. This paper is published with the permission of the Director of KEMRI.

Supplementary material
Supplementary File 1. Biodata questionnaire used in this study.

Click here to access the data.
References


Open Peer Review

Current Peer Review Status: ✔️ ✔️

Version 1

Reviewer Report 06 August 2018

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David P. Moore

1 Medical Research Council, Respiratory and Meningeal Pathogens Research Unit, University of the Witwatersrand, Johannesburg, South Africa
2 Department of Paediatrics, Chris Hani Baragwath Academic Hospital and University of the Witwatersrand, Johannesburg, South Africa

This study on the molecular epidemiology of respiratory viral illness in community and hospital settings in coastal rural Kenya is an important addition to the growing literature on respiratory viral epidemiology in sub-Saharan African children. Future publications from this group of researchers is eagerly anticipated, particularly with regards tracking temporality and seasonality of viral prevalence over multiple years, and detailing the frequently-occurring viral-viral co-detections in this cohort.

Kenya is generally known to be a low human immunodeficiency virus type 1 (HIV) prevalence setting. It would, however, be useful to factor in HIV status of study participants in future studies from this well-delineated catchment area.

Point for clarification in the analysis:

1. In the first sentence of the Results section, 5647 participants were enrolled – considerably fewer than the 7000 participants sought in the sample size calculation: how this ‘under-count’ impacted on the ability to establish certainty of statistical inference needs to be mentioned in the Discussion section;

2. Figures 6 B and 6 C seem superfluous, as there is no text which describes the statistical comparison between pathogen density and inpatient or outpatient status. Suggest add in one or two sentences regarding this analysis in the last paragraph of your Results section.

Recommended minor stylistic adjustments:

- In the first sentence of the Introduction, rather state ‘and with the majority of the burden’;
- In the second sentence of the Introduction, rather state ‘are recognized as the major cause’;
In the first sentence of the second paragraph of the Introduction, rather state ‘Given this context, it is likely that greater emphasis will be placed on the control of virus-associated ARI.’;

3. In order to strengthen the important assertions made in sentence four of the second paragraph of the Introduction, especially with regards ‘social distancing’, I would suggest that the authors reference this sentence;

4. In the last sentence of paragraph three of the Introduction, suggest reword as ‘Here we present the baseline results of 1 year of respiratory virus epidemiological surveillance at health facilities within a well-defined coastal Kenyan population, and compare them with contemporaneous data from inpatient admissions to Kilifi County Hospital (KCH).’;

5. In the fourth sentence of the first paragraph of Patient recruitment and specimen collection, reword as ‘they awaited review’;

6. In the last sentence of the second paragraph of Patient recruitment and specimen collection, I would suggest that the authors reference their proportional split between respiratory viruses;

7. In the second sentence of the paragraph on Statistical analysis, suggest reword as ‘proportion of virus positives’;

8. In the third sentence of the paragraph on Statistical analysis, insert a space between the preceding full stop and the words ‘Chi-squared’;

9. In Table 1, it makes logical sense to reorder the columns according to order in which a clinician would examine a patient, starting with history and moving to physical examination: I propose that an optimised order would be: ‘Cough’, ‘Difficulty breathing’, ‘Fever’, ‘Nasal discharge’, ‘Nasal flare’, ‘Chest indrawing’, ‘Crackles’, ‘Wheeze’;

10. On page 6, in the second last sentence of column one (starting ‘The number of participants in each facility...’ gives a \( \chi^2 \) degree of freedom of 11: is this in fact correct? My reading is that this should be 8 (in view of 9 included outpatient facilities having been included in the sampling strategy). Please correct if appropriate;

11. Also, close the gap between ‘\( \chi^2 \)(8/11)’ and ‘\( \chi =78.26 \);’

12. In the first sentence under Virus detection, reword as ‘respiratory virus targets’;

13. In Table 2, add a space between ‘positive’ and ‘(n)’ in the second cell in the top row, and between ‘Total’ and ‘(n)’ in the sixth cell. Widen the Table slightly to accommodate a slightly wider columns;

14. In the first line on page 7, suggest close the gaps between ‘\( \chi^2 \); ‘(1)’, and ‘\( \chi =177.3 \);’

15. On page 7, the last sentence of the first paragraph (which describes the pilot phase of the study in December 2015) does not seem to add to the ‘meat’ of the results: suggest delete this sentence;
16. In the last sentence of the fourth paragraph on page 7, suggest reword to 'except with HMPV.';

17. In the second sentence of the second column on page 7, suggest reword influenza A activity to ‘while influenza A had biphasic activity in January/February and November/December’;

18. In the first sentence under the heading **Comparison of inpatient vs outpatient virus detections**, suggest reword to ‘inpatient settings’;

19. In the fourth sentence under the heading **Comparison of inpatient vs outpatient virus detections**, suggest close the gap between ‘χ²(1)’ and ‘=56.6’;

20. In the last line on page 7, suggest qualify as ',', both in Kenya,’;

21. In the legend to **Figure 2**, replace the full stop after ‘RSVB’ with a comma and change ‘E229’ to ‘229E’;

22. In the legend to **Figure 3**, replace the lower case ‘e’ in ‘229e’ with an upper case ‘E’;

23. In the first line on page 8, suggest reword as ‘under 5 years old with ARI were 49.8% and 54%, respectively.’;

24. In the first sentence of the second paragraph on page 8, place the full stop outside the closing parentheses: ‘kindergarten or primary).’;

25. In the first sentence of the third paragraph on page 8, suggest change wording from ‘in accordance’ to ‘in keeping’;

26. In the legend to **Figure 4**, replace the full stop after ‘RSVB’ with a comma and change ‘E229’ to ‘229E’;

27. In **Figure 5**, the same seasonal (grey) line plot is included in each of the graphs and represents the total number of samples collected per month in the study: this should be mentioned in the Figure legend;

28. In legend to **Figure 5**, replace the full stop after ‘RSVB’ with a comma and change ‘E229’ to ‘229E’;

29. In **Figure 6 A**, suggest quote the percentage ‘49%’ to one decimal place, and add a space between ‘Inpatients:’ and ‘Virus’ in the text superimposed on the plot area;

30. In the legend to **Figure 6**, replace the full stop after ‘RSVB’ with a comma and change ‘E229’ to ‘229E’;

31. In the first sentence of the second paragraph on page 13, suggest reword as ‘We observed a high prevalence of antibiotic prescribing to outpatients presenting with ARI most likely cause by viruses.’;

32. In the second sentence of the third paragraph in page 13, suggest reword to ‘Based on long-
term in-patient surveillance';

33. In the third sentence of the third paragraph in page 13, suggest replace ‘MPX’ with ‘multiplex’, as the abbreviation ‘MPX’ has not been defined before in the text;

34. In the third sentence of the third paragraph in page 13, suggest reword to ‘both in outpatients and inpatients’;

35. In the third sentence of the third paragraph in page 13, correct ‘CoV 22E’ to ‘CoV 229E’;

36. In the second sentence of the second column on page 13, suggest reword as ‘respiratory virus among children hospitalized with LRTI’.

**Is the work clearly and accurately presented and does it cite the current literature?**

Yes

**Is the study design appropriate and is the work technically sound?**

Yes

**Are sufficient details of methods and analysis provided to allow replication by others?**

Yes

**If applicable, is the statistical analysis and its interpretation appropriate?**

Yes

**Are all the source data underlying the results available to ensure full reproducibility?**

Yes

**Are the conclusions drawn adequately supported by the results?**

Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:**

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Reviewer Report 31 July 2018

https://doi.org/10.21956/wellcomeopenres.15964.r33577

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Ting Shi
University of Edinburgh, Edinburgh, UK

This is a well described manuscript reporting the epidemiological data of respiratory viruses at nine outpatient health facilities in rural coastal Kenya. More than half (53.7%) of participants with ARI symptoms were from children younger than 5 years. The most common respiratory viruses detected were rhinovirus, influenza virus, coronavirus, parainfluenza virus, respiratory syncytial virus (RSV) and adenovirus. In comparison group with young children admitted to hospitals, the frequency of RSV and adenovirus was significantly higher. However, a few points need to be clarified.

1. Would be nice to know the severity of participants with multiple viruses and see whether there is an association. What about the multiple viruses infection in inpatients?
2. Newborns aged less than 7 days were excluded. Would this affect the detection rate of respiratory viruses in infants? Would this underestimate the burden?

The detection of viruses in participants with ARI symptoms does not necessarily indicate a causal attribution of these viruses (association ≠ causality). Would be nice to add a few sentences discussing this point.

Is the work clearly and accurately presented and does it cite the current literature?
Yes

Is the study design appropriate and is the work technically sound?
Yes

Are sufficient details of methods and analysis provided to allow replication by others?
Yes

If applicable, is the statistical analysis and its interpretation appropriate?
Yes

Are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions drawn adequately supported by the results?
Yes

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Epidemiology, global health, child health, respiratory disease

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.
Thank you for taking time to review this manuscript. Please find responses to the issues raised written in italics font below.

1. Would be nice to know the severity of participants with multiple viruses and see whether there is an association. What about the multiple viruses infection in inpatients?

*The outpatient participants were all mild cases presenting for care. Occurrence of multiple respiratory viruses infection did not change severity of symptoms among children hospitalized with pneumonia or severe pneumonia. By using pediatric inpatient data and comparing individuals with multiple viruses and those with presence of only one virus, there was no statistically significant difference in disease severity distribution (p=0.075).*

2. Newborns aged less than 7 days were excluded. Would this affect the detection rate of respiratory viruses in infants? Would this underestimate the burden?

*We excluded newborns less than 7 days because of difficulties associated with collecting a nasopharyngeal swab sample among neonates. Care was given priority to sick newborns of less than 7 days brought to the outpatient facility and not research. However, for the inpatient surveillance, newborns to 1 day old were included. We are not sure if this would have underestimated the detection rates of respiratory viruses or disease burden among infants in the outpatient setting.*

3. The detection of viruses in participants with ARI symptoms does not necessarily indicate a causal attribution of these viruses (association ≠ causality). Would be nice to add a few sentences discussing this point.

*Agreed. This was an observational study and had several limitations since we did not test for other pathogens likely to cause respiratory illness.*

**Competing Interests:** None